



WafoL– a Wafo module for Analysis of Random Lagrange Waves

Tutorial for WafoL version 1.2

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Information on WAFO and Wafol

- This is the second version of a tutorial for how to use the MATLAB WAFO-module `Wafol` for analysis and simulation of random Lagrange waves. The module is (to be) included in the WAFO toolbox in the folder `lagrange`. The module consists of a number of MATLAB m-files and it requires a standard MATLAB setup together with the WAFO25 toolbox. In some example we use routines from other MATLAB toolboxes, like the `signal` toolbox.

Compared to Version 1.0.2 from April 2015, the present Wafol version 1.2 contains routines for generation of non-linear, second order 3D Stokes-Lagrange waves. They are based on MATLAB and Fortran routines written by Marc Prevosto, IFREMER, Brest, France. The routines have been adapted to work together with WAFOL. The option to use the MATLAB Parallel Computing toolbox has also been introduced in this routine.

New in Ver 1.2 are also some hints on how to export wave movies from WAFOL.

- On WAFOL:
 - Download the `Wafol.zip` file, unzip it to a suitable folder, and add to your MATLAB-path. Be sure to put it first in the path to avoid name conflicts.
 - The MATLAB code used for the examples in this tutorial can be found in three script files `WafolChx.m`. Some editing has been made on figures, and some simulations have been run with more replicates than in `WafolChx.m`.
 - The standard for random number generator initiation was changed in MATLAB 2011. The simulation routines in WAFOL work with both old and new standard.
 - Some changes in the graphics system were made with MATLAB 2014b, which caused warnings in some plotting routines in WAFO and WAFOL. An update of `Wafol25` will come, but in the meantime you can comment out all occurrences of the command `mkdeleteproxy` in plotting routines to avoid the warnings.
 - There are two routines for 2^{nd} order Stokes-Lagrange waves, `spec21dat3DM` and `spec21dat3DP`. They use a compiled Fortran routine that comes in two versions, for 32-bit and for 64-bit MATLAB called `partkern.mexw32` and `partkern.mexw64`. They come with the `Wafol.zip` and you have to put them in the appropriate folders in your `Wafol/exec` folder.

- The routine `spec21dat3DP` is a version of `spec21dat3DM` adapted for parallel processing with the Parallel Computing Toolbox in MATLAB.
 - If you have many MATLAB-toolboxes installed, name-conflicts may occur – for example, a call to `smooth` will cause errors with wrong path priority. Solution: arrange the MATLAB-path with `WAFO` first. The routine `initwafo` can do that for you.
 - Valuable comments on the tutorial and the use of `WAFOL` by associate professor Ying-guang Wang at Shanghai Jiao Tong University, Department of Naval Architecture and Ocean Engineering, are gratefully acknowledged.
 - Comments on the tutorial and on the routines are appreciated to `georg@maths.lth.se`
- On `WAFO`:
 - The word `WAFO`, when used in path specifications, means the full name of the `WAFO` main catalogue, for instance
`C:/wafo25`
 - `WAFO` is built of modules of platform independent MATLAB m-files and a set of executable files from `C++` and `Fortran` source files. These executables are platform and MATLAB-version dependent, and they have been tested with recent MATLAB and WINDOWS installations.
 - `WAFO` Version 2.5, was released in beta version in January 2009 and in stable version in February 2011. It can be downloaded from
<https://github.com/wafo-project>
There you also find `PYWAFO`, a Python version. The old repository,
<http://code.google.com/p/wafo/>
does not work any more.
Older versions of the toolbox can be downloaded from the `WAFO` homepage
<http://www.maths.lth.se/matstat/wafo/>
There you can also find links to exercises and articles using `WAFO`, and notes about its history.
 - For help on the toolbox, write `help wafo25`. Note, that in Windows, some of the routines in Chapter 4 do not work with MATLAB 2006 or earlier.
 - Comments and suggestions on `WAFO` are solicited — send to
`wafo@maths.lth.se`
 - The owners of the `WAFO` package are
Per Andreas Brodtkorb: `per.andreas.brodtkorb@gmail.com`
Georg Lindgren: `georg.lindgren@gmail.com`
Igor Rychlik: `igor.rychlik@gmail.com`
 - New contributor: `Marc.Prevosto@ifremer.fr`

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CHAPTER 1

2D Lagrange waves

1.1 The 2D Gauss-Lagrange model

1.1.1 The Gaussian model

The 2D Gaussian wave model is a stationary homogeneous Gaussian process $w(t, u)$ depending on time t and location u . It describes the height of the water surface at location u observed at time t . In WAFO the space coordinate $u = u_0$ is assumed fixed and $w(t, u_0)$ then resembles the waves observed by a stationary wave gauge, we denote them $w(t)$ for short.

The standard way in WAFO to generate a Gaussian time wave at location $u_0 = 0$ is as a sum of harmonics,

$$w(t) = m + \sum_{j=0}^N \sqrt{S_j} R_j \cos(-\omega_j t + \theta_j), \quad (1.1)$$

with independent Rayleigh distributed amplitudes $R_j = \sqrt{A_j^2 + B_j^2}$, A_j, B_j independent and standard normal variables. The phases Θ_j are random, uniformly distributed in $(0, 2\pi)$, cf. the WAFO-tutorial [14, Sec. 2.2.2]¹. The weight factors S_j are given by the one-sided spectral density $S(\omega), \omega \geq 0$, with frequency spacing $\Delta\omega$, $S_j = \Delta\omega S(\omega_j) = \Delta\omega S(j\Delta\omega)$. Note that both phases and amplitudes are random in the WAFO model. In the future we set the mean level to $m = 0$.

The Gaussian wave in time t and space u is similar to (1.1),

$$w(t, u) = m + \sum_{j=0}^N \sqrt{S_j} R_j \cos(\kappa_j u - \omega_j t + \Theta_j), \quad (1.2)$$

¹Note that the representation (1.1) has a minus sign for the frequency $\omega_j = j\Delta\omega > 0$ where the WAFO tutorial has a plus sign.

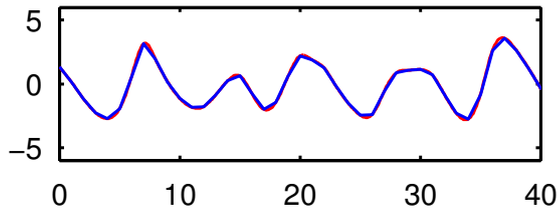


Figure 1.1: *Lagrange wave in time with slight front-back asymmetry.*

with wave-number κ_j related to the frequency ω_j through the depth dependent dispersion relation,

$$\omega^2 = g\kappa \tanh h\kappa, \quad (1.3)$$

with gravity g and water depth h . By convention in WAFOL we choose κ and ω to both be positive, leading to 2D waves moving “from left to right”.

1.1.2 The Gauss-Lagrange model

The stochastic Gaussian model only describes the random variations of the water surface. The Gauss-Lagrange model describes the combined vertical and horizontal movements of individual water particles. In WAFOL we only consider particles on the surface, and we further assume that there is no interaction between harmonics – this is “the 1st order model”.

The 2D Gauss-Lagrange wave model consists of two correlated Gauss processes, $w(t, u)$ and $x(t, u)$ which describe the vertical and horizontal movements of individual particles with time t . The space parameter u is called the *reference location*, indicating that it is the particle’s location “at rest” on the constant mean surface $m = 0$. The process $x(t, u)$ is the horizontal deviation of the particle from its reference location. A particle with original still water location $(u, 0)$ is, at time t located at $(u+x(t, u), w(t, u))$.

Here is a first example of how to generate a Lagrange wave in WAFOL.

```
S = jonswap; S.h=20;
opt=simoptset('dt',1);
[w,x]=spec2ldat(S,opt,'lalpha',1);
[L,L0] = ldat2lwav(w,x,'time',[],10);
subplot(211)
plot(L.t,L.Z); hold on
plot(L0.t,L0.Z,'r')
axis([0 40 -6 6]); hold off
```

Figure 1.1 shows a Lagrange time wave sampled at 1 Hz in blue and a spline smoothed version in red. The parameter `lalpha` with value 1 gives the waves a slightly steeper increase phase than decrease phase. The water depth is `S.h=20` [m].

The horizontal process is completely determined by the vertical process.² If $w(t, u)$

²In the most general form of a Lagrange model, extra independent random harmonics can be added to the terms in $x(t, u)$.

is given by (1.2), then

$$x(t, u) = \sum_{j=0}^N \sqrt{S_j \Delta \omega} \rho_j R_j \cos(\kappa_j u - \omega_j t + \Theta_j + \theta_j), \quad (1.4)$$

where ρ_j is called the *amplitude response* and θ_j is the *phase response* or “phase shift”.

The response functions

The horizontal process $x(t, u)$ is a *linear filtration* of the vertical process $w(t, u)$ and the filter is defined by a frequency/wavenumber dependent complex response function

$$H(\omega) = \rho(\omega) e^{i\theta(\omega)}.$$

The amplitude and phase responses in (1.4) are then obtained as

$$\rho_j = \rho(\omega_j), \quad \theta_j = \theta(\omega_j).$$

The response functions determine the non-Gaussian characters of the Lagrange waves, in particular crest-trough and front-back asymmetry. In the standard Lagrange model the filter response is depth and frequency dependent, given by

$$H_M(\omega) = i \frac{\cosh(h\kappa)}{\sinh(h\kappa)}, \quad (1.5)$$

leading to a frequency independent phase shift of $\pi/2$ between vertical and horizontal movements. The subscript M in (1.5) stands for *Miche waves*; see [10]. This choice of response function results in waves with crest-trough asymmetry, with more peaked crests and shallower troughs compared to the Gaussian waves. However, Miche waves are front-back statistically symmetric; wave fronts and wave back distributions are mirror images of each other; [1, 2].

In order to give wave front-back asymmetry the response function must give a frequency dependent phase shift. In WAFOL this is realized by adding a term α/ω^2 to the Miche response,

$$H(\omega) = H_M(\omega) + \frac{\alpha}{\omega^2}. \quad (1.6)$$

This choice corresponds to a direct relation between the horizontal particle acceleration and the vertical height,

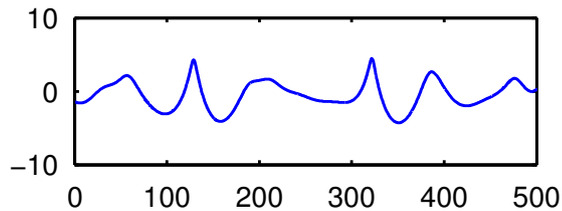
$$\frac{\partial^2 x(t, u)}{\partial t^2} = \frac{\partial^2 x_M(t, u)}{\partial t^2} - \alpha w(t, u) \quad (1.7)$$

where $x_M(t, u)$ is the Miche solution; [5, 6, 7].

1.2 Generating Lagrange waves with WAFOL

The basic routines in WAFOL for simulation of Lagrange waves are `spec21dat` and `1dat21wav`.

Figure 1.2: *Front-back symmetric Lagrange space wave on shallow water ($h = 8$ m); direct plot.*



1.2.1 The simulation options

The parameters for the simulations are set by an options structure. The default option `opt = simoptset` gives the result

`opt =`

```

    Nt: 2048
    Nu: 2048
    Nv: []
    dt: 0.5000
    du: 0.5000
    dv: []
    lalpha: 0
    lbeta: 0
    ffttype: 'ffttime'
    iseed: 'shuffle'
    plotflag: 0

```

with number of time and space points, the corresponding time and space steps, and the value of the α parameter in (1.6), (β is not used in this tutorial). The `ffttype` determines the hierarchy in the simulation – `ffttime` uses the FFT routine in time, stepping over the different space values. The alternatives are `fftspac` and `ffttwodim`. The `iseed` option `shuffle` sets the random number generator to, just, random.

1.2.2 Generating the elementary processes

The simulation routine `spec2ldat` is an expansion of the WAFO routines `spec2sd` and `seasim`. Figure 1.2 was generated by the following commands:

```

S = jonswap(1.5); S.h = 8;
opt = simoptset('Nt',256,'dt',0.125,'Nu',256*8,'du',0.25,'iseed',123791);
[w,x] = spec2ldat(S,opt,'iseed','shuffle') ; % Keep [w,x]
subplot(211)
plot(x.u+x.Z(:,128),w.Z(:,128));
axis([0 500 -10 10]) % Keep the figure

```

Note how `simoptset` changes the default values of `opt` and how `spec2ldat` resets seed option to `'shuffle'`. The argument 1.5 in `jonswap` is a cut off frequency to remove small ripples. The output `[w,x]` gives the vertical and horizontal fields as structures with values at the selected space and time coordinates as fields `'Z'`, `'u'`, `'t'`:

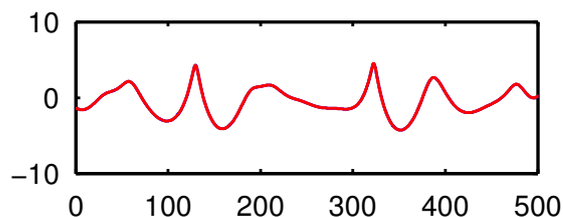


Figure 1.3: *Front-back symmetric Lagrange space wave on shallow water ($h = 8$ m) generated by `ldat2lwav`.*

```
w =
    Z: [2048x256 double]
    u: [2048x1 double]
    t: [1x256 double]
    note: 'JONSWAP, Hm0 = 7, Tp = 11, gamma = 2.3853'
```

```
x =
    Z: [2048x256 double]
    u: [2048x1 double]
    t: [1x256 double]
    note1: 'Horizontal Lagrange component'
    note2: 'alpha=0, beta =0'
```

and the plot command `plot(x.u+x.Z(:,128),w.Z(:,128))` plots the space wave according to the definition, observed at time $t_0 = 128 * dt$.

1.2.3 Generating the Lagrange wave

The WAFOL routine `ldat2lwav` is used to construct Lagrange time or space waves from the elementary processes. Keep the `[w,x]` from the previous session and compute the space wave at time $t_0 = 16$.³

```
[L,L0] = ldat2lwav(w,x,'space',16)
subplot(212)
plot(L.u,L.Z,L0.u,L0.Z,'r')
axis([0 500 -10 10])
```

The blue curve (almost invisible) in Figure 1.3 is the same space wave as is plotted “by hand” in Figure 1.2. The red curve is a smoothed version; the two are almost identical. The full call to the routine is

```
[L,Lsmooth]=ldat2lwav(w,x,type,tu0,dense)
```

where `type` can be `'time'` or `'space'` and `tu0` is the space or time (in absolute units `meter` or `seconds`), respectively, for which the wave is computed. The parameter `dense` is the interpolation rate (a positive integer). The output `Lsmooth` is an interpolated and smoothed version of `L`; see also Subsection 1.2.4 for exceptions where the Lagrange model give unphysical results.

³If you run into MATLAB error, make sure you use the `smooth` routine from WAFOL; check your MATLAB path.

The mean water level

The Lagrange transformation acts on the Gaussian field $w(t, u)$ by compressing the crest parts of the waves, making them shorter in space or time, and expanding the trough parts, making them longer. This affects the empirical mean surface level and makes it (in general) negative. We illustrate this and take the mean of the Gaussian part and of the generated space Lagrange wave, and obtain, for the example above,

```

MGauss = mean(w.Z(:))
MGauss = 1.2794e-16
MLagrange = mean(L0.Z)
MLagrange = -0.5025

```

The standard deviation

The distortion of the Gauss field also affects the standard deviation, making it smaller for the Lagrange wave than for the Gauss wave. The theoretical standard deviation for the Gaussian wave is obtained by the commands `mom=spec2mom(S)` and `std = sqrt{mom(1)}`. Our simulation yielded

```

SGauss = std(w.Z(:))
SGauss = 2.0529
SLagrange = std(L0.Z)
SLagrange = 1.9376

```

For crest-trough asymmetric waves with peaked crests and flat troughs is it a general rule that the ratio between standard deviation and average crest-trough wave height is smaller than for a corresponding Gauss field.

1.2.4 Depth dependence and loops

The Lagrange transformation is sensitive to the water depth. For infinite water depth the water particle will move in randomly perturbed circles. For decreasing depth the particle paths will be more and more elongated and randomly deformed ellipses, [10]. For very small depths the model may produce typical loops at the wave crests, where the water surface is folded. This is a consequence of the absence of physical constraints in the model.

The interpolation and smoothing in the WAFO routine `ldat2lwav` does not accept loops and will produce a smoothed version `Lsmooth` up to about a wave period/wave length before the first loop. If the first loop occurs early in the series, then `L0` will be empty. The routine `looptest(S,opt)` simulates independent samples of `[w,x]` and gives as output the observed number of `x`-fields in which loops has occurred.

The depth dependence for space waves is illustrated in Figure 1.4, generated by the following code – to get time waves, just change the type option from `'space'` to `'time'` and plot with `L0.t`, etc. Note that the shallow water case `S.h=3` will easily produce folding. In the plot we therefore use a direct plot routine for that case.

