

Euler Simulation, Python

Contents

1	Introduction	1
2	Code Documentation	2
2.1	simulator.py	2
2.1.1	Simulator1D: Constructor	2
2.1.2	Simulator1D: step()	3
2.1.3	Simulator1D: calculate_time_derivatives()	3
2.1.4	Simulator1D: volume()	4
2.1.5	Simulator1D: peak_location()	4
2.1.6	Simulator1D: zeta_at()	4
2.1.7	Simulator1D: run_simulation()	4
2.1.8	Simulator1D: data_save_params()	6
2.1.9	Simulator1D: soliton()	6
2.1.10	Simulator1D: KY_bathym()	7
2.1.11	Simulator1D: KY_sim()	7
2.1.12	Simulator1D: fields	7
2.2	integrator.py	9
2.2.1	Integrator1D: euler	9
2.2.2	Integrator1D: RK4	9
2.2.3	Integrator1D: implicit_midpoint	9
2.2.4	Integrator1D: AM1	10
2.2.5	Integrator1D: DIRK3	10

1 Introduction

The Euler Simulation project applies the methods used in Knowles and Yeh (2018) to simulate, in one dimension, the shoaling process of a shallow-water wave. The original paper uses such a simulation to predict the wave amplification process, that is, how a , the amplitude of the wave, relates to h , the local water depth.

This project uses the same algorithm, in python, to handle other cases, e.g. the wind's effect on a shoaling solitary wave.

2 Code Documentation

Simulation involves two python files.

2.1 simulator.py

simulator.py contains one relevant class: `Simulator1D`, which wraps necessary functionality to run a simulation.

2.1.1 Simulator1D: Constructor

```
sim = Simulator1D(bathymetry, dt, dx, eta0, phiS0)
```

Initializes a Simulator instance with the given parameters:

argument	description
bathymetry	numpy array of the bathymetry, must have an even number of nodes. 0 is expected to be water level.
dt	time resolution of the simulation
dx	spatial resolution of the simulation (distance between points of bathymetry)
eta0	initial free surface heights (with spatial resolution dx, 0 is expected for still water) expected to be a numpy array
phiS0	velocity potential at the free surface, expected to be a numpy array.

`Simulator1D` has the following keyword arguments:

argument	description	default
zeta_x	First derivative (gradient) of the bathymetry, if a higher order approximation of gradient is desired. If no argument for zeta_x is passed, it is instead calculated from the finite difference $f'(x) = \frac{f(x+h) - f(x-h)}{2h} + O(h^2)$ where the edges are assumed to have a gradient of 0	(see description)
M	Terms in the pertubation expansion (higher number is more accurate, but requires more computation)	5
v	lowpass threshold. If v is a number, any wavenumber greater than the largest wavenumber times v is clipped off each timestep. v can also be a function that takes a wavenumber magnitude and the peak wavenumber, and returns how it should be scaled	0.7
g	acceleration due to