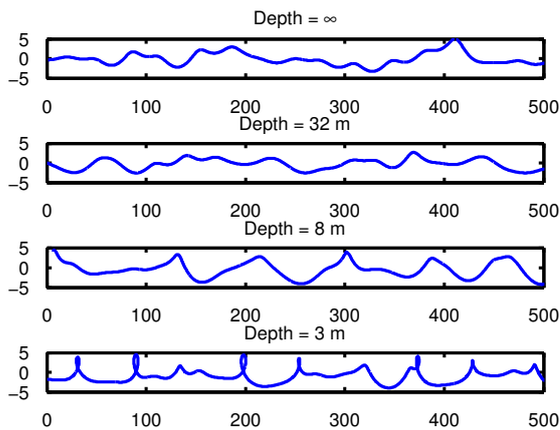


Figure 1.4: *Crest-trough asymmetric Lagrange space wave on different water depth.*

```

opt = simoptset('Nt',128,'Nu',2048,'du',0.25);
S = jonswap(1.5); S3 = S; S3.h = 3;
S8 = S; S8.h = 8; S32 = S; S32.h = 32;
[w,x] = spec2ldat(S,opt);
[w3,x3] = spec2ldat(S3,opt);
[w8,x8] = spec2ldat(S8,opt);
[w32,x32] = spec2ldat(S32,opt);
[L,L0] = ldat2lwav(w,x,'space');
[L3,L03] = ldat2lwav(w3,x3,'space');
[L8,L08] = ldat2lwav(w8,x8,'space');
[L32,L032] = ldat2lwav(w32,x32,'space');

figure(1); clf
subplot(411)
plot(L0.u,L0.Z); axis([0 500 -5 5]);
title('Depth = \infty')
subplot(412)
plot(L032.u,L032.Z); axis([0 500 -5 5]);
title('Depth = 32 m')
subplot(413)
plot(L08.u,L08.Z); axis([0 500 -5 5]);
title('Depth = 8 m')
subplot(414)
% 3m water depth may cause loops and ldat2lwav
% then gives only a short piece of L03.
% We plot space wave directly from w3,x3 as in Figure 1.2
plot(x3.u+x3.Z(:,64),w3.Z(:,64))
axis([0 500 -5 5])
title('Depth = 3 m')

```

Figure 1.5 shows the truncation effect of `ldat2lwav` with the following code.

```

S=jonswap(1.5); S.h=20;
opt=simoptset('dt',0.125,'lalpha',2);

```

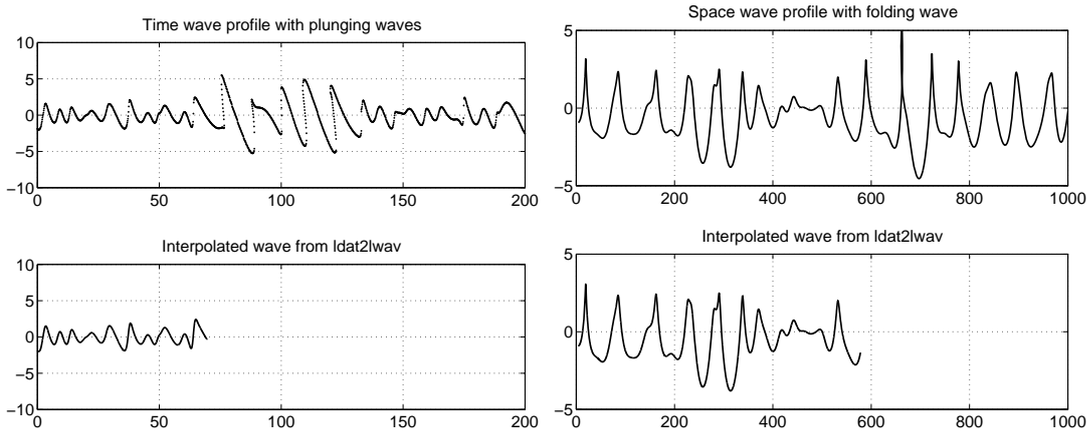


Figure 1.5: *Left: Time wave profile with plunging wave truncated by ldat2lwav. Right: Space wave profile with folded wave truncated by ldat2lwav.*

```
[w,x]=spec2ldat(S,opt);
[L,L0]=ldat2lwav(w,x,'time',[ ],10,1)
pause
S=jonswap(1.5); S.h=4;
opt=simoptset('dt',0.125,'lalpha',0,'ffttype','fftspace');
[w,x]=spec2ldat(S,opt);
[L,L0]=ldat2lwav(w,x,'space',[ ],10,1)
```

If no subplot(212) appears, L0 was empty – try again.

1.2.5 Front-back asymmetry

The main feature of the Lagrange approach is that it can easily generate waves with realistic front-back asymmetry.⁴ The asymmetry is controlled by the parameter `opt.lalpha`. This example illustrates, in Figure 1.6, the effect of different α -values on the front-back asymmetry,

```
opt = simoptset('Nt',2048,'dt',0.125,'Nu',512,'du',0.25);
S = jonswap(1.5); S.h=20;
[w0,x0] = spec2ldat(S,opt);
[w1,x1] = spec2ldat(S,opt,'lalpha',0.75);
[w2,x2] = spec2ldat(S,opt,'lalpha',1.5);
[L0,L00] = ldat2lwav(w0,x0,'time');
[L1,L01] = ldat2lwav(w1,x1,'time');
[L2,L02] = ldat2lwav(w2,x2,'time');
figure(1); clf
subplot(311); plot(L00.t,L00.Z);title('\alpha = 0');
axis([0 50 -10 10])
```

⁴The crest-trough asymmetry generated by the Lagrange transformation can also be found in the 2nd order Stokes waves; see Chapter 3.

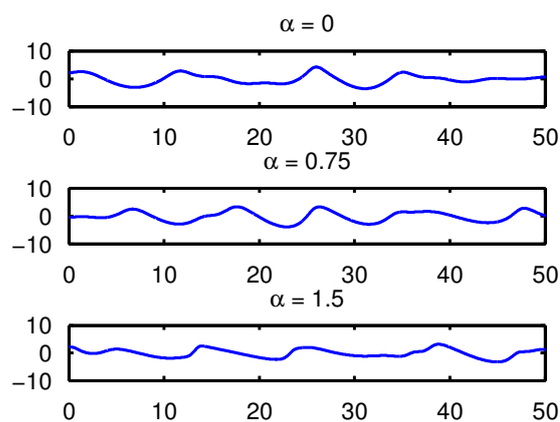


Figure 1.6: *Front-back asymmetric Lagrange time waves with different α -values. Water depth: $S.h = 20$ [m].*

```
subplot(312); plot(L01.t,L01.Z);title('\alpha = 0.75');
axis([0 50 -10 10])
subplot(313); plot(L02.t,L02.Z);title('\alpha = 1.5');
axis([0 50 -10 10])
```

Note that Figure 1.6 shows time waves where the steep wave side is facing left. You should generate the corresponding space waves – then you can chose a smaller Nt and a larger Nu -value.

1.3 Slopes and asymmetry measures

An argument for the Gauss-Lagrange model

To be really useful in practice any physical, mathematical or stochastic wave model has to be calibrated against empirical data. The stochastic Gauss-Lagrange wave model offers unique features, not shared by deterministic models, which can be used for calibration. Some of these features are also useful in their own right, and most important of these are the statistical distributions of different geometrical wave characteristics.

1.3.1 Extracting slope characteristics from wave data

WAFOL offers two ways to extract slope values from wave data, one general and one based on the special Lagrange format with separate vertical and horizontal components.

The general routine `wav2slope` can be applied to all types of wave data with the call

```
Slopes = wav2slope(L,absolutelevels,dense,p)
```

It takes a time or space wave `L` and extracts slope values at up- and downcrossings of the levels defined by the vector `absolutelevels`, after interpolation with rate `dense` and smoothing with parameter `p` with default value `p = 1`. The wave data `L` must be a structure with values in `L.Z` and arguments in `L.t` or `L.u`, for time and space waves, respectively. The arguments must be in increasing order, but need not be equidistant.

The second route is intended for simulated Lagrange wave data and it uses the basic structure of the model. The call is

```
Slopes = ldat2lslope(w,x,type,absolutelevels)
```

Here w , x are the output of the simulation routine `spec2ldat`. This routine identifies the x - and w -combinations that generate a time or space wave and computes the slope according to equations (9) and (10) in [6]. It disregards crossings that are combined with folding of plunging waves.

A special routine is available for simulation experiments with slope distributions directly from the orbital spectrum S . The call

```
Slopes = spec2slopedat(S,Nsim,type,relativelevels,opt)
```

simulates N_{sim} independent samples of Lagrange waves with specified spectrum and extracts the slopes, using `wav2slope`. The default choice of levels is `relativelevels = [-1 0 1 2]` which will give slopes at crossing of the absolute levels `[-1 0 1 2]*Hs/4` relative to the still water level `mw1 = 0`; note $H_s = 4*std$. The routine `spec2steepdat` is an extension of `spec2slopedat` and it will be described in Section 1.3.3.

Empirical slope distribution

To illustrate slope distributions in front-back asymmetric wave data we first generate synthetic waves and then use the first routine on the data. We use the same models as for Figure 1.6 and extract the slopes at the up- and downcrossings of levels `[0 1 2]*std` defined in units of standard deviations relative to the still water level. The call

```
opt = simoptset('Nt',2048*16,'dt',0.125,'Nu',512,'du',0.25);
S = jonswap(1.5,[6 10]); S.h=20;
[w0,x0] = spec2ldat(S,opt);
[w1,x1] = spec2ldat(S,opt,'lalpha',0.5);
[w2,x2] = spec2ldat(S,opt,'lalpha',1);
[L0,L00] = ldat2lwav(w0,x0,'time');
[L1,L01] = ldat2lwav(w1,x1,'time');
[L2,L02] = ldat2lwav(w2,x2,'time');
mom = spec2mom(S);
levels=[0 1 2]*sqrt(mom(1)); % wav2slope requires absolute levels
Slope0 = wav2slope(L00,levels);
Slope1 = wav2slope(L01,levels);
Slope2 = wav2slope(L02,levels)
```

will produce a result like (depending on the randomness)

Slope2 =

```
up: {[993x1 double] [560x1 double] [110x1 double]}
down: {[993x1 double] [560x1 double] [110x1 double]}
levels: [0 1.4800 2.9600]
```

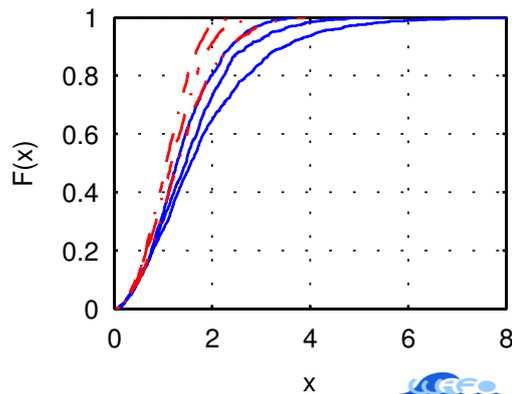


Figure 1.7: Empirical distributions of up- and downcrossing (absolute values) slopes at still water level for Lagrange time waves with different degrees of front-back asymmetry; $\alpha = 0, 0.5, 1$.

The plot the empirical distributions in Figure 1.7 of the slopes of the mean level we use the WAFO routine `plottedf`,

```
plottedf(Slope0.up{1}); hold on
plottedf(Slope1.up{1});
plottedf(Slope2.up{1});
plottedf(-Slope0.down{1}, 'r-.');
plottedf(-Slope1.down{1}, 'r-.');
plottedf(-Slope2.down{1}, 'r-.');
axis([0 8 0 1])
```

Simulated slope distributions can be generated directly from the spectrum by the routine `spec2slopedat`. As a by-product we also get the average shape of the waves in the neighborhood of upcrossings and downcrossings of the mean water level. We generate some such empirical distributions before we compute the exact theoretical ones. Figure 1.8 shows, in the upper row, simulated CDF of slopes at up- and downcrossings of different levels. The lower row shows the average wave shape near up- and downcrossing of the mean water level. (The downcrossing curves are mirrored in the origin to be comparable with upcrossings ditto.) First compute the distributions,

```
S=jonswap(1.5, [6 10]); S.h=20;
opt=simoptset(opt, 'Nt', 64, 'dt', 0.25, ...
    'Nu', 2048*16, 'du', 0.25, 'fftype', 'fft', 'fftsize', 1);
Nsim=100; % Takes time; Nsim = 10 in the command script
Slopes=spec2slopedat(S, Nsim, 'space', [], opt)
Slopes1=spec2slopedat(S, Nsim, 'space', [], opt, 'lalpha', 1)
```

and then plot the empirical CDF:s and the average waves; Fig 1.8.

```
figure(1); clf
subplot(221); box; hold on
for f=1:length(Slopes.levels),
    if ~isempty(Slopes.up{f})
        plottedf(Slopes.up{f})
    end
    if ~isempty(Slopes.down{f})
```

```

        plottedf(-Slopes.down{f},'-.')
    end
end
axis([0 0.8 0 1])
subplot(222); box; hold on
for f=1:length(Slopes1.levels),
    if ~isempty(Slopes1.up{f})
        plottedf(Slopes1.up{f})
    end
    if ~isempty(Slopes1.down{f})
        plottedf(-Slopes1.down{f},'-.')
    end
end
axis([0 0.8 0 1])
subplot(223); box; hold on
plot(Slopes.meanwavex,Slopes.meanwaveup); ax=axis;
axis([Slopes.meanwavex(1) Slopes.meanwavex(end) ax(3) ax(4)])
plot(Slopes.meanwavex,Slopes.meanwavedown,'-'); grid
subplot(224); box; hold on
plot(Slopes1.meanwavex,Slopes1.meanwaveup)
axis([Slopes.meanwavex(1) Slopes.meanwavex(end) ax(3) ax(4)])
plot(Slopes1.meanwavex,Slopes1.meanwavedown,'-'); grid

```

One can see from the upper left plot in Figure 1.8 that the slopes become steeper with increasing level, in agreement with the crest-trough asymmetry of the Lagrange waves.

Simulation with $l\alpha > 1$ may generate warnings for double crossings (= folding waves) and empty output. Such waves are discarded in the simulation which means that the generated slopes are slopes in waves where no folding occurred. Each repetition in the example corresponds to 8192 m of waves, and in a run with 400 repetitions with $\alpha = 1.5$ a total of 34 fields were discarded.⁵ To obtain sufficient accuracy, 50 repetitions are sufficient.

Run a similar example with time waves with a larger `opt.Nt` and smaller `opt.Nu` – see code in `WAFOL.m`. Notice how downcrossing slopes are smaller than the upcrossing ones for time waves, while the opposite holds for space waves.

1.3.2 Theoretical slope distribution

The `WAFOL` module contains routines for computation of the exact theoretical slope distribution at level crossings, based on the crossing theory in [5, 6].

The main routines to get the exact theoretical slope distribution CDF:s directly from the orbital spectrum are `spec2timeslopecdf` and `spec2spaceslopecdf` and the related

⁵The reason to disregard folded waves is that slope distribution becomes less meaningful for such waves.

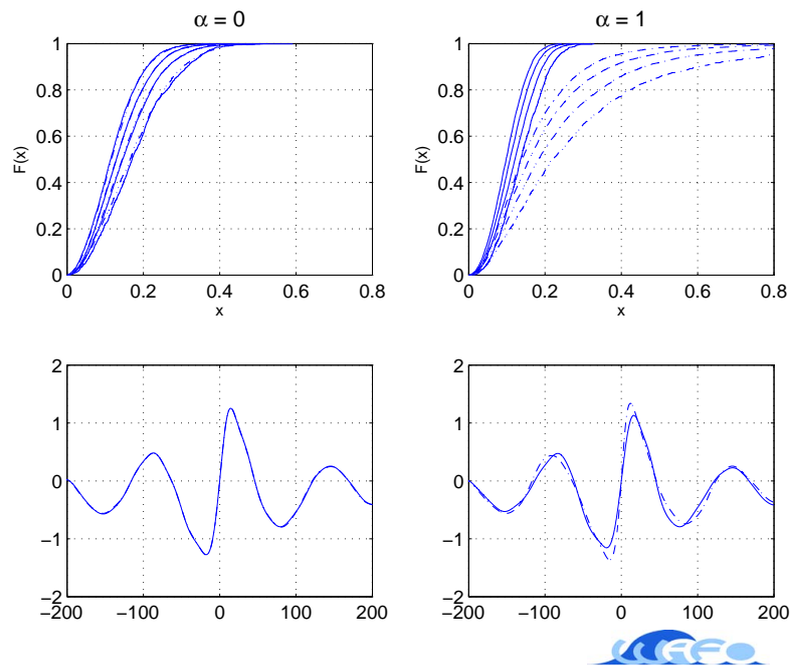


Figure 1.8: *Space waves. Upper row: Simulated CDF of slopes at up (solid) and down (dash-dotted) crossings of levels $[-1 \ 0 \ 1 \ 2] \times$ standard deviation. Left: front-back symmetric waves, $\alpha = 0$, where up- and downcrossing profiles are identical. Right: asymmetric waves, $\alpha = 1$. Lower row: Average wave shape near up- (solid) and down- (dash-dotted) crossings.*

routines for the PDF-functions. We can now compare the empirical slop distributions that we just obtained with the theoretical ones.

We use the observed (simulated) `Slopes1` values from the previous example and compare with the theoretical distribution. For time waves we use the following commands. First, simulate slopes:

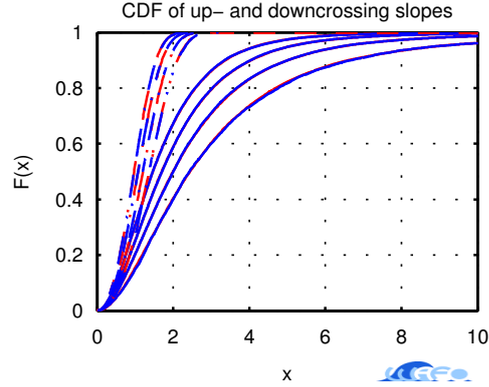
```
S=jonswap(1.5,[6 10]); S.h=20;
opt=simoptset('Nt',2048*16,'dt',0.125,...
    'Nu',128,'du',0.25,'fftype','fftime');
Nsim=100; % Takes time; {\tt Nsim = 10} in the command script
lev=[-1:2]; % Relative levels
% Absolute levels will be computed from spectral moment
Slopes=spec2slopedat(S,Nsim,'time',lev,opt)
Slopes1=spec2slopedat(S,Nsim,'time',lev,opt,'lalpha',1)
```

Then compute the theoretical CDF:s,

```
y=0:0.01:10;
[Fu,Fd] = spec2timeslopecdf(S,y,lev,opt);
[Fu1,Fd1] = spec2timeslopecdf(S,y,lev,opt,'lalpha',1);
```

and compare the results for $\alpha = 1$ in Figure 1.9 (for code, see `WafolCh1.m`).

Figure 1.9: CDF for up- and downcrossing (absolute values) slopes in asymmetric time waves with asymmetry parameter $\alpha = 1$: red = theoretical, blue = simulated, solid = upcrossing, dash-dotted = downcrossing. Crossed levels = $[-1 \ 0 \ 1 \ 2]$ *standard deviation. Outer curves = highest level.



1.3.3 Asymmetry measures

Wave asymmetry and wave steepness can be summarised in many different ways, and many measures have been suggested in the literature. To define the measures we need to define some geometric wave characteristics. With notations as in Figure 1.10, we define for *time waves*,

Slope ratio at down/up-crossings: $\lambda_{AL} = -\frac{E(L_t(t_{down}))}{E(L_t(t_{up}))} \approx \frac{E(H_{cb}/T_{cb})}{E(H_{cf}/T_{cf})}$,

Front/back period ratio: $\lambda_{NLS} = E(T_{cf})/E(T_{cb})$,

Hilbert transform: $A_H = E(\widehat{L}(t)^3)/\sigma^3$.

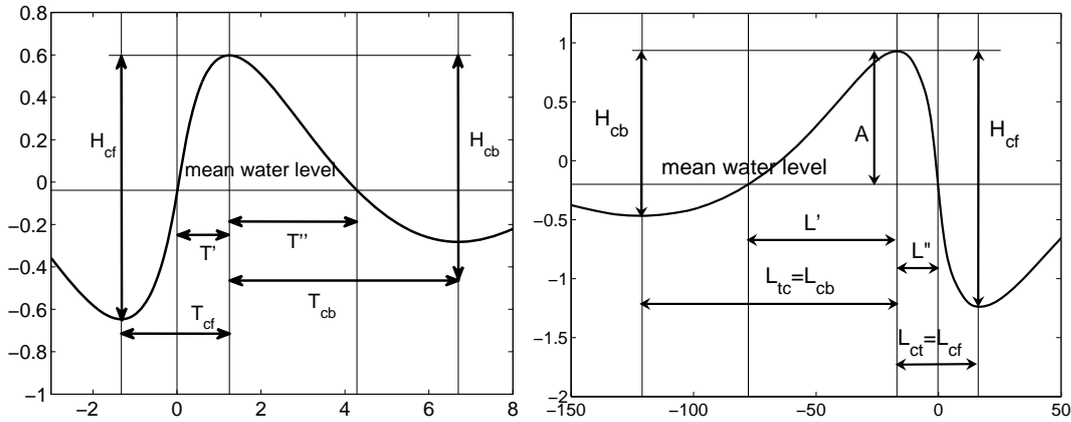


Figure 1.10: Asymmetry characteristics for time (left) and space (right) waves moving from left to right. The subscripts *cb, cf* refer to crest-back and crest-front definitions, and the subscripts *tc, ct* refer to trough-crest and crest-trough definitions.

The measure λ_{AL} was defined in [7] and is based on the slopes $L_t(t_{down})$ and $L_t(t_{up})$ at down- and upcrossings of the mean level. If front and back crest amplitudes are about the same, it is approximately equal to the index λ_{NLS} , proposed in [11]. The third measure A_H , introduced in [4], is based on the Hilbert transform $\widehat{L}(t)$ and is defined from its third moment and standard deviation.

The three asymmetry measures can be computed by simulation from the spectrum by the WAFOL routine `spec2steepdat` with the call

```
[Slope, Steep, Data] = spec2steepdat(S,Nsim,type,relativelevels,opt)
```

The routine is an extended (and more time-consuming) version of `spec2slopedat`, the routine that we used in Section 1.3.1. It generates, besides the slope characteristics at level crossings, crest front/back period data, and a data structure with simulated Lagrange data, together with first and second derivatives.

The structure `Slope` has fields with indices `lambdaAL` and `AH`, while the index `lambdaNLS` is a field in `Steep`. The following time consuming commands were used to generate the asymmetry data in Table 1.1. You may want to experiment with a smaller (or larger!) `Nsim`.

```
S = jonswap(1.5,[6 10]); S.h=20;
opt = simoptset('Nt',2048*16,'dt',0.125,...
    'Nu',321,'du',0.125,'fftype','fftime');
Nsim = 50; % Change this to Nsim = 10 or 20
[Slope0,Steep0,~] = spec2steepdat(S,Nsim,'time',[],opt);
[Slope05,Steep05,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',0.5);
[Slope10,Steep10,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',1);
[Slope15,Steep15,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',1.5);
[Slope20,Steep20,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',2.0);
```

The λ -indices in Table 1.1 are easy to interpret. For example, $\lambda_{AL} = 0.529$ for $\alpha = 1$ means that the time wave rate of increase is about twice as fast as the decrease at the still water level. The value $\lambda_{NLS} = 0.552$ for $\alpha = 2$ means that the time from trough to crest is about half the time from crest to trough, on average.

	λ_{AL}	λ_{NLS}	A_H
$\alpha = 0.0$	1.005	1.002	0.005
$\alpha = 0.5$	0.737	0.855	-0.191
$\alpha = 1.0$	0.529	0.734	-0.386
$\alpha = 1.5$	0.350	0.630	-0.588
$\alpha = 2.0$	0.226	0.552	-0.644

Table 1.1: *Simulated asymmetry measures in time waves for different α -values.*

CHAPTER 2

3D Lagrange waves

2.1 The 3D Lagrange model

The 3D Lagrange model consists of a Gaussian homogeneous random field $w(t, u, v)$ for the vertical movements of water particles and two correlated Gaussian fields $x(t, u, v), y(t, u, v)$ for the horizontal movements. The total energy of $w(t, u, v)$ is distributed over elementary waves with frequency ω and direction θ according to a *directional orbital spectrum* $S(\omega, \theta)$, $\omega > 0$, $-\pi < \theta \leq \pi$. The x -process describes the horizontal movements in the directions $\theta = 0, \pi$, and y the movements in the directions $\theta = \pm\pi/2$.

For simulation purposes in WAFOL the spectrum is discretized over wave-numbers $\kappa_{jk}^u, \kappa_{jk}^v$, and with related frequencies $\omega_{jk} > 0$, given by the dispersion relation,

$$\omega^2 = g|\kappa| \tanh h|\kappa|, \quad |\kappa| = \sqrt{\kappa_u^2 + \kappa_v^2}.$$

In analogy with (1.2) and (1.4) the three components in the 3D model are

$$w(t, u, v) = \sum_{j,k} \sqrt{S_{jk}} R_{jk} \cos(\kappa_{jk}^u u + \kappa_{jk}^v v - \omega_{jk} t + \Theta_{jk}), \quad (2.1)$$

$$x(t, u, v) = \sum_{j,k} \sqrt{S_{jk}} R_{jk} \rho_{jk}^u \cos(\kappa_{jk}^u u + \kappa_{jk}^v v - \omega_{jk} t + \Theta_{jk} + \psi_{jk}^u), \quad (2.2)$$

$$y(t, u, v) = \sum_{j,k} \sqrt{S_{jk}} R_{jk} \rho_{jk}^v \cos(\kappa_{jk}^u u + \kappa_{jk}^v v - \omega_{jk} t + \Theta_{jk} + \psi_{jk}^v), \quad (2.3)$$

The 3D Lagrange wave field

The 3D lagrange wave field is a random time varying field $L(t; x, y), t \in \mathbb{R}, (x, y) \in \mathbb{R}^2$ that, at time t , takes the value $w(t, u, v)$ at location $(x = x(t, u, v), y = y(t, u, v))$:

$$L(t; x(t, u, v), y(t, u, v)) = w(t, u, v). \quad (2.4)$$

It may happen that there are more than one (u, v) for which $(x(t, u, v), y(t, u, v)) = (x, y)$; in that case L takes multiple values.

The response functions

The response functions are allowed to depend on frequency ω and direction θ , and wave-number $\boldsymbol{\kappa} = (\kappa^u, \kappa^v)$ as in the 2D case. Now, there is more freedom to introduce asymmetry, depending on wave direction, and the user can construct a model for any special purpose. The standard implementation of WAFOL uses the following generalization of (1.6), introduced in [8, Eqn. (10)],

$$\mathbf{H}(\theta, \boldsymbol{\kappa}) = \begin{pmatrix} \rho^u e^{i\psi^u} \\ \rho^v e^{i\psi^v} \end{pmatrix} = \frac{\alpha}{\omega^2} \cdot \begin{pmatrix} \cos^2(\theta) |\cos(\theta)| \\ \cos^2(\theta) \sin(\theta) \operatorname{sign}(\cos \theta) \end{pmatrix} + i \frac{\cosh |\boldsymbol{\kappa}|h}{\sinh |\boldsymbol{\kappa}|h} \cdot \begin{pmatrix} \cos \theta \\ \sin \theta \end{pmatrix}. \quad (2.5)$$

This choice modifies the front-back asymmetry for waves with direction away from $\theta = 0, \pi$ and blocks it completely at $\theta = \pm\pi/2$.

2.2 Generating 3D Lagrange waves with WAFOL

2.2.1 Generating the elementary processes by spec2ldat3D

WAFOL offers two routines for simulation of the 3D elementary processes, and in this chapter we describe the 1st order routine, `spec2ldat3D`. The alternative routine is called `spec2ldat3DM` and it can also generate non-linear, 2nd order Stokes variation, both in vertical and horizontal dimension. That routine and its parallel companion `spec2ldat3DP` will be described in Chapter 3.

The directional spectrum

The most important input variable is of course the directional orbital power spectrum for the Gaussian wave field $w(t, u, v)$. It can be given it either frequency-direction form or in wave-number form, both supported by WAFO and WAFOL.

The simplest way to produce a directional spectrum is illustrated by the example in the WAFO-routine `mkdspec`,

```
S = jonswap
D = spreading(linspace(-pi,pi,51), 'cos2s')
Snew = mkdspec(S,D,1)
```

that will produce the left plot in Figure 2.1. The wave-number spectrum to the right is produced by

```
Sk2d = spec2spec(Snew, 'k2d')
plotspec(Sk2d)
axis([-0.05 0.05 -0.05 0.05])
```

The general form of the `spreading` command is

```
D = spreading(directions, type, maindirection, spreading, frequencies, fdep)
```

For experiments with front-back asymmetry it is recommended to set `maindirection = 0`. The parameter `spreading` defines the degree of directional spreading; `spreading = 0` gives isotropic waves with equal energy in all directions, a high value gives almost uni-directional 3D waves moving from “right to left” – this convention differs from that in Section 1.1.1. The default value is `spreading = 15` as in the example. Observe that `frequencies` must be the same as the frequencies in the spectrum, `S.w`; the standard spectra in WAFO have the same default values as the `spreading` function. The last parameter, `fdep` takes values 0,1 for frequency independent or frequency dependent spreading (default), respectively.

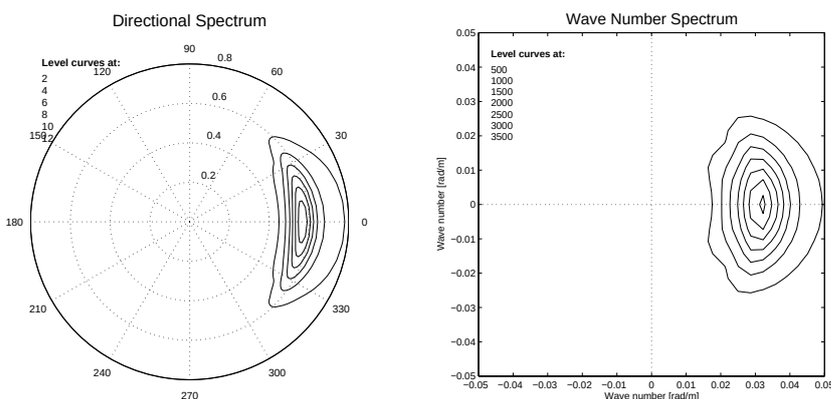


Figure 2.1: *Left: directional spectrum in frequency-direction form. Right: the same spectrum in wave-number form.*

The simulation options in `spec2ldat3D`

The options command `opt = simoptset` is used to define the number of time and space coordinates, `Nt`, `Nu`, `Nv`, their spacings, `dt`, `du`, `dv`, the asymmetry parameter `lalpha`, the seed for the random number generator `iseed`, (default `shuffle`), and the `plotflag`. The simulation in `spec2ldat3D` is always done with a 2D FFT over space, then looping over time, so the `ffttype` option is not relevant for the 3D model. The simulation generates a large data set with three fields of size `Nt x Nu x Nv` each, and there is a memory trade-off between time and space.

If one seeks a large wave field at only one time point, one can choose `Nt = 1` and `Nu`, `Nv` as large as needed. But if the goal is to get a set of long wave time series, i.e. `Nt` large, at a modest number of locations, one must be sure not to take `Nu x Nv` too small, and this for two reasons. First, both `Nu` and `Nv` must be big enough to cover the full variation of the horizontal movements within the studied region. This means, for example, that if one wants to generate wave a field over an area of size `200x100 meters` then one should have `Nu x du` at least `200 + 6 std(X)` and similar for the Y-field.¹

Second, the number of space points defines in a non-systematic way the periodicity of the time waves. At present, there is no systematic study of how large space field is

¹`std(X)` can be estimated by `mean(mean(std(X.Z)))`.

needed to avoid bias caused by time periodicity. The user is advised not to generate excessively long time series but rather generate many independent series.

In the examples we use the following options for time and space oriented problems:

```
optTime = simoptset('Nt',1001,'dt',0.2,...
                  'Nu',128,'du',5,'Nv',64,'dv',5);
optSpace = simoptset('Nt',1,'dt',0.5,...
                   'Nu',1024,'du',1,'Nv',512,'dv',1);
```

Generation of the elementary processes

The elementary processes are generated by the routine `spec2ldat3D`. First we define the directional spectrum and then generate the elementary processes. We define two spreading functions, one, `D5`, that gives moderate spreading, and one, `D15`, that gives waves with more concentrated direction.

```
S = jonswap(1.5); S.h = 20;
D5 = spreading(101,'cos',0,5,S.w,0);
SD5 = mkdspec(S,D5);
D15 = spreading(101,'cos',0,15,S.w,0);
SD15 = mkdspec(S,D15);
```

The time option

```
[W,X,Y] = spec2ldat3D(SD5,optTime);
```

takes about 30 seconds and generates about 200 MB of wave data. The space option

```
[W,X,Y] = spec2ldat3D(SD5,optSpace);
```

takes less than 1 second and generates about 25 MB of wave data.

2.2.2 Generating the Lagrange waves from the 3D fields

The generation options

The 3D analogue to `ldat2lwav` in `WAFOL` to generate wave fields from elementary wave data is `ldat2ldat3D` with call

```
L = ldat2lwav3D(W,X,Y,options3D)
```

where the argument `options3D` defines the type and dimension of the output. The routine can be used to generate single Lagrange fields or a series of fields that can be used to create a movie of the time dependent wave field. It is also possible to extract individual time series of point wave data from sets of locations.

The `options3D` structure can be set by an `genoptset` command as in this example:

```

opt3D=genoptset('type','field','t0',20)

opt3D =

    type: 'field'
      t0: 20
      PP: []
  start: [10 10]
      end: []
      rate: 1
  plotflag: 'on'

```

There are three different types: `field`, `moviedata`, `timeseries`. In the example the parameter `t0` sets the time of observation of a single field. The parameters `start`, `end` defines the absolute coordinates of the field or movie, that should be well inside area spanned by $N_u \times du$ and $N_v \times dv$; see text and footnote on page 19. The parameter `PP` is a $2 \times np$ array of coordinates for `np` observation points of time series, and `rate` is a parameter for interpolation in time series.

Generating a single field

We use the options

```

optSpace = simoptset('Nt',20,'dt',1,...
                    'Nu',256,'du',1,'Nv',128,'dv',1);
opt3D = genoptset('type','field','t0',10)

```

to generate 20 front-back asymmetric fields, estimate the standard deviations to decide on cut off points and extract the field at time `t0 = 10`:

```

[W,X,Y] = spec2ldat3D(SD15,optSpace,'lalpha',1)
Sx = mean(mean(std(X.Z))) % result = 4.6
Sy = mean(mean(std(Y.Z))) % result = 1.9
opt3D = genoptset(opt3D,'start',[20 10])
Lfield = ldat2lwav3D(W,X,Y,opt3D)

```

Figure 2.2 shows the result – note the vertical scale.

Generating a movie

The option `opt3D.type = 'moviedata'` (or simply `'movie'`) generates data for a 3D wave movie. The output from is a structure that can generate wave movie by means of the routine `seamovieL`; this is a version of the `WAFOL` routine `seamovie` adapted for the dimension standard in `WAFOL`.

We use the options

```

optTime = simoptset('Nt',1001,'dt',0.2,...
                    'Nu',128,'du',10,'Nv',64,'dv',10);
opt3D = genoptset('type','movie')

```

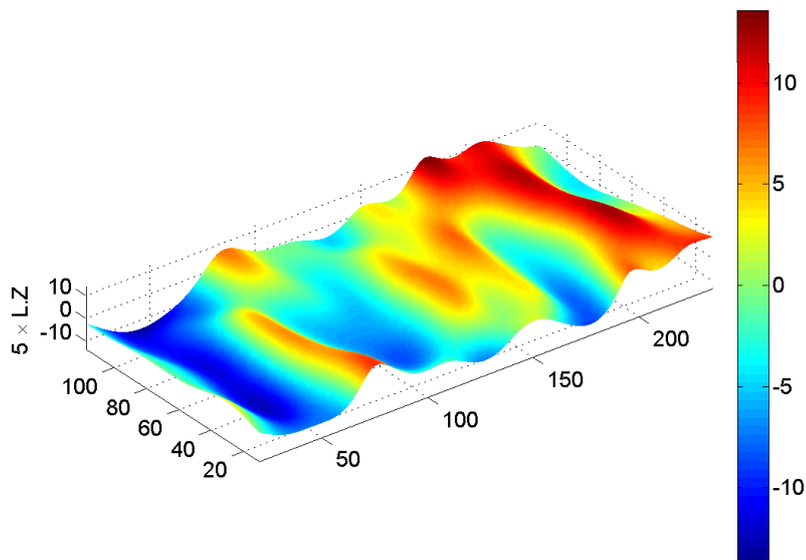


Figure 2.2: *A front-back asymmetric Lagrange field with strong directionality, generated by `ldat2lwav3D`.*

to generate wave movie data

```
[W,X,Y] = spec2ldat3D(SD15,optTime,'lalpha',1.5)
opt3D = genoptset(opt3D,'start',[20 10])
Lmovie = ldat2lwav3D(W,X,Y,opt3D)
```

that, besides producing the movie structure `Lmovie`, plots that last frame of the movie.

The movie is generated and displayed by a command

```
Mv = seamovieL(Lmovie,displaytype)
```

where `displaytype` can take values 1, 2, 3. A value 1 gives a surf-type perspective movie like field in Figure 2.2, a value 2 gives moving contours at still water level. `displaytype = 3` give a grayscale movie. The last frames with the two latter options are shown in Figure 2.3.

```
Mv2 = seamovieL(Lmovie,2); pause
Mv3 = seamovieL(Lmovie,3); pause
```

Beware that the movie structures are about 10 times bigger than the data structure `Lmovie` that is used to generate the movie.

Generating time series

The third type option in `genoptset` is `timeseries`, used to extract pointwise time series wave data obtained at one or more fixed locations. We illustrate how to obtain three time series, observed at three locations along the center line of the area:

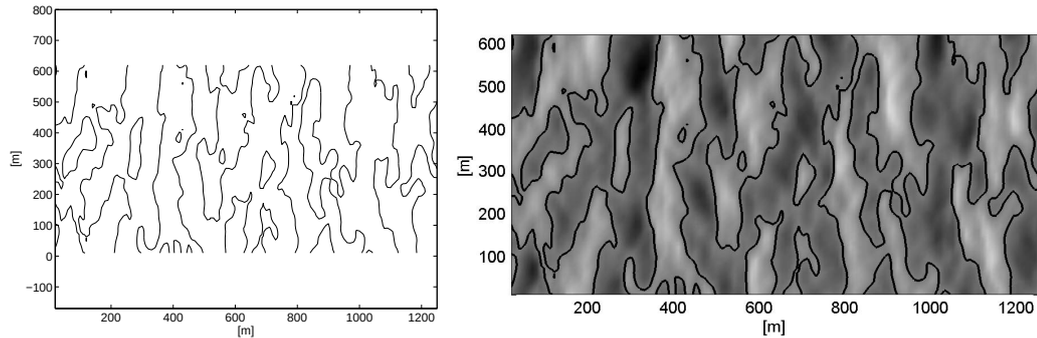


Figure 2.3: Frames of wave movie; option `displaytype = 2` (left) and `3` (right).

```
opt3D=genoptset('type','timeseries','PP',[400 425 450; 300 300 300])
```

```
opt3D =
```

```

type: 'timeseries'
t0: []
PP: [2x3 double]
start: [10 10]
end: []
rate: 1
plotflag: 'on'
```

We use the elementary wave data W, X, Y from the movie example and generate the three time series, and plot the result in Figure 2.4:

```

Lseries = ldat2lwav3D(W,X,Y,opt3D,'plotflag','off')
subplot(311)
plot(Lseries.t,Lseries.Z{1},'r')
grid on; hold on
plot(Lseries.t,Lseries.Z{2},'g')
plot(Lseries.t,Lseries.Z{3},'b')
```

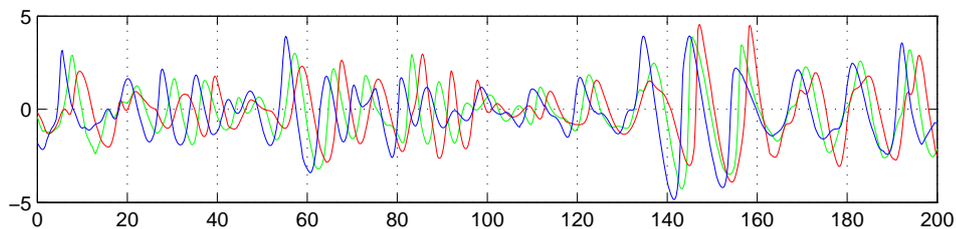


Figure 2.4: Three wave time series separated 25 meters in wind direction. Blue curve: upwind station, red curve: downwind station.

Figure 2.5 shows the cross-correlation function between the upwind and the downwind series, taken at a distance of 50 meters. We use the MATLAB routine `xcov` in the signal processing toolbox.

```
Rx = xcov(Lseries.Z{1},Lseries.Z{3},200,'biased')
subplot(211)
plot((-200:200)/5,Rx,'LineWidth',2,'FontSize',15)
```

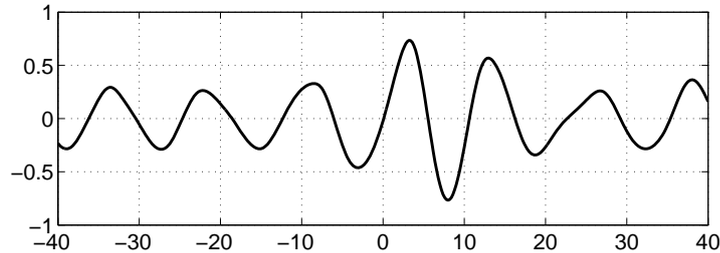


Figure 2.5: *Cross-correlation function between upwind and downwind time series; maximum correlation at time lag 3 seconds, in agreement with “average wave speed” of 17 seconds and peak period of 11 seconds.*

2.2.3 Generation of multiple time series from 3D spectrum

The routine `spec2lseries` generates several time series directly from the directional 3D spectrum without the route via `spec2l3d` and `l3d2lwav`. The call is

```
L = spec2lseries(Spec3D,PP,options)
```

where `PP = [x1 x2 ... xn; y1 y2 ... yn]` defines the coordinates for the observations. This direct routine will generate the same result and it takes about the same total time as the combined route, but it is much less memory demanding. In particular, the memory requirement is almost independent of the length and number of the time series.

If you want to compare the two methods, be sure to set `options.iseed` to a suitable integer – the default value is `shuffle`, i.e. `random` and set `opt3d.start = [0 0]`. Note that `opt3d.start` is not used in the direct routine; be sure that the observation points are well inside the generated region.

2.2.4 Run and export movies

You might want to export a wave movie for use outside MATLAB, for example, publish it on a web-page or send it by e-mail. Then you can use the `seamovieL` to save the movie as an avi-file. Just add a name string, e.g. `'Wave.avi'` as a third argument:

```
Mv1 = seamovieL(L,1,'Wave.avi');
```

If the current folder already contains a movie `Wave.avi` then a movie with a random name is produced. The extension `.avi` will be automatically added if missing.

The routine `seamovieL` uses the MATLAB-command `movie2avi` which puts restrictions on the MATLAB-installation. It works without extra work, at least, with version 8.5. The movie structures are usually large files and the avi-files generated are even larger. It is recommended that you edit them with a video editing program and save in some other format, e.g. `mp4`, to facilitate distribution.

CHAPTER 3

2^{nd} order non-linear Lagrange waves

3.1 Euler, Gauss, Lagrange, and Stokes waves

3.1.1 Gauss and 2^{nd} order Stokes interaction

The Gaussian waves model (1.2) describes the water surface as the sum of statistically independent harmonic components, acting without interaction between distinct frequencies. The 2^{nd} order Stokes model also contains sums of harmonics with frequencies equal to the sums and differences, in simplified form,

$$1^{st} \text{ order term : } w_1(t) = m + \sum A_j \cos(\omega_j t + \theta_j) = m + \Re \sum Z_j e^{i\omega_j t}, \quad (3.1)$$

$$2^{nd} \text{ order term : } w_2(t) = \Re \sum Z_j Z_k H_{jk}^+ e^{i(\omega_j + \omega_k)t} + \Re \sum Z_j Z_k^* H_{jk}^- e^{i(\omega_j - \omega_k)t}, \quad (3.2)$$

$$\text{Total wave : } w(t) = w_1(t) + w_2(t).$$

Here, the complex variables Z_j define both amplitudes and phases of the harmonics and * denotes complex conjugate. The factors H_{jk}^+ and H_{jk}^- are “ 2^{nd} order transfer factors”, and they are quite complicated depth dependent functions of the frequencies ω_j and the corresponding wave-numbers κ_j ; [9, 12, 13]. Equations (3.1-3.2) give an “Euler” description of the water elevation at a fixed location.

3.1.2 Lagrange and 2^{nd} order Stokes interaction

In the Gauss-Lagrange model the vertical and horizontal displacements are correlated Gaussian processes. In the Stokes-Lagrange model, also the horizontal processes have an added 2^{nd} order term of the same type as (3.2), but with special transfer factors, [3, 12, 13]. The horizontal components furthermore contain a random drift term, $d_x(t), d_y(t)$, constant in time, called the “Stokes drift”.

Figure 3.1 illustrates the linear filters, $\mathcal{L}_{zw}^1, \mathcal{L}_{zw}^1$, between the generating elements $\{Z_j\}$ and the Gaussian components, and the quadratic filters from w_1, x_1 to w_2, x_2 .

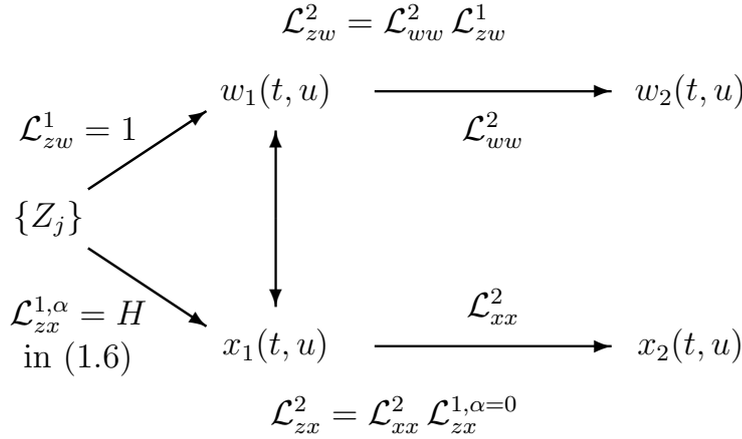


Figure 3.1: Schematic view of the generation of wave components from $\{Z_j\}$ in (3.1).

3.1.3 WAFO and WAFOL routines for 1st and 2nd order waves

The WAFO25 toolbox contains the routine `spec2nlsdat` for simulation of 2D Stokes-Euler waves. It is an extension of the (Euler) routine `spec2sdat` whose WAFOL version is `spec2l1dat`. The WAFO routine `seasim` has the WAFOL extension `spec2l1dat3D`; both generate 3D waves.

The WAFOL routines `spec2l1dat3DM` and `spec2l1dat3DP` are 3D Stokes-Lagrange companions to `spec2nlsdat`. They can be used to produce the components for 2D and 3D Stokes-Lagrange waves, including the Stokes drift. `spec2l1dat3DM` is a (M)odification and extension of the WAFOL routine `spec2l1dat3D` and `spec2l1dat3DP` is a version arranged for (P)arallel processing with the Parallel Computing Toolbox in MATLAB.

Table 3.1 lists the capabilities of different WAFO and WAFOL routines to simulate the components of linear and non-linear waves, based on the Gaussian generator $\{Z_j\}$.

	Gauss- Euler 2D	Gauss- Euler 3D	Gauss- Lagrange 2D	Gauss- Lagrange 3D	Stokes- Euler 2D	Stokes- Euler 3D	Stokes- Lagrange 2D	Stokes- Lagrange 3D
WafO								
<code>spec2sdat</code>	+	-	-	-	-	-	-	-
<code>seasim</code>	+	+	-	-	-	-	-	-
<code>spec2nlsdat</code>	+	-	-	-	+	-	-	-
WafOL								
<code>spec2l1dat</code>	(+)	-	+	-	-	-	-	-
<code>spec2l1dat3D</code>	(+)	(+)	+	+	-	-	-	-
<code>spec2l1dat3DM</code>	(+)	(+)	+	+	(+)	(+)	+	+
<code>spec2l1dat3DP</code>	(+)	(+)	+	+	(+)	(+)	+	+

Table 3.1: Spectral based simulation routines in WAFO and WAFOL

3.1.4 A comment on terminology

In the table we have used the terms “Euler” and “Lagrange” in an un-orthodox way. In hydrodynamics they denote two equivalent ways to define the physics of water waves, where an Euler description is focused on the velocity field at each fixed coordinate, while the Lagrange description integrates the velocity fields to get the trajectories of individual particles. The WAFOL routines are basically stochastic Euler routines, dealing only with the statistical properties of the vertical variation of the free surface, in the simplest form as a Gaussian process.

In the Lagrange wave model, as described in this tutorial, and in most publications on the topic, one simply borrows the Gaussian process from the Euler description and uses it as a Lagrange description of particle movements, with vertical and horizontal processes related by a hydrodynamically motivated linear filter equation. A quadratic filter then brings the Gauss model into a Stokes model. The WAFOL routines `spec2ldat`, `spec2ldat3D`, `spec2ldat3DM/P` generate such 1st and 2nd order Lagrange data `ldat`.

The routines `ldat2lwav` and `ldat2lwav3D` bring Lagrange data (of any order) into an easily observable Euler model. In fact, empirical Lagrange data on particle trajectories are rare, compared to the abundance of Euler data. Thus, our Lagrange routines not only simulate trajectories, but, with some extra effort, also the Euler observables. This explains the (+) notations in Table 3.1.

3.2 Generating 1st order processes

3.2.1 Introduction to the routines

The routine `spec2ldat3DM` generates 1st and 2nd order ingredients to a 3D Lagrange field, with a call similar to that of `spec2ldat3D` but with an extra parameter, `order`:

```
[W,X,Y] = spec2ldat3DM(Spec,order,options); % order = 1
[W,X,Y,W2,X2,Y2] = spec2ldat3DM(Spec,order,options); % order = 2
```

The input directional spectrum is defined by a one-sided spectrum structure with the direction dependence specified by a separate field.¹ Set up the spectrum information and simulation options as follows:

```
S = jonswap;
D = spreading(linspace(-pi,pi,91),'cos2s',0,10,S.w,0); % Here, the last
    % parameter is set to 0 to give frequency independent spreading
S.h = 40;
S.D = D; % This extra field carries the spreading information
Sdir = mkdspec(S,D);
plotspec(Sdir)
option = simoptset('Nt',1001,'dt',0.2,'Nu',300,'du',1,'Nv',100,'dv',1);
```

¹The WAFOL version 1.1.1 of `spec2ldat3DM` does not allow input of a directional spectrum with frequency dependent spreading. Thus, only frequency independent spreading is possible at present.

Note that the spectrum structure \mathbf{S} with the extra field $\mathbf{S.D} = \mathbf{D}$ carries the same information as the directional spectrum \mathbf{Sdir} . They are used as input in `spec2ldat3D` and in `spec2ldat3DM`, `spec2ldat3DP`, respectively.

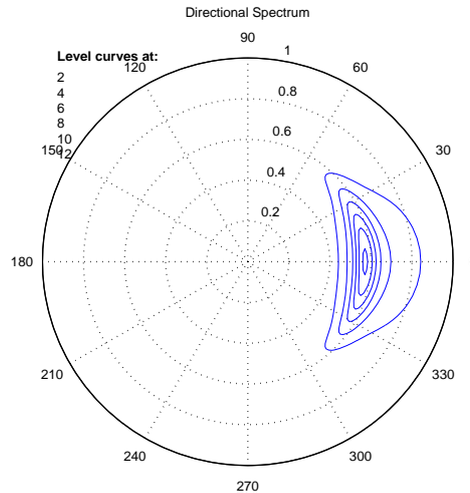


Figure 3.2: *Directional spectrum with frequency independent spreading; cf. Figure 2.1.*

3.2.2 A comparison of execution times for 1st order models

We start by comparing the execution times obtained with `spec2ldat3D`, `spec2ldat3DM`, and `spec2ldat3DP` with parallel computing activated. The first routine uses a two-dimensional FFT over space, looping over time, the two others use a one-dimensional FFT over time, looping over x- and y-space directions.

First generate the elements by the different routines and compare the distributions; they should be equal, but small differences must be expected.

```
[W1,X1,Y1] = spec2ldat3D(Sdir, option);
[W2,X2,Y2] = spec2ldat3DM(S,1, option);
% matlabpool local 4 % Run if PCT is available
[%W3,X3,Y3] = spec2ldat3DP(S,1, option);
```

```
figure(2), clf
subplot(221)
qqplot(W1.Z(:),W2.Z(:)); grid on
xlabel('W1.Z quantiles')
ylabel('W2.Z quantiles')
subplot(222)
plotnorm(W1.Z(:))
```

We have run each of the commands 100 times to get the execution time and total standard deviation for each of the components, `std(W.Z(:))`, etc., for each simulation, and then calculated the average and standard deviations.

We can draw two conclusions from the results in Table 3.2. As seen, the execution time is about the same for `spec2ldat3D`, with two-dimensional FFT, and `spec2ldat3DP`, with one-dimensional FFT and parallelization, while the non-parallelized version takes

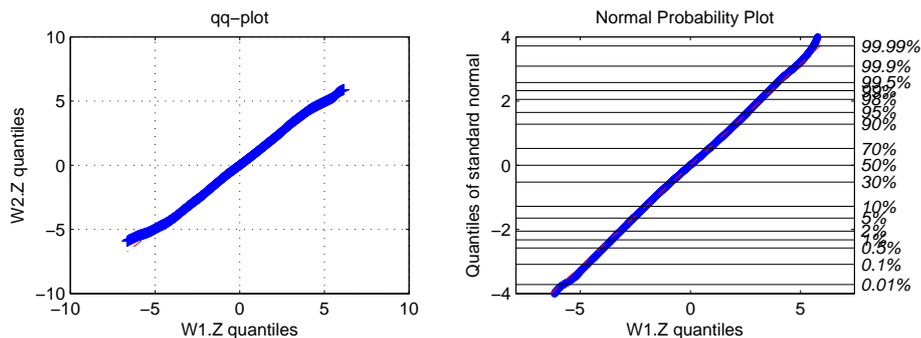


Figure 3.3: Right: normal plot of $W1.Z$ generated by `spec2ldat3D`. Left: a qqplot comparison with $W2.Z$ generated by `spec2ldat3DM`.

Routine	<code>spec2ldat3D</code>	<code>spec2ldat3DM</code>	<code>spec2ldat3DP</code>
Relative exec time	1 (2.3%)	1.32 (0.9 %)	1.05 (1.5%)
Mean std W.Z	1.75 (0.26)	1.72 (0.19)	1.70 (0.17)
Mean std X.Z	4.36 (0.89)	4.26 (0.64)	4.20 (0.57)
Mean std Y.Z	1.53 (0.32)	1.46 (0.19)	1.46 (0.18)

Table 3.2: Relative execution times and obtained average standard deviations for the three components, $W.Z$, $X.Z$, and $Y.Z$, simulated 100 times; numbers in parenthesis are the observed relative standard deviation, and standard deviations of the measured quantities, respectively. The absolute execution time is about 35 seconds per simulation with `spec2ldat3D`.

32% more time. One possible reason that `spec2ldat3D` can compete with `spec2ldat3DP` is that in the first routine, the space grid internally is optimized for `fft2`, stepping only in time, while in the second routine the parallelization is used in the u -variable, stepping over the v -variable, to perform a less time-consuming `fft` in the time-variable. As a consequence, the relative speed of the two routines will vary with the dimensions in time and space.

One can also note that the observed standard deviation of $W.Z$ is slightly less than $1.75 = Hs/4$ for two of the routines. This is an effect of the relative sparsity of the spectral density discretization, which is automatic from the time parameters, and leads to a small loss in energy. This phenomenon can also occur for `spec2ldat3D`.

Regardless of how the `ldat`-structures W, X, Y are generated, they can be used as in Chapter 2 by `ldat2lwav` and `ldat2lwav3D` to generate a Lagrange wave or wave field.

The motivation for the routines `spec2ldat3DM` and `spec2ldat3DP` is that they can generate 2nd order corrections both in the vertical component and in the horizontal components; then the parallelization is very effective.

3.2.3 Some differences between the two methods for 2D waves

The routines `spec2ldat3DM/P` can take a non-directional spectrum structure as spectral argument and generate the vertical, `W.Z`, and horizontal, `X.Z`, components for a 2D Lagrange wave. One should expect the result would be the same as that produced by `spec2ldat`, but there are some differences.

- `spec2ldat3DM/P` use the number of frequencies in the spectrum to decide on the number of time points to be generated. The spectrum is interpolated to reach the approximate time span as specified by `W.t(end) = (opt.Nt-1) * opt.dt`.
- `spec2ldat` change an odd value of `Nt` to the nearest greater even value; (similar with `Nu` if `fftttype = 'fftspac'`).

We issue the following commands in order to simulate `W`, `X` over 250 seconds with sampling rate 2 Hz, i.e. we set `Nu=501`, `dt=0.5`, and ask for the resulting `W`, `Wm`:

```
S = jonswap; opt = simoptset('Nt',501,'dt',0.5,'Nu',1001,'du',1);
[W,X] = spec2ldat(S,opt); [Wm,Xm] = spec2ldat3DM(S,1,opt);
```

`W =`

```
      Z: [1001x502 double]
      u: [1001x1 double]
      t: [1x502 double]
  note: [1x41 char]
    std: 1.6650
  meanperiod: 8.4879
  meanwavelength: 112.4406
```

`Wm =`

```
      Z: [1001x516 double]
      u: [1001x1 double]
      t: [516x1 double]
```

We note the change in the number of time steps, while the number of space steps is unchanged. The generated time spans are `W.t(end) = 250.5000` and `Wm.t(end) = 250.0145`. Thus, neither routine gives the exact time span, 250, that we expected!

3.2.4 Using the data from `spec2ldat3DM/P`

The `ldat` produced by `spec2ldat3DM/P` have the same structure as that produced by `spec2ldat3D` and it can be used as described in Chapter 2. However, since `spec2ldat3D` usually is the faster routine one may prefer that for 1st order simulation.

3.3 Generating 2^{nd} order 2D waves

We now turn to the main theme of this chapter, generation of 2^{nd} order Lagrange waves. The WAFO toolbox contains the routine `spec2nlsdat` for generation of 2^{nd} order 2D Stokes-Euler waves. With `spec2lmat3DM/P` one can generate both 2D and 3D Stokes-Lagrange waves, and we start with the 2D case.

3.3.1 2D Stokes time waves with `spec2nlsdat`

As listed in Table 3.1 there are two routines that generate 2D Stokes waves, the WAFO routine `spec2nllmat`, which directly generate a Stokes time wave at a single point, and the more general WAFO routines `spec2lmat3DM/P`, which give the Lagrange components, which combined with `lmat2lwav` give the time wave.

We use the JONSWAP spectrum without spreading, and set the depth to `S.h=20`.

```
S = jonswap; S.h = 20;  np = 250; dt = 0.2;
[xs2 xs1] = spec2nlsdat(S,np,dt);
figure(1); clf
waveplot(xs1,'r',xs2,'b',1,1)
```

The result is the Gaussian time wave `xs1`, a two-column array with time in the first and the vertical height in the second column, and the 2^{nd} order Stokes wave `xs2`. We use `waveplot` to show the result in Figure 3.4; see `help waveplot`.

3.3.2 2D Stokes waves with `spec2lmat3DM/P` and `lmat2lwav`

As shown in Chapter 2 the routine `lmat2lwav3D` can give all kinds of observable wave data from the Lagrange components `W,X,Y`. The option `opt3D.type = 'timeseries'` will lead to one or more timeseries, of the same type as the `xs1,xs2` waves produced by `spec2nlsdat`. In this section we want only 2D waves and use a uni-directional spectrum and set `opt.Nv = 1`. Note the result, that the Stokes drift in `X2` is negative directed in the main wave direction.

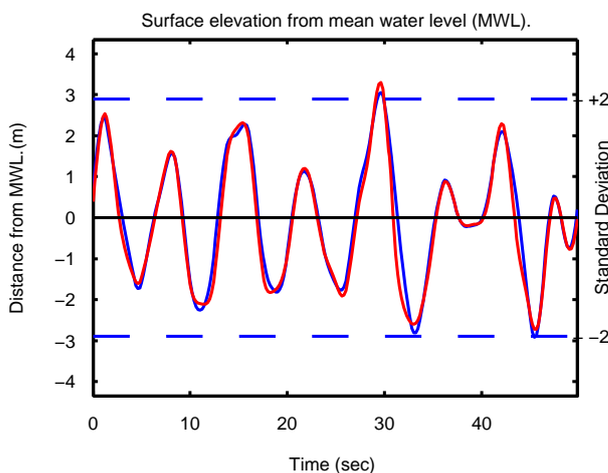


Figure 3.4: A waveplot of linear (blue) and non-linear (red) time wave generated by `spec2nlsdat`.

```

S = jonswap; S.h = 20;
opt = simoptset('Nt',250,'dt',0.2,'Nu',1001,'du',1,'Nv',1);
[W,X,~,W2,X2,~] = spec2ldat3DM(S,2,opt);
X2.drift(end,end)

```

Time waves

We now construct two sets of 2^{nd} order components, one with Stokes drift and one without, and generate the timeseries observed at location 250 and plot the three series:

```

Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Wtot2d = Wtot2; % No Stokes drift in the vertical direction !
Xtot2 = X; Xtot2.Z = X.Z + X2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;
[L,L0] = ldat2lwav(W,X,'time',250,1,0);
[L2,L20] = ldat2lwav(Wtot2,Xtot2,'time',250,1,0);
[L2d,L20d] = ldat2lwav(Wtot2d,Xtot2d,'time',250,1,0);

plot(L0.t,L0.Z,'b'); hold on, grid on;
plot(L20.t,L20.Z,'r-');
plot(L20d.t,L20d.Z,'r'); hold off

```

Figure 3.5 shows the effect of the Stokes drift, shifting the peaks in time. Observe that the shapes of shifted and non-shifted waves are different, since they have different reference coordinates, i.e. they have different origin in space.

Space waves

We use the component processes to generate space waves by means of `ldat2lwav`, but first we increase the time interval:

```

S = jonswap; S.h = 20;
opt = simoptset('Nt',1000,'dt',0.2,'Nu',1001,'du',1,'Nv',1);
[W,X,~,W2,X2,~] = spec2ldat3DM(S,2,opt);

```

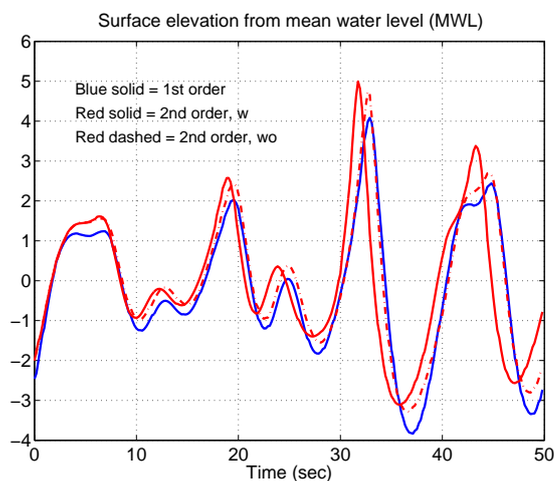


Figure 3.5: *Time waves: 1st order (blue) and 2nd order with (red, solid) and without (red, dashed) Stokes drift.*

We generate the space waves at three timepoints, 50, 100, 150 to see the effect of the Stokes in relation to the 2^{nd} order effect, and plot the results in Figure 3.6, (for plotting commands, see `WafolCh3.m`).

```

Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;

[L50,L050] = ldat2lwav(W,X,'space',50,1,0);
[L2d50,L20d50] = ldat2lwav(Wtot2d,Xtot2d,'space',50,1,0);
[L100,L0100] = ldat2lwav(W,X,'space',100,1,0);
[L2d100,L20d100] = ldat2lwav(Wtot2d,Xtot2d,'space',100,1,0);
[L150,L0150] = ldat2lwav(W,X,'space',150,1,0);
[L2d150,L20d150] = ldat2lwav(Wtot2d,Xtot2d,'space',150,1,0);

```

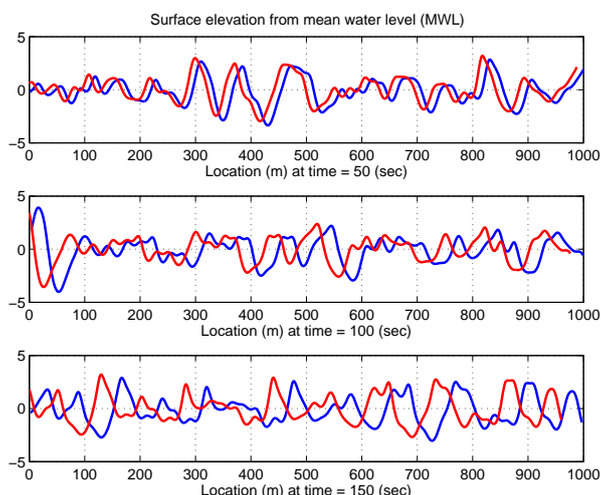


Figure 3.6: *Space waves: 1st order (blue) and 2nd order with Stokes drift (red) at three different time points. Waves move from right to left.*

3.3.3 Wave asymmetry in 2^{nd} order waves

The WAFO routine `spec2nlmdat` generates 2^{nd} order time waves with typical crest-trough asymmetry, and we saw one example in Section 3.3.1.

To generate front-back asymmetric waves we need the Lagrange technique in WAFOL, as was described in Sections 1.1.2 and 1.2.5. The degree of asymmetry is determined by the parameter α , which enters into the transfer function $\mathcal{L}_{zx}^{1,\alpha} = H$ in Figure 3.1. As illustrated in that figure, the α affects only the 1^{st} order and not the 2^{nd} order X -component; of course also the Y -component is affected in an analogous way.

We illustrate the effect by simulating the components,

```
[W,X,~,W2,X2,~] = spec2lmdat3DM(S,2,opt,'Nt',250,'lalpha',1.5);
```

and repeating the commands for the time waves in Figure 3.5 to get Figure 3.7.

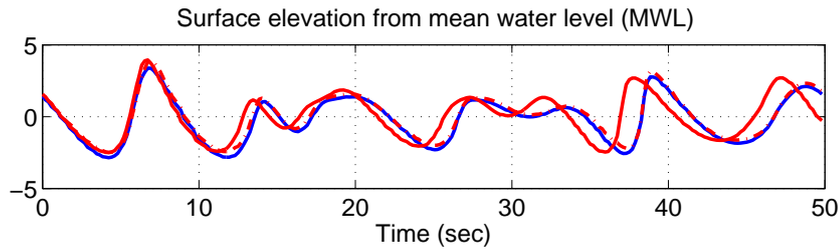


Figure 3.7: *Front-back asymmetric time wave with 2^{nd} order components.*

3.4 Generating 2^{nd} order 3D waves

To simulate 2^{nd} order 3D Lagrange waves with `spec2ldat3DM/P` we need a directional spectrum specified by a spectral density structure `S` with a direction field `.D`. We also set a preliminary options structure, make a simulation and take a look at the drift in the main wave direction:

```
S = jonswap; S.h=20;
D = spreading(linspace(-pi,pi,51),'cos2s',0,15,S.w,0);
S.D = D;
opt=simoptset('Nt',250,'dt',0.2,'Nu',250,'du',1,'Nv',50,'dv',1);
[W,X,W2,X2,Y2] = spec2ldat3DM(S,2,opt);
surf(X2.drift(1:2:end,1:2:end,end)) % To see the drift magnitude
```

We see in Figure 3.8 that in this simulation the maximal extension of the Stokes drift in the x -direction at the end of the time interval is almost 20 meters. This means that if we want to make a movie of the field over a certain region we have to extend the reference region with at least this amount to be sure to capture the variation over the whole region of interest. The drift in the y -direction is smaller and less than ± 5 meters.

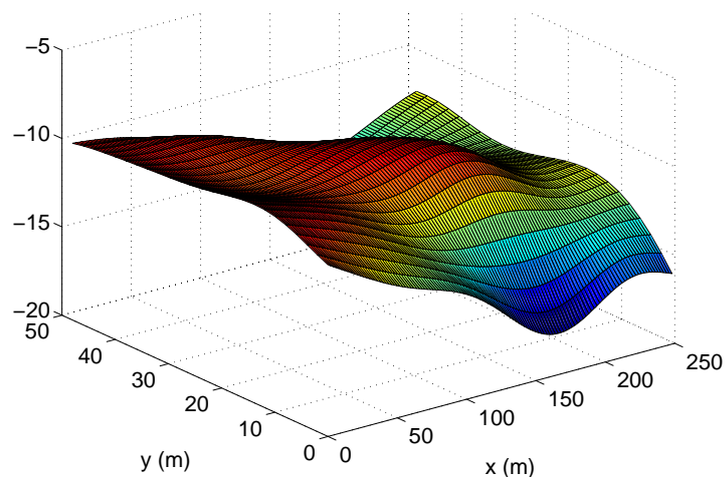


Figure 3.8: *Stokes drift at the end of the observation interval*

To be on the safe side, we decide to extend the reference region by 100 meters at the far end, by 10 meters at the near end, and by 15 meters both sideways. Thus, to simulate a 50 seconds wave movie over a region 250 x 100 meters we choose `opt.Nu=360`, `opt.du=1` and `opt.Nv=130`, `opt.dv=1`.

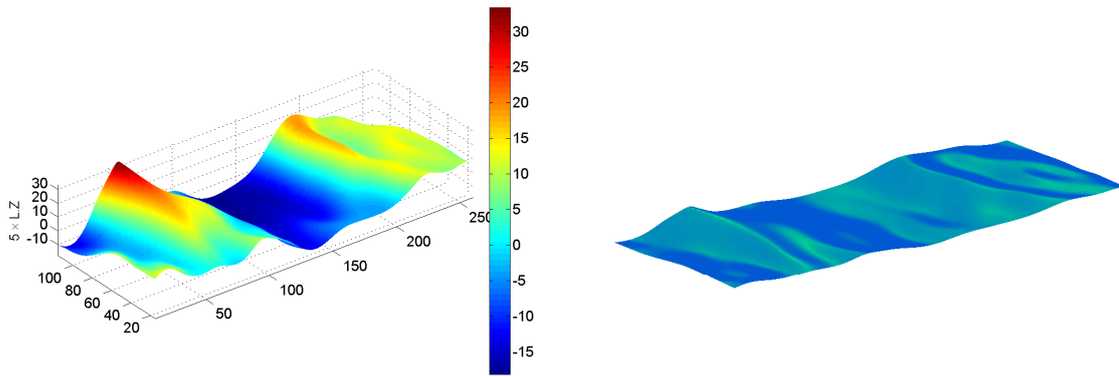


Figure 3.9: Last wave views from `ldat2lwav3D` and `seamovieL`; symmetric waves.

Note that if the reference region is defined too small strange patterns can appear at the borders of the simulated region. Then one has to extend the reference region to the expense of increased simulation time.

```
opt=simoptset('Nt',250,'dt',0.2,'Nu',360,'du',1,'Nv',130,'dv',1);
[W,X,Y,W2,X2,Y2] = spec2ldat3DM(S,2,opt);
Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;
Ytot2d = Y; Ytot2d.Z = Y.Z + Y2.Z + Y2.drift;
opt3D = genoptset('start',[10 15],'end',[260 115]);
figure(1); L=ldat2lwav3D(Wtot2,Xtot2d,Ytot2d,opt3D);
figure(2); Mv=seamovieL(L,1);
```

Figure 3.9 shows the last wave views in `L` and in `Mv` with different vertical scaling. The right hand view has the correct vertical and horizontal scaling while the left hand view exaggerates the vertical variation.

For comparison, we can generate front-back asymmetric waves by replacing `opt` by, e.g.,

```
optasym = simoptset(opt,'alpha',1.5);
```

and repeating the simulation. Figure 3.10 shows the result.

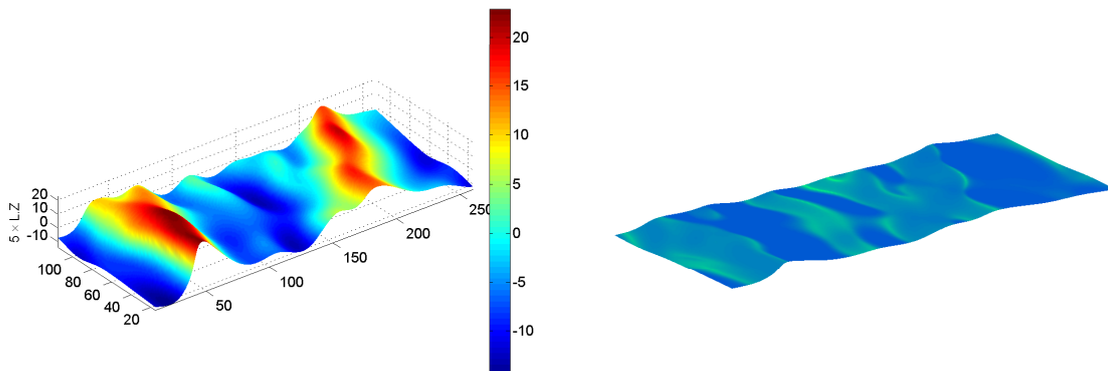


Figure 3.10: Last wave views from `ldat2lwav3D` and `seamovieL`; asymmetric waves.

APPENDIX A

Commands for the examples

The command files `WafolChx.m` contain the code for the examples in the tutorial. Some of the examples take some time and generate big data arrays. You may need to reduce the size of the problems, for example by using a smaller `Nsim` or reduce the dimension. Figure titles and other editing are often not included.

The time to run `WafolCh1` in Windows 7 with Matlab 2013a on an Intel Core i7-870, 2.93GHz, 16GB RAM, is about 30 minutes, the time for `WafolCh2` is less than one minute, and that for `WafolCh3` is about 15 minutes.

The “pause-state” is set to `off` in the script. Set it to `on` if you want to pause between tasks.

A.1 WafolCh1, Commands for Chapter 1

```
% Script with commands for Wafol tutorial, Chapter 1
```

```
pstate = pause('off');
TSTART = cputime;
disp(' Chapter 1: 2D Lagrange waves')

disp(' Section 1.1.2')
% Figure: Lagrange time wave, example of asymmetry
S = jonswap; S.h = 20;
opt = simoptset('dt',1);
[w,x] = spec2ldat(S,opt,'lalpha',1);
[L,L0] = ldat2lwav(w,x,'time',[],10);
subplot(211)
plot(L.t,L.Z); hold on
plot(L0.t,L0.Z,'r')
axis([0 40 -6 6]); hold off
pause
```

```

disp(' Section 1.2.2')
% Figure: Lagrange space wave, direct plot
S = jonswap(1.5); S.h = 8;
opt = simoptset('Nt',256,'dt',0.125,'Nu',...
    256*8,'du',0.25,'iseed',123791);
[w,x] = spec2ldat(S,opt,'iseed','shuffle')
subplot(211)
plot(x.u+x.Z(:,128),w.Z(:,128))
axis([0 500 -10 10])
pause

disp(' Section 1.2.3')
% Figure: Lagrange space wave from [L,L0]
[L,L0] = ldat2lwav(w,x,'space',16)
subplot(212)
plot(L.u,L.Z,L0.u,L0.Z,'r')
axis([0 500 -10 10])
pause

MGauss = mean(w.Z(:))
MLagrange = mean(L0.Z)
SGauss = std(w.Z(:))
SLagrange = std(L0.Z)
pause

disp(' Section 1.2.4')
% Figure: Crest-trough asymmetry in space, depth dependence
opt = simoptset('Nt',128,'Nu',2048,'du',0.25);
S = jonswap(1.5);
S3 = S; S3.h = 3;
S8 = S; S8.h = 8;
S32 = S; S32.h = 32;
[w,x] = spec2ldat(S,opt);
[w3,x3] = spec2ldat(S3,opt);
[w8,x8] = spec2ldat(S8,opt);
[w32,x32] = spec2ldat(S32,opt);
[L,L0] = ldat2lwav(w,x,'space');
[L3,L03] = ldat2lwav(w3,x3,'space');
[L8,L08] = ldat2lwav(w8,x8,'space');
[L32,L032] = ldat2lwav(w32,x32,'space');
figure(1)
clf
subplot(411)
plot(L0.u,L0.Z); axis([0 500 -5 5]);
title('Depth = \infty')

```

```

subplot(412)
plot(L032.u,L032.Z); axis([0 500 -5 5]);
title('Depth = 32 m')
subplot(413)
plot(L08.u,L08.Z); axis([0 500 -5 5]);
title('Depth = 8 m')
subplot(414)
% 3m water depth may cause loops and ldat2lwav give empty L3,L03
% We plot space wave directly from w3,x3
plot(x3.u+x3.Z(:,64),w3.Z(:,64))
axis([0 500 -5 5])
title('Depth = 3 m')
clear w x w3 x3 w8 x8 w32 x32
pause

% Figure: Truncation of time and space waves
S = jonswap(1.5); S.h=20;
opt = simoptset('dt',0.125,'lalpha',2);
[w,x] = spec2ldat(S,opt);
[L,L0] = ldat2lwav(w,x,'time',[ ],10,1)
pause
S = jonswap(1.5); S.h=4;
opt = simoptset('dt',0.125,'lalpha',0,'ffttype','fftspace');
[w,x] = spec2ldat(S,opt);
[L,L0] = ldat2lwav(w,x,'space',[ ],10,1)
clear w x
pause

disp(' Section 1.2.5')
% Figure: Front-back asymmetric time waves, different alpha
opt = simoptset('Nt',2048,'dt',0.125,'Nu',512,'du',0.25);
S = jonswap(1.5); S.h=20;
[w0,x0] = spec2ldat(S,opt);
[w1,x1] = spec2ldat(S,opt,'lalpha',0.75);
[w2,x2] = spec2ldat(S,opt,'lalpha',1.5);
[L,L0] = ldat2lwav(w0,x0,'time');
[L1,L01] = ldat2lwav(w1,x1,'time');
[L2,L02] = ldat2lwav(w2,x2,'time');
figure(1)
clf
subplot(311)
plot(L0.t,L0.Z)
title('\alpha = 0')
axis([0 50 -10 10])
subplot(312)
plot(L01.t,L01.Z)

```

```

title('\alpha = 0.75')
axis([0 50 -10 10])
subplot(313)
if ~isempty(L02),
    plot(L02.t,L02.Z)
    title('\alpha = 1.5')
    axis([0 50 -10 10])
end
pause

disp(' Section 1.3.1')
% Figure: Empirical time slope CDF at crossings - different alpha
opt = simoptset('Nt',2048*16,'dt',0.125,'Nu',256,'du',0.25);
S = jonswap(1.5,[6 10]); S.h=20;
[w0,x0] = spec2ldat(S,opt);
[w1,x1] = spec2ldat(S,opt,'lalpha',0.5);
[w2,x2] = spec2ldat(S,opt,'lalpha',1);
[L0,L00] = ldat2lwav(w0,x0,'time'); clear w0 x0
[L1,L01] = ldat2lwav(w1,x1,'time'); clear w1 x1
[L2,L02] = ldat2lwav(w2,x2,'time'); clear w2 x2
mom = spec2mom(S);
levels=[0 1 2]*sqrt(mom(1)); % wav2slope requires absolute levels
Slope0 = wav2slope(L00,levels); clear L00
Slope1 = wav2slope(L01,levels); clear L01
Slope2 = wav2slope(L02,levels)
clear L02
pause

% Plotting

figure(10)
clf
plottedf(Slope0.up{1}); hold on
%get(H101,'Children')
plottedf(Slope1.up{1});
plottedf(Slope2.up{1});
plottedf(-Slope0.down{1},'r-.');
plottedf(-Slope1.down{1},'r-.');
plottedf(-Slope2.down{1},'r-.');
axis([0 8 0 1])
pause

% Figure: Empirical space slope CDF at crossings - different alpha
% S=jonswap(1.5); S.h=20;
S=jonswap(1.5,[6 10]); S.h=20;
opt=simoptset('Nt',64,'dt',0.25,...

```

```

        'Nu',2048*16,'du',0.25,'ffttype','fftspace');
Nsim=10;
Slopes = spec2slopedat(S,Nsim,'space',[],opt)
Slopes1 = spec2slopedat(S,Nsim,'space',[],opt,'lalpha',1)
figure(1); clf
subplot(221); box; hold on
for f=1:length(Slopes.levels),
    if ~isempty(Slopes.up{f})
        plottedf(Slopes.up{f}); grid on
    end
    if ~isempty(Slopes.down{f})
        plottedf(-Slopes.down{f},'-'); grid on
    end
end
axis([0 0.8 0 1])
title('\alpha = 0')
subplot(222); box; hold on
for f=1:length(Slopes1.levels),
    if ~isempty(Slopes1.up{f})
        plottedf(Slopes1.up{f}); grid on
    end
    if ~isempty(Slopes1.down{f})
        plottedf(-Slopes1.down{f},'-'); grid on
    end
end
axis([0 0.8 0 1])
title('\alpha = 1')
subplot(223); box; hold on
plot(Slopes.meanwavex,Slopes.meanwaveup)
ax=axis;
axis([Slopes.meanwavex(1) Slopes.meanwavex(end) ax(3) ax(4)]);
plot(Slopes.meanwavex,Slopes.meanwavedown,'-'); grid on
subplot(224); box; hold on
plot(Slopes1.meanwavex,Slopes1.meanwaveup)
axis([Slopes1.meanwavex(1) Slopes1.meanwavex(end) ax(3) ax(4)]);
plot(Slopes1.meanwavex,Slopes1.meanwavedown,'-'); grid on
pause

% Figure: Same as previous, but for time waves
S=jonswap(1.5,[6 10]); S.h=20;
opt=simoptset('Nt',2048*8,'dt',0.125,...
    'Nu',321,'du',0.125,'ffttype','fftttime');
Nsim=10;
Slopes=spec2slopedat(S,Nsim,'time',[],opt)
Slopes1=spec2slopedat(S,Nsim,'time',[],opt,'lalpha',1)
figure(1); clf

```

```

subplot(221); box; hold on
for f=1:length(Slopes.levels),
    if ~isempty(Slopes.up{f})
        plottedf(Slopes.up{f}); grid on
    end
    if ~isempty(Slopes.down{f})
        plottedf(-Slopes.down{f},'-'); grid on
    end
end
axis([0 10 0 1])
title('\alpha = 0')
subplot(222); box; hold on
for f=1:length(Slopes1.levels),
    if ~isempty(Slopes1.up{f})
        plottedf(Slopes1.up{f}); grid on
    end
    if ~isempty(Slopes1.down{f})
        plottedf(-Slopes1.down{f},'-'); grid on
    end
end
axis([0 10 0 1])
title('\alpha = 1')
subplot(223); box; hold on
plot(Slopes.meanwavex,Slopes.meanwaveup);
ax=axis;
axis([Slopes.meanwavex(1) Slopes.meanwavex(end) ax(3) ax(4)]);
plot(Slopes.meanwavex,Slopes.meanwavedown,'-'); grid
subplot(224); box; hold on
plot(Slopes1.meanwavex,Slopes1.meanwaveup);
axis([Slopes.meanwavex(1) Slopes.meanwavex(end) ax(3) ax(4)]);
plot(Slopes1.meanwavex,Slopes1.meanwavedown,'-'); grid on
pause

disp(' Section 1.3.2')
% Figure: Comparison between empirical and theoretical slope CDF
S = jonswap(1.5,[6 10]); S.h=20;
opt = simoptset('Nt',2048*8,'dt',0.125,...
    'Nu',321,'du',0.125,'ffttype','ffttime');
Nsim = 10;
Slopes = spec2slopedat(S,Nsim,'time',[],opt);
Slopes1 = spec2slopedat(S,Nsim,'time',[],opt,'lalpha',1);
relativelevels = [-1:2];
y=0:0.01:10;
[Fu,Fd] = spec2timeslopecdf(S,y,relativelevels,opt);
[Fu1,Fd1] = spec2timeslopecdf(S,y,relativelevels,opt,'lalpha',1);
pause

```

```

% Plot the results
figure(1); clf
plottedf(Slopes1.up{4}); % Some very large values set the axis
clf
axis([0 10 0 1])
hold on; grid on
plot(Fu1.x,Fu1.f{4}, 'r')
plot(Fu1.x,Fu1.f{3}, 'r')
plot(Fu1.x,Fu1.f{2}, 'r')
plot(Fu1.x,Fu1.f{1}, 'r')
plottedf(Slopes1.up{4});
plottedf(Slopes1.up{3});
plottedf(Slopes1.up{2});
plottedf(Slopes1.up{1});
plot(Fd1.x,Fd1.f{1}, 'r-.')
plot(Fd1.x,Fd1.f{2}, 'r-.')
plot(Fd1.x,Fd1.f{3}, 'r-.')
plot(Fd1.x,Fd1.f{4}, 'r-.')
plottedf(-Slopes1.down{1}, '-. ');
plottedf(-Slopes1.down{2}, '-. ');
plottedf(-Slopes1.down{3}, '-. ');
plottedf(-Slopes1.down{4}, '-. ');
pause

disp(' Section 1.3.3')
% Table: Asymmetry measures
S = jonswap(1.5,[6 10]); S.h=20;
opt = simoptset('Nt',2048*8,'dt',0.125,...
    'Nu',321,'du',0.125,'fftype','fftime');
Nsim = 10;
[Slope0,Steep0,~] = spec2steepdat(S,Nsim,'time',[],opt);
[Slope05,Steep05,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',0.5);
[Slope10,Steep10,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',1);
[Slope15,Steep15,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',1.5);
[Slope20,Steep20,~] = spec2steepdat(S,Nsim,'time',[],opt,'lalpha',2.0);

disp('End of Chapter 1')

TSTOP=cputime
disp(['Total time = ' num2str(TSTOP-TSTART) ' sec']);
pause(pstate)

```

A.2 WafoLCh2, Commands for Chapter 2

```

% Script with commands for WafoL tutorial, Chapter 2

rng('default') % Reset the random number generator
pstate = pause('off');
TSTART = cputime;
disp(' Chapter 2: 3D Lagrange waves')

disp(' Section 2.2.1 Spectrum and simulation options')
% Directional spectrum
% Figure: Directional spectra in polar and Cartesian form
S = jonswap
D = spreading(linspace(-pi,pi,51),'cos2s',0,15,S.w,0)
figure(1); clf
Snew = mkdspec(S,D,1)

Sk2d = spec2spec(Snew,'k2d')
figure(2); clf
plotspec(Sk2d); clear Sk2d
axis([-0.05 0.05 -0.05 0.05])
pause

% Simulation options
optTime = simoptset('Nt',1001,'dt',0.2,...
    'Nu',128,'du',5,'Nv',64,'dv',5);
optSpace = simoptset('Nt',1,'dt',0.5,...
    'Nu',1024,'du',1,'Nv',512,'dv',1);

% Generation of the elementary processes
S = jonswap(1.5); S.h = 20;
D5 = spreading(101,'cos',0,5,S.w,0);
SD5 = mkdspec(S,D5);
D15 = spreading(101,'cos',0,15,S.w,0);
SD15 = mkdspec(S,D15);

% [W,X,Y] = spec2ldat3D(SD5,optTime);
[W,X,Y] = spec2ldat3D(SD5,optSpace);
pause

disp(' Section 2.2.2 Generating the Lagrange waves from the 3D fields')

% Generating a single field
optSpace = simoptset('Nt',20,'dt',1,...
    'Nu',256,'du',1,'Nv',128,'dv',1);
opt3D = genoptset('type','field','t0',10)

```

```

% Figure: One front-back asymmetric field
[W,X,Y] = spec2ldat3D(SD15,optSpace,'lalpha',1)
Sx = mean(mean(std(X.Z))) % result = 4.6
Sy = mean(mean(std(Y.Z))) % result = 1.9
opt3D = genoptset(opt3D,'start',[20 10])
Lfield = ldat2lwav3D(W,X,Y,opt3D)
pause

% Generating a movie
optTime = simoptset('Nt',101,'dt',0.2,...
                   'Nu',128,'du',10,'Nv',64,'dv',10);
opt3D = genoptset('type','movie')

[W,X,Y] = spec2ldat3D(SD15,optTime,'lalpha',1.5)
opt3D = genoptset(opt3D,'start',[20 10])
Lmovie = ldat2lwav3D(W,X,Y,opt3D)
pause

% Figure: Last frame of seamovie - two displays
Mv2 = seamovieL(Lmovie,2)
pause
Mv3 = seamovieL(Lmovie,3)
pause

% Generating time series
opt3D=genoptset('type','timeseries','PP',[400 425 450; 300 300 300])

% Figure: Three point time series at distance
Lseries = ldat2lwav3D(W,X,Y,opt3D,'plotflag','off')
subplot(311)
plot(Lseries.t,Lseries.Z{1},'r')
grid on; hold on
plot(Lseries.t,Lseries.Z{2},'g')
plot(Lseries.t,Lseries.Z{3},'b')
pause

% Figure: Cross-correlation function between time series
Rx = xcov(Lseries.Z{1},Lseries.Z{3},200,'biased');
subplot(211)
plot((-200:200)/5,Rx)

disp('End of Chapter 2')

TSTOP=cputime
disp(['Total time = ' num2str(TSTOP-TSTART) ' sec']);

```

```
pause(pstate)
```

A.3 WafolCh3, Commands for Chapter 3

```
% Script with commands for Wafol tutorial, Chapter 3

rng('default')
pstate = pause('off');
TSTART = cputime;
disp(' Chapter 3: 2nd order non-linear Lagrange waves')

disp(' Section 3.2.1')
% Figure: Directional spectrum
S = jonswap;
D = spreading(linspace(-pi,pi,91),'cos2s',0,10,S.w,0);
S.h = 40;
S.D = D;
Sdir = mkdspec(S,D);
figure(1), clf
plotspec(Sdir)
option = simoptset('Nt',1001,'dt',0.2,'Nu',300,'du',1,'Nv',100,'dv',1);
pause

disp(' Section 3.2.2')
% Figure: qq-plot
[W1,X1,Y1] = spec2ldat3D(Sdir,option);
[W2,X2,Y2] = spec2ldat3DM(S,1,option);

% Run if PCT is available
% matlabpool local 4
% [W3,X3,Y3] = spec2ldat3DP(S,1,option);

figure(2), clf
subplot(221)
plotqq(W1.Z(:),W2.Z(:)); grid on
xlabel('W1.Z quantiles')
ylabel('W2.Z quantiles')
subplot(222)
plotnorm(W1.Z(:))
pause
clear W1 W2 X1 X2 Y1 Y2

disp(' Section 3.2.3')
% Check changes in region
S = jonswap;
opt = simoptset('Nt',501,'dt',0.5,'Nu',501,'du',1);
```

```

[W,X] = spec2ldat(S,opt);
[Wm,Xm] = spec2ldat3DM(S,1,opt);
pause

disp(' Section 3.3.1')
% Figure: Linear and non-lines with spec2nlsdat
S = jonswap; S.h = 20;
np = 250; dt = 0.2;
[xs2, xs1] = spec2nlsdat(S,np,dt);
figure(1), clf
waveplot(xs1,'b',xs2,'r',1,1)
pause

disp(' Section 3.3.2')
% Time waves
S = jonswap; S.h = 20;
opt = simoptset('Nt',250,'dt',0.2,'Nu',1001,'du',1,'Nv',1);
[W,X,~,W2,X2,~] = spec2ldat3DM(S,2,opt);
X2 % The 2nd order x-components has a field 'drift'
pause

% Figure: Time waves with and without Stokes drift
Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Wtot2d = Wtot2; % No Stokes drift in the vertical direction !
Xtot2 = X; Xtot2.Z = X.Z + X2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;

[L,L0] = ldat2lwav(W,X,'time',250,1,0);
[L2,L20] = ldat2lwav(Wtot2,Xtot2,'time',250,1,0);
[L2d,L20d] = ldat2lwav(Wtot2d,Xtot2d,'time',250,1,0);

figure(1), clf
plot(L0.t,L0.Z,'b','LineWidth',2); hold on, grid on;
plot(L20.t,L20.Z,'r-','LineWidth',2)
plot(L20d.t,L20d.Z,'r','LineWidth',2)
set(gca,'FontSize',14)
title('Surface elevation from mean water level (MWL)')
xlabel('Time (sec)'); hold off
pause

% Space waves
S = jonswap; S.h = 20;
opt = simoptset('Nt',1000,'dt',0.2,'Nu',1001,'du',1,'Nv',1);
[W,X,~,W2,X2,~] = spec2ldat3DM(S,2,opt);

% Figure: Space wave with Stokes drift

```

```

Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Wtot2d = Wtot2; % No Stokes drift in the vertical direction !
Xtot2 = X; Xtot2.Z = X.Z + X2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;

```

```

[L50,L050] = ldat2lwav(W,X,'space',50,1,0);
%[L250,L2050] = ldat2lwav(Wtot2,Xtot2,'space',50,1,0);
[L2d50,L20d50] = ldat2lwav(Wtot2d,Xtot2d,'space',50,1,0);
[L100,L0100] = ldat2lwav(W,X,'space',100,1,0);
%[L2100,L20100] = ldat2lwav(Wtot2,Xtot2,'space',100,1,0);
[L2d100,L20d100] = ldat2lwav(Wtot2d,Xtot2d,'space',100,1,0);
[L150,L0150] = ldat2lwav(W,X,'space',150,1,0);
%[L2150,L20150] = ldat2lwav(Wtot2,Xtot2,'space',150,1,0);
[L2d150,L20d150] = ldat2lwav(Wtot2d,Xtot2d,'space',150,1,0);

```

```

figure(1), clf
subplot(311)
plot(L050.u,L050.Z,'b','LineWidth',2); hold on, grid on;
plot(L20d50.u,L20d50.Z,'r','LineWidth',2)
set(gca,'FontSize',12)
title('Surface elevation from mean water level (MWL)')
xlabel('Location (m) at time = 50 (sec)')
subplot(312)
plot(L0100.u,L0100.Z,'b','LineWidth',2); hold on, grid on;
plot(L20d100.u,L20d100.Z,'r','LineWidth',2)
set(gca,'FontSize',12)
xlabel('Location (m) at time = 100 (sec)')
subplot(313)
plot(L0150.u,L0150.Z,'b','LineWidth',2); hold on, grid on;
plot(L20d150.u,L20d150.Z,'r','LineWidth',2)
set(gca,'FontSize',12)
xlabel('Location (m) at time = 150 (sec)')
pause

```

```

disp(' Section 3.3.3, Front-back asymmetry')
% Figure: Front-back asymmetric 2nd order waves
S = jonswap; S.h = 20;
[W,X,~,W2,X2,~] = spec2ldat3DM(S,2,opt,'Nt',250,'lalpha',1.5);
Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Wtot2d = Wtot2;
Xtot2 = X; Xtot2.Z = X.Z + X2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;

[L,L0] = ldat2lwav(W,X,'time',250,1,0);
[L2,L20] = ldat2lwav(Wtot2,Xtot2,'time',250,1,0);
[L2d,L20d] = ldat2lwav(Wtot2d,Xtot2d,'time',250,1,0);

```

```

figure(1), clf
subplot(311)
plot(L0.t,L0.Z,'b','LineWidth',2); hold on, grid on;
plot(L20.t,L20.Z,'r-','LineWidth',2)
plot(L20d.t,L20d.Z,'r','LineWidth',2)
set(gca,'FontSize',14)
title('Surface elevation from mean water level (MWL)')
xlabel('Time (sec)')
pause

disp(' Section 3.4')
S = jonswap; S.h=20;
D = spreading(linspace(-pi,pi,51),'cos2s',0,15,S.w,0);
S.D = D;
opt=simoptset('Nt',250,'dt',0.2,'Nu',250,'du',1,'Nv',50,'dv',1);
[W,X,Y,W2,X2,Y2] = spec2ldat3DM(S,2,opt);

% Figure: Check the Stokes drift
figure(1), clf
surf(X2.u(1:2:end),X2.v(1:2:end),X2.drift(1:2:end,1:2:end,end))

% Extend the reference region
opt=simoptset('Nt',250,'dt',0.2,'Nu',360,'du',1,'Nv',130,'dv',1);
[W,X,Y,W2,X2,Y2] = spec2ldat3DM(S,2,opt);
Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;
Ytot2d = Y; Ytot2d.Z = Y.Z + Y2.Z + Y2.drift;
% and set the region of interest
opt3D = genoptyset('start',[10 15],'end',[260 115]);

% Figures: Fields and movies
figure(1), clf
L=ldat2lwav3D(Wtot2,Xtot2d,Ytot2d,opt3D)
drawnow
figure(2), clf
Mv=seamovieL(L,1)
pause

% Front-back asymmetry
optasym = simoptset(opt,'lalpha',1.5);
[W,X,Y,W2,X2,Y2] = spec2ldat3DM(S,2,optasym);
Wtot2 = W; Wtot2.Z = W.Z + W2.Z;
Xtot2d = X; Xtot2d.Z = X.Z + X2.Z + X2.drift;
Ytot2d = Y; Ytot2d.Z = Y.Z + Y2.Z + Y2.drift;
figure(3), clf

```

```
L2=lodat2lwav3D(Wtot2,Xtot2d,Ytot2d,opt3D)
drawnow
figure(4), clf
Mv2=seamovieL(L2,1)

disp('End of Chapter 3')

TSTOP=cputime
disp(['Total time = ' num2str(TSTOP-TSTART) ' sec']);
pause(pstate)
```

APPENDIX B

Wafol routines

Matlab m-routines

```
% WAFOL = lagrange module for WAFO toolbox
% Version 1.2.1 05-01-2016
%
% Wafol contains routines for 1st and 2nd order random Lagrange waves
%
%   dat2crossind      - Finds indices to level v down and/or upcrossings from data
%   disper2          - Dispersion relation with possible mean flow
%   genoptset         - Creates or alters 3D generation options structure
%   ldat2lslope       - Extracts slopes at level crossings in Lagrange model
%   ldat2lwav         - Finds time/space Lagrange process from simulated components
%   ldat2lwav3D       - Generates Lagrange 3D wave process from simulated components
%   looptest          - Simulates 2D Lagrange waves to estimate folding rate
%   lwav2frontback    - Gives front/back crest periods/wavelength of wave data
%   pdfnorm2d         - Bivariate Gaussian distribution
%   seamovieL         - Makes a movie of a 2D or 3D simulated sea structure
%   simoptset         - Creates or alters simulation options structure
%   spec2lasym        - Simulates asymmetry measures for Lagrange waves from spectrum
%   spec2lcov         - Calculates auto- and cross-covariance functions
%   spec2ldat         - Simulates w and x components of 2D Lagrange wave
%   spec2ldat3D       - Spectral simulation of components in 3D Lagrangian sea
%   spec2ldat3DM      - Particle trajectory simulation according to Marc Prevosto
%   spec2ldat3DP      - Parallel version of spec2ldat3DM for trajectory simulation
%   spec2lseries      - Spectral simulation of time series in 3D Lagrangian sea
%   spec2slcomp       - Compares 2nd order Stokes and 1st order Lagrange time waves
%   spec2slopedat     - Simulates Lagrange waves and extracts slopes at crossings
%   spec2slopedat3D   - Simulates values and slopes in 3D Lagrange field
%   spec2spaceslopecdf - Computes cdf for slope at crossings of space waves
%   spec2spaceslopepdf - Computes pdf for slope at crossings of space waves
%   spec2steepdat     - Simulates Lagrange waves and extracts steepness and slopes
%   spec2timeslopecdf - Computes cdf for slopes at crossings of time waves
%   spec2timeslopepdf - Computes pdf for slopes at crossings of time waves
%   wav2slope         - Extracts slopes at up- and downcrossings after smoothing
```

Scripts for examples

```
% WafolCh1      - Script with commands for Wafol tutorial, Chapter 1
% WafolCh2      - Script with commands for Wafol tutorial, Chapter 2
% WafolCh3      - Script with commands for Wafol tutorial, Chapter 3
```

Executables

```
% partkern.mexw32 - mexw32 file for use with 32 bit systems
% partkern.mexw64 - mexw64 file for use with 64 bit systems
```

```

function [ind, Nc]= dat2crossind(x,v,wdef,nowarning)
%DAT2CROSSIND Finds indices to level v down and/or upcrossings from data
%
%CALL: [ind, Nc]= dat2crossind(x,v,wdef/cdef,warning);
%
% ind = indices to the level v crossings of the original sequence x
% Nc = number of crossings (i.e.length of ind)
%
% x = the surface elevation data
% v = the reference level (default v = mean of x)
% wdef = defines the type of wave. Possible options are
%       'dw', 'uw', 'cw', 'tw' or 'none'. (Default 'none').
%       If wdef='none' all crossings will be returned,
%       otherwise only the crossings which defines a
%       wave according to the wave definition will be returned.
% cdef = defines the type crossings returned. Possible options are
%       'd' 'u' or 'all'. (Default 'all').
%       If def='d' all down-crossings will be returned.
%       Similarly if def='u' only the up-crossings will be returned
%       otherwise 'all' the crossings will be returned.
% nowarning = true suppresses warning for no crossings (default = false)
%
% Example:
% t = linspace(0,7*pi,250);
% x = sin(t);
% [ind, Nc] = dat2crossind(x,0.75,'u')
% plot(t,x,'.',t(ind),x(ind),'o')
%
% See also findcross, wavedef, crossdef

```

```

function [l,res] = disper2(t,dt,niter,dx_rel)
%DISPER2 Dispersion relation with possible mean flow
%
%   disper2: dispersion relation, Newton-Raphson
%           with possible mean flow
%
%            $(2.\pi.f-k.v)^2 = g.k.\tanh(k.d)$ 
%
%CALL: [l,res] = disper2(t[,d[,niter[,dx_rel]])
%
%   input:  t : period vector
%           dt : water depth vector [water depth d, mean flow v]
%              or matrix same number of lines as length of t
%              vector or matrix [water depth d, surface current ...
%                               speed, bottom current speed, type of profile]
%           type of profile = 0 => uniform (bottom current ...
%                               speed is not used) (default value)
%           type of profile = 1 => linear
%           type of profile = 2 => exponential
%           (default value [Inf,0])
%
%   output: l : wavelength (same size as t)
%           res : residue  $(\omega_0 - xk \cdot v)^2 - g_0 \cdot xk \cdot \tanh(xk \cdot d)$ 

```

```
function options3D = genoptset(varargin)
%GENOPTSET Creates or alters 3D generation options structure
%
%CALL: options3D = genoptset(funcname,opts1,opts2,...,par1,val1,par2,val2,...);
%
% options3D    = options structure in which the named
%               parameters have the specified values.
% funcname     = string giving the name of the function for which default
%               values for the options structure should be extracted.
%
% par1,par2..= strings identifying the parameter to alter
% val1,val2..= corresponding values the parameters are altered to.
%
% GENOPTSET sets options for transformation of ldat3D (W,X,Y) to
% 3D Lagrange fields or 2D timeseries
%
% GENOPTSET with no input arguments and no output arguments displays all
% parameter names and their possible values.
%
% See also troptset and simoptset
```

```
function Slopes=ldat2lslope(w,x,typ,levels)
%LDAT2LSLOPE Extracts slopes at level crossings in Lagrange model
%
%Call: Slopes = ldat2lslope(w,x,type,levels)
%
% Slopes = struct array with observed slopes at the up- and
%         downcrossings of specified levels
%
% w,x     = vertical and horizontal component in Lagrange model
% type    = 'space' gives slopes in space waves
%         'time' gives time slopes
% levels  = vector of absolute levels relative mwl=0
%         (no default)
%
% Example:
% S=jonswap; mom=spec2mom(S);
% opt=simoptset;
% opt=simoptset(opt,'dt',0.25,'du',0.25)
% [w,x]=spec2ldat(S,opt)
% levels=[0 1 2]*sqrt(mom(1));
% Slopes=ldat2lslope(w,x,'time',levels)
```

```

function [L,L0]=ldat2lwav(w_in,x_in,type,tu0,dense,plott)
%LDAT2LWAV Finds time/space Lagrange process from simulated components
% This version returns true time or 2D space profile and
% the smoothed initial part until the first loop/break
%
%CALL: [L,L0]=ldat2lwav(w,x,type,tu0,dense,plotting)
%
% L      = Lagrange process structure with fields
%         L.type and L.t/L.u and L.Z
%         L can contain loops and breaking waves
% L0     = a pruned and smoothed version of L without loops
%
% w      = Gaussian vertical process structure w.Z, w.u, w.t
% x      = Gaussian horizontal variation process structure
%         x.Z, x.u, x.t
% type   = 'time' or 'space'
% tu0    = time t0 for space wave,
%         = space coordinate u0 for time wave
% dense  = interpolation rate for smoothing
% plotting = 0/false, no plotting (default),
%         = 1/true, plot L (and L0 if not empty)

```

```

function L=ldat2lwav3D(W,X,Y,opt3D,varargin)
%LDAT2LWAV3D Generates Lagrange 3D wave process from simulated components
% from W,X,Y fields (as output from spec2ldat3D)
%
%CALL: L =ldat2lwav3D(W,X,Y,opt3D)
%
% L      = Lagrange structure with fields
%         L.Z, L.x, L.y, L.t, L.type
%
% W      = Gaussian vertical process structure w.Z, w.u, w.t
% X/Y    = Gaussian horizontal variation structures
%         X.Z, X.u, X.v, X.t
%         Y.Z, Y.u, Y.v, Y.t
%         if X and Y are empty then output is Gaussian
% opt3D  = structure (set by genoptset) with fields
%         .type   = 'movie' gives time dependent wave fields over times W.t
%                 = 'field' give wave field(s) at time(s) given by
%         .t0     = 't0' (string) or empty: gives t0 = W.t(end)/2 (default)
%                 = t0 (numeric): gives field at time t0
%         .start  = [startx starty] lower left corner of fields
%         .end    = [endx endy] upper right corner of fields
%                 if empty end is generated from start coordinates
%         .plotflag = 'on' plots one field - the last one
%
%         OR
%         .type   = 'timeseries' gives n time series at points
%                 with coordinates
%         .PP     = 2 x n array [p1,...,pn; q1,...,qn]'
%                 If n > 1 then output L.Z is is
%                 a cell array L.Z{1}, ..., L.Z{n}
%         .rate   = interpolation rate (default = 1) not yet implemented
%         .plotflag = 'on' plots one time series - the last one
%
%         OR (NOT YET AVAILABLE)
%         .type   = 'swath' gives encountered Lagrange sea elevation
%                 observed from a moving object with speed(s) --
%                 possible further fields for this option are
%         .v [m/s]= along straight line(s) from
%         .start  = default = (W.u(1),W.v(1)/2) to
%         .end    = default = (W.u(end),W.v(1)/2)
%         OR (NOT YET AVAILABLE)
%         .type   = 'vfield' gives velocity field

```

```
function Nloops = looptest(S,opt,varargin)
%LOOPTEST Simulates 2D waves to estimate folding rate
```

```
function Steep = lwav2frontback(L)
%LWAV2FRONTBACK Gives front/back crest periods/wavelength of wave data
%
%CALL: Steep =lwav2frontback(L)
%
% Steep = struct array with fields
%   .ffull = full front period/wavelength
%   .fhalf = half front period/wavelength
%   .bfull = full back period/wavelength
%   .bhalf = half back period/wavelength
%
% L      = 2D wave L.t (L.u) and L.Z
```

```
function pdf = pdfnorm2d(X,m,S)
%PDFNORM2D Bivariate Gaussian distribution
%
%CALL: pdf = pdfnorm2d(X,m,S)
%
% X = 1x2 or n x 2 array of arguments
% m = 1x2 or nn x 2 array of mean values (default zero vector)
% S = 2x2 covariance matrix (default identity matrix)
% If nn=1 then m is filled to size n x 2
% else if n ~= nn then X is filled with first row to size nn x 2
%
% Example:
% x = linspace(-5,5);
% [X1 X2] = meshgrid(x);
% f = reshape(pdfnorm2d([X1(:),X2(:)]),100,100);
% [area,epsi] = simpson(x,f);
% [area2,epsi2] = simpson(x,area);
%
% See also: pdfnorm, pdfnormnd
```

```
function Mv=seamovieL(Y,s,Wavename)
%SEAMOVIEL Makes a movie of a 2D or 3D simulated sea structure
%           and optionally saves it as an avi-file
%
%CALL: Mv = seamovieL(Y,s,Wavename)
%
%       Mv = movie
%
%       Y = struct with 2d or 3d simulation (from seasim)
%       s = type of plot if 3d: if s=1 then surf-plot, if s=2 contour,
%           else gray-scale overview with troughs dark and crests light
%           (default 1)
%       Wavename = optional namestring for avi-file (e.g. 'MyWave.avi')
%               If absent, no avi-file is produced
%               The avi-option works for Matlab ver 8.5 but not for ver 8.1
%
% See also seasim, movie, getframe
```

```

function options = simoptset(varargin)
%SIMOPTSET Creates or alters simulation options structure
%
%CALL: options = simoptset(funcname,opts1,opts2,...,par1,val1,par2,val2,...);
%
% options      = transformation options structure in which the named
%               parameters have the specified values.
% funcname     = string giving the name of the function for which default
%               values for the options structure should be extracted.
%               Options are 'dat2tr', 'lc2tr', 'reconstruct'.
% opts1,
% opts2..     = options structures
% par1,par2.. = strings identifying the parameter to alter
% val1,val2.. = corresponding values the parameters are altered to.
%
% SIMOPTSET combines the default options for a function given by FUNCNAME
% with new options structures (OPTS1,OPTS2,...) and/or with the named
% parameters (PAR1,PAR2,...) with the corresponding values (VAL1, VAL2,...).
% The parameters are set in the same order as the input arguments.
% Any parameters with non-empty values of the options struct overwrite
% the corresponding old parameters.
% The input arguments can be given in any order with one exception:
% PARx and VALx must be given in pairs in that order.
% Any unspecified parameters for PARx are set to [].
% Parameters with value [] indicate to use the default value for that
% parameter when OPTIONS is passed to the function. It is sufficient to
% type only the 2 first characters to uniquely identify the parameter
% or function name. Upper case letters for parameter names and values
% that are strings are ignored. If an invalid string is provided, the
% default is used.
%
% SIMOPTSET with no input arguments and no output arguments displays all
% parameter names and their possible values.
%
% SIMOPTSET with no input arguments creates an options structure
% OPTIONS where all the fields are set to []. ???
%
% See also troptset, genoptsetfunction options = simoptset(varargin)

```

```
function [Aa,Average]=spec2lasym(S,opt,alpha,Nsim)
%SPEC2LASYM Estimates asymmetry measures for Lagrange waves
%
% Useful to compute asymmetry measures for many degrees of asymmetry
% as specified by alpha = input vector of lalpha-values
% See help text for spec2slopedat
```

```

function [rww,rwx,rxx]=spec2lcov(spec,t,u,type,alpha,beta)
%SPEC2LCOV Calculates auto- and cross-covariance functions
%      between W(0,0) and X(t,u)
%      or selected derivatives in the 2D Lagrange model
%
%CALL: [rww,rwx,rxx]=spec2lcov(spec,t,u,type,alpha,beta)%
% For type==1
%      rww      = structure with fields  rww.R, rww.t, rww.u
%              with covariance values Cov(W(0,0),W(t,u))
%      rwx, rxx = similar structures with Cov(W(0,0),X(t,u)) and
%              Cov(X(0,0),X(t,u))
% For the other types the three covariance structures contain the
% covariance and cross-covariances indicated below
%      spec      = orbital spectrum structure with depth  h
%      t          = vector of time values
%      u          = vector of space values
%      type      = 1,2,3,4,5,6,7
%      alpha,beta = parameters in the linked model
%
% if type=1: W(0,0),X(0,0) and W(t,u),X(t,u)
%             gives r^(ww),r^(wx),r^(xx)
% if type=2: dW/dt(0,0),dX/dt(0,0) and W(t,u),X(t,u)
%             gives r^(ww)_t0,r^(wx)_t0,r^(xx)_t0
% if type=3: dW/du(0,0),dX/du(0,0) and W(t,u),X(t,u)
%             gives r^(ww)_u0,r^(wx)_u0,r^(xx)_u0
% if type=4: dW/dt(0,0),dX/dt(0,0) and dW/dt(t,u),dX/dt(t,u)
%             gives r^(ww)_tt,r^(wx)_tt,r^(xx)_tt
% if type=5: dW/du(0,0),dX/du(0,0) and dW/du(t,u),dX/du(t,u)
%             gives r^(ww)_uu,r^(wx)_uu,r^(xx)_uu
% if type=6: dW/dt(0,0),dX/du(0,0) and dW/dt(t,u),dX/du(t,u)
%             gives r^(ww)_tu,r^(wx)_tu,r^(xx)_tu
%
% NOTE: This routine works only for one-sided spectra

```

```

function [w,x]=spec2ldat(spec,options,varargin)
%SPEC2LDAT Simulates w and x components of 2D Lagrange wave
%
%CALL: [w,x] = spec2ldat(spec,options);
%
% w      = Gaussian vertical process structure w.Z,w.u,w.t
% x      = Gaussian horizontal process structure x.Z,x.u,x.t
%
% spec =S   a frequency spectral density structure in
%           angular frequency ('w') or frequency ('f') form
% options = struct with fields
%   .Nt     = giving Nt time points. (default length(S)-1=n-1).
%           If Nt>n-1 it is assumed that S.S(k)=0 for k>n-1
%   .Nu     = giving Nu space points (default = Nt)
%   .dt     = step in grid (default dt is defined by the Nyquist freq)
%   .du     = step in grid (default du is defined by the Nyquist freq)
%   (.u     = if non-empty and = [u1 u2 Nu] the u-vector will be set to
%           u = linspace(u1,u2,Nu), ONLY TESTED for ffttype='ffttime'
%           if empty, then u = linspace(0,(Nu-1)*du,Nu))
%   .lalpha = alpha value for modified Lagrange (default = 0)
%   .lbeta  = beta value for modified Lagrange (default =0)
%   .iseed  - method or starting seed for the random number generator
%           (default = 'shuffle')
%   .ffttype - 'fft-space', fft over space, loop over time
%           generate space series with evolvment
%           over time (useful if Nu > Nt),
%   - 'ffttime', fft over time, loop over space (default)
%           generate time series with evolvment
%           over space (useful if Nt > Nu),
%   - 'ffttwodim', 2D-fft over time and space.
%
% Example of spec2ldat and ldat2lwav
%
% S=jonswap; opt=simoptset;
% opt=simoptset(opt,'dt',0.25,'du',0.25);
% type='time';
% [w,x]=spec2ldat(S,opt)
% [L,Lsmooth]=ldat2lwav([w,x],type,[],10);
% subplot(211)
% plot(Lsmooth.t,Lsmooth.Z); axis([0 50 -6 6])
% [w,x]=spec2ldat(S,opt,'lalpha',1)
% [L1,Lsmooth1]=ldat2lwav([w,x],type,[],10);
% subplot(223)
% plot(Lsmooth1.t,Lsmooth1.Z); axis([0 50 -6 6])
% [w,x]=spec2ldat(S,opt,'lalpha',2)
% [L2,Lsmooth2]=ldat2lwav([w,x],type,[],10);
% subplot(224)
% plot(Lsmooth2.t,Lsmooth2.Z); axis([0 50 -6 6])
%
% Version corresponding to Applied Ocean Research 2009 with respect
% to .lalpha and .lbeta
% See also: spec2sdat,cov2sdat, gaus2dat

```

```

function [W,X,Y]=spec2ldat3D(Spec,options,varargin)
%SPEC2LDAT3D Spectral simulation of components in 3D Lagrangian sea
%
%CALL: [W,X,Y]=spec2ldat3D(Spec,options)
%
% W      = Gaussian vertical process structure W.Z,W.u,W.v,W.t
% X      = Gaussian horizontal process structure x.Z,x.u,x.v,x.t
% Y      = Gaussian horizontal process structure y.Z,y.u,y.v,y.t
%
% Spec   = a directional frequency spectral density structure in
%          angular frequency ('w') and directional ('theta') form
%          Alt. in wave number ('k', 'k2') form
% options = struct with fields
%   .Nt   = giving Nt time points. (default length(S)-1=n-1).
%          If Nt>n-1 it is assumed that S.S(k)=0 for all k>n-1
%   .Nu   = giving Nu space points along x-axis (default = Nt)
%   .Nv   = giving Nv space points along y-axis (default = Nt)
%   .dt   = step in grid (default dt is defined by the Nyquist freq)
%   .du   = step in grid (default du is defined by the Nyquist freq)
%   .dv   = step in grid (default dv is defined by the Nyquist freq)
%   .lalpha = alpha value for modified Lagrange (default = 0)
%   .iseed = starting seed number for the random number generator
%           (default 'shuffle')
%   .plotflag = 0 (no plotting)
%
% Version corresponding to Applied Ocean Research 2009 with respect
% to .lalpha
%
% Based on WAFO-routine seasim, version sept 2014, see documentation
% See also: spec2ldat,spec2sdat,cov2sdat, gaus2dat

```

```

function [W,X,Y,W2,X2,Y2] = spec2ldat3DM(Spec,order,options,varargin)
%SPEC2LDAT3DM Particle trajectory simulation according to Marc Prevosto
%           2D or 3D, first or second order Lagrange waves
%
%CALL: [W,X,Y,W2,X2,Y2] = spec2ldat3DM(spec,order,options)
%
% I - Spec      : one-sided spectral structure with fields
%     'S'       : spectral density values
%     'freq'    : 1D frequency spectrum over 'w'/'f'
%     'D'       : spreading structure from D = spreading
% I - order    : 1, first order (default), or 2, second order, waves
% I - options  : struct with fields
%     .Nt      = minimum number of time points
%     .Nu/.Nv  = giving Nu/Nv space points along x/y-axis (default = 100)
%     .dt      = approximate time-step
%     .du/.dv  = step in grid (default du=dv=1)
%     .iseed   = starting seed number for the random number generator
%               (default 'shuffle')
%     h        = water depth (field in Spec, default = Inf)
%
%     z0       = secret parameter = particle depth between 0 and -h
%               can be set inside program, default = 0
%     trf,tls  = secret parameters for to truncate spectrum to avoid
%               instability and increase speed,
%               can be set inside program, default = 0.01,0.001
%
% 0 - W,X,Y    = first order components
% 0 - W2,X2,Y2 = second order components

```

```

function [W,X,Y,W2,X2,Y2] = spec2ldat3DP(Spec,order,options,varargin)
%SPEC2LDAT3DP Parallel version of spec2ldat3DM trajectory simulation
%           2D or 3D, first or second order Lagrange waves.
%           Same as spec2ldat3DM but with parallel looping in space
%           using matlabpool - can reduce simulation time to
%           the expense of heavy memory usage
%
%CALL: [W,X,Y,W2,X2,Y2] = spec2ldat3DP(spec,order,options)
%
% I - Spec      : one-sided spectral density structure fields
%       'S'      : spectral density values
%       'freq'   : 1D frequency spectrum over 'w'/'f'
%       'D'      : spreading structure from D = spreading
% I - order     : 1, first order (default), or 2, second order, waves
% I - options   : struct with fields
%       .Nt      = minimum number of time points
%       .Nu/.Nv = giving Nu/Nv space points along x/y-axis (default = 100)
%       .dt      = approximate time-step
%       .du/.dv = step in grid (default du=dv=1)
%       .iseed   = starting seed number for the random number generator
%                   (default 'shuffle')
%       h        = water depth (field in Spec, default = Inf)
%
%       z0       = secret parameter = particle depth between 0 and -h
%                   can be set inside program, default = 0
%       trf,tls  = secret parameters for to truncate spectrum to avoid
%                   instability and increase speed,
%                   can be set inside program, default = 0.01,0.001
%
% 0 - W,X,Y     = first order components
% 0 - W2,X2,Y2  = second order components

```

```

function L=spec2lseries(Spec,PP,options,varargin)
%SPEC2LSERIES Spectral simulation of time series in 3D Lagrangian sea
%
%CALL: L=spec2lseries(Spec,Points,options)
%
% L      = struct with n time series
%
% Spec   = a directional frequency spectral density structure in
%          angular frequency ('w') and directional ('theta') form
%          Alt. in wave number ('k', 'k2') form
%
% Points = [x1, ..., xn; y1, ..., yn] array with coordinates of
%          measurement points
%
% options = struct with fields
%   .Nt    = giving Nt time points. (default length(S)-1=n-1).
%           If Nt>n-1 it is assumed that S.S(k)=0 for all k>n-1
%   .Nu    = giving Nu space points along x-axis (default = Nt)
%   .Nv    = giving Nv space points along y-axis (default = Nt)
%   .dt    = step in grid (default dt is defined by the Nyquist freq)
%   .du    = step in grid (default du is defined by the Nyquist freq)
%   .dv    = step in grid (default dv is defined by the Nyquist freq)
%   .lalpha = alpha value for modified Lagrange (default = 0)
%   .iseed = starting seed number for the random number generator
%           (default 'shuffle')
%   .plotflag = 0 (no plotting)
%
% Version corresponding to Applied Ocean Research 2009 with respect
% to .lalpha

```

```

function [s2,s1,L0]=spec2slcomp(S,options,varargin)
%SPEC2SLCOMP Compares 2nd order Stokes and 1st order Lagrange time waves
%
%CALL: [s2,s1,L0] = spec2slcomp(S,options,varargin)
%
% s2    = 2nd order Stokes wave structure s2.Z,s2.t
% s1    = Gaussian wave structure s1.Z,s1.t
% L0    = 1st order (smoothed) Lagrange wave structure L.Z,L.t
% spec  = S    a frequency spectral density structure in
%           angular frequency ('w') or frequency ('f') form
% options = struct with fields
%   .Nt = giving Nt time points. (default length(S)-1=n-1).
%         If Nt>n-1 it is assumed that S.S(k)=0 for k>n-1
%   .dt = step in grid (default dt is defined by the Nyquist freq)
%   .u  = [-u1 u1 Nu] gives u = linspace(-u1,u1,Nu) grid
%         Nu should be an odd integer
%         Generated waves are time waves observed at u = 0
%
% The routine is a combination of
%   spec2nldat (Stokes) and spec2ldat/ldat2lwav (Lagrange)
%
% See also spec2nldat spec2linspec, spec2ldat

```

```

function Slopes=spec2slopedat(S,Nsim,type,lev,options,varargin)
%SPEC2SLOPEDAT Simulates Lagrange waves and extracts slopes at crossings
%
%CALL: Slopes = spec2slopedat(S,Nsim,type,levels,options)
%
% Slopes = struct array with observed slopes at the up- and
%           down-crossings of specified levels
%
% S        = orbital spectrum
% Nsim     = number of replicates in simulation (default = 1)
% levels   = vector of standardized levels relative to zero
%           (default = [-1 0 1 2]*standard deviation)
% type     = 'space' (default), 'time', or 'approxtime'
% options  = struct with fields for individual replicates
%   .Nt    = giving Nt time points. (default length(S)-1=n-1).
%           If Nt>n-1 it is assumed that S.S(k)=0 for all k>n-1
%   .Nu    = giving Nu space points (default = Nt)
%   .du    = step in grid (default dt is defined by the Nyquist freq)
%   .dt    = step in grid (default dt is defined by the Nyquist freq)
%   .lalpha = alpha value for modified Lagrange
%   .lbeta  = beta value for modified Lagrange
%   .ffttype = 'ffttime' (default), 'fftspace', 'ffttwodim'
%   .iseed  - setting for random number generator,
%           default = 'shuffle', [ int32 ]
%   .plotflag - 'off', no plotting (default)
%           - 'on'
%
% Example:
% S=jonswap; opt=simoptset; mom=spec2mom(S);
% opt=simoptset(opt,'dt',0.25,'du',0.25)
% Nsim=100;
% levels=[0 1 2];
% Slopes=spec2slopedat(S,Nsim,'time',levels,opt)
%
% Used by spec2lasym
% See also: spec2ldat, ldat2lwav, wav2slope

```

```
function Slope = spec2slopedat3D(Spec,Nsim,Points,options,varargin)
%SPEC2SLOPEDAT3D Simulates values and slopes in 3D Lagrange field
%
%   with choice between
%   GL, fft2 over space, stepping over t, if Spec is dirspec
%   MP, fft in time, stepping over space, if Spec is onedim with field D
```

```

function [Fu,Fd]=spec2spaceslopecdf(S,y,lev,opt,varargin)
%SPEC2SPACESLOPECDF Computes cdf for slope at crossings of space waves
%
%CALL: [Fu,Fd] = spec2spaceslopecdf(S,y,type,levels,options,varargin)
%
% Fu, Fd = cdf for Lagrange space wave slopes
%         at up- and down-crossings of specified levels
%         For time waves, use spec2timeslopecdf
%
% y       = cdf calculated at y
% levels  = vector of relative levels (default = [-1 0 1 2])
% options = struct with fields (plus some more)
%         .lp      = if .p and .lbeta exist, the output x is modified
%                   Lagrange with extra  $-\beta/(-i\omega)^p$  in the transfer
%                   function
%         .lalpha  = alpha value for modified Lagrange
%         .lbeta   = beta value for modified Lagrange
%
% Example:
% S=jonswap; mom=spec2mom(S);
% opt=simoptset('du',0.125,'Nt',512,'dt',0.25);
% levels=[0 1 2];
% y=linspace(-2,2,1001);
% [Fu,Fd]=spec2spaceslopecdf(S,y,levels,opt)
% clf
% subplot(211)
% plot(Fu.x,Fu.f{1},Fu.x,Fu.f{2},Fu.x,Fu.f{3}); hold on
% plot(Fd.x,Fd.f{1},'-.',Fd.x,Fd.f{2},'-.',Fd.x,Fd.f{3},'-.')
% [Fu,Fd]=spec2spaceslopecdf(S,y,levels,opt,'lalpha',1)
% title('Slope CDF at space up- and downcrossings, symmetric waves')
% axis([-2 2 0 1])
% subplot(212)
% plot(Fu.x,Fu.f{1},Fu.x,Fu.f{2},Fu.x,Fu.f{3}); hold on
% plot(Fd.x,Fd.f{1},'-.',Fd.x,Fd.f{2},'-.',Fd.x,Fd.f{3},'-.')
% title('Slope CDF at space up- and downcrossings, asymmetric waves')
% axis([-2 2 0 1])
%
% See also: spec2spaceslopecdf, spec2timeslopecdf, spec2ldat,
%          ldat2lwav, wav2slope

```

```

function [fu,fd]=spec2spaceslopepdf(S,y,levels,opt,varargin)
%SPEC2SPACESLOPEPDF Computes pdf for slope at crossings of space waves
%
%CALL: [fu,fd] = spec2spaceslopepdf(S,y,levels,options,varargin)
%
% fu, fd = pdf for Lagrange space wave slopes
%          at up- and down-crossings of specified levels
%          For time waves, use spec2timeslopecdf
%
% y       = pdf calculated at y
% levels  = vector of relative levels (default = [-2 -1 0 1 2])
% options = struct with fields (plus some more)
%   .lp    = if .p and .lbeta exist, the output x is modified
%            Lagrange with extra  $-\beta/(-i\omega)^p$  in the transfer
%            function
%   .lalpha = alpha value for modified Lagrange
%   .lbeta  = beta value for modified Lagrange
%   .ltheta = if exist produces [w,x] to be transformed, theta =
%            angle for the transformation
%   .lp, .lbeta and .ltheta are empty ([]) by default
%   .plotflag - 'off', no plotting (default)
%             - 'on'
%
% Example: See example for spec2spaceslopepdf
%
% Ref: Lindgren & Aberg, JOMAE, 131 (2009) 031602-1
% See also: spec2ldat, spec2timeslopecdf, ldat2lwav, wav2slope

```

```

function [Slopes,Steep,Data]=spec2steepdat(S,Nsim,type,lev,opt,varargin)
%SPEC2STEEP DAT Simulates Lagrange waves and extracts steepness and slopes
%
%CALL: [Slopes,Steep,Data] = spec2steepdat(S,Nsim,type,levels,options)
%
% Slopes      = struct with fields
%   .up       = observed slopes at the up- and
%   .down     = down-crossings of specified levels
%   .meanup   = average wave profiles near up- and crossings
%   .meandown = down-crossings of mean water level
%   .meanx    = at corresponding times
%   .A        = asymmetry measure by Hilbert transform skewness
%   .lambdaAL= asymmetry measure slope ratio at mean crossing
% Steep       = struct with fields
%   .ffull    = full front steepness as measured by L',L'' etc
%   .bfull    = full back steepness
%   .fhalf    = half front steepness
%   .bhalf    = half back steepness
%   .lambdaN  = asymmetry measure according to front/back half period
% Data        = struct Lagrange waves and two derivatives
%
% S           = orbital spectrum
% Nsim        = number of replicates in simulation (default = 1)
% levels      = vector of standardized levels relative to
%              (default levels = [-1 0 1 2 3]*standard deviation )
% type        = 'time', or 'space'
% options     = struct with fields for individual replicates
%   .plotflag - 0, no plotting (default)
%             - 1, plotting of waves and cross-covariance
%             - 2, plotting of average waves
%             - 3, both the above
%
% See also: spec2ldat, ldat2lwav, wav2slope

```

```

function [TTFu,varargout]=spec2timeslopecdf(S,y,lev,opt,varargin)
%SPEC2TIMESLOPECDF Computes cdf for slopes at crossings of time waves
%
%CALL: [TTFu,TTFd,STFu,STFd,VTFu,VTFd] = ...
%           spec2timeslopecdf(S,y,levels,options,varargin)
%
% XTFu, XTFd = cdf for slopes at up- and down-crossings of levels
%             according to Sections 5.1.1, 5.1.2, 5.1.3 in
%             Adv Appl Probab 42 (2010) 489-508.
%             TT = time slopes at time crossings
%             ST = space slopes at time crossings
%             VT = velocity at time crossings
%             Any number of output cdf:s can be specified,
%             starting with TTFu, [with optional TTFd, ...]
%             Note: Crossings with retrograd x-movement not included.
%
%             For space waves, use spec2spaceslopecdf
%
% y           = cdf calculated at y
% levels      = vector of relative levels (default = [-1 0 1 2])
% options     = struct with fields (plus some more)
%   .lalpha   = alpha value for modified Lagrange
%   .lbeta    = beta value for modified Lagrange
%   .plotflag = 'off', no plotting (default)
%             - 'on'
%
% Example:
% S=jonswap; mom=spec2mom(S);
% opt=simoptset('du',0.125,'Nu',512,'dt',0.125);
% levels=[0 1 2]*sqrt(mom(1));
% y=linspace(0,8,1001);
% [TTFu,TTFd]=spec2timeslopecdf(S,y,levels,opt,'lalpha',1)
% clf
% plot(TTFu.x,TTFu.f{1},TTFu.x,TTFu.f{2},TTFu.x,TTFu.f{3}); hold on
% plot(TTFd.x,TTFd.f{1},'-.',TTFd.x,TTFd.f{2},'-.',TTFd.x,TTFd.f{3},'-.')
% title('Slope CDF at up- and downcrossings, asymmetric time waves')
% axis([0 8 0 1])
%
% See also: spec2ldat, spec2slopepdf, ldat2lwav, wav2slope

```

```

function [TTfu,varargout]=spec2timeslopepdf(S,y,levels,opt,varargin)
%SPEC2TIMESLOPEPDF Computes pdf for slopes at crossings of time waves
%
%CALL: [TTfu,TTfd,STfu,STfd,VTfu,VTfd] = ...
%           spec2timeslopepdf(S,y,levels,options,varargin)
%
% XTfu, XTfd = pdf for slopes at up- and down-crossings of levels
%             according to Sections 5.1.1, 5.1.2, 5.1.3 in
%             Adv Appl Probab 42 (2010) 489-508.
%             TT = time slopes at time crossings
%             ST = space slopes at time crossings
%             VT = velocity at time crossings
%             Any number of output cdf:s can be specified,
%             starting with TTFu, [with optional TTFd, ...]
%             pdf is computed from a smoothed gradient of the
%             simulated cdf; see spec2timeslopecdf
%
%             For space waves, use spec2spaceslopepdf
%
% y           = pdf calculated at y
%             Note: the accuracy will depend on the y-spacing
%             and on an interior smoothing parameter smoothp
% levels      = vector of relative levels
%             (default = [-1 0 1 2])
% options     = struct with fields (plus some more)
%             .lalpha = alpha value for modified Lagrange
%             .lbeta  = beta value for modified Lagrange
%
% Example:
% S=jonswap; mom=spec2mom(S);
% opt=simoptset('du',0.125,'Nu',256,'dt',0.125);
% levels=[0 1 2]*sqrt(mom(1));
% y=linspace(0,8,101);
% [TTfu,TTfd]=spec2timeslopepdf(S,y,levels,opt,'lalpha',1)
% clf
% plot(TTfu.x,TTfu.f{1},TTfu.x,TTfu.f{2},TTfu.x,TTfu.f{3}); hold on
% plot(TTfd.x,TTfd.f{1},'-.',TTfd.x,TTfd.f{2},'-.',TTfd.x,TTfd.f{3},'-.')
% title('Slope PDF at up- and downcrossings, asymmetric time waves')
% axis([0 8 0 1])
%
% See also: spec2ldat, spec2slopepdf, ldat2lwav, wav2slope

```

```
function Slope=wav2slope(L,lev,dense,p)
%WAV2SLOPE Extracts slopes at up- and downcrossings after smoothing
%
%CALL: Slope=wav2slope(L,levels,dense,p)
%
% Slope = structure with observed up- and down-crossing slopes
%
% L = data with fields L.t/L.u and L.Z
% levels = vector with absolute levels; (required field, no default)
% dense = interpolation rate,
%         []: no interpolation or smoothing
%         positive integer: data are interpolated at rate
%         dense at equidistant points (default: dense=20)
% p = [0...1] is a smoothing parameter (default=1)
%     0 -> LS-straight line
%     1 -> cubic spline interpolation
```


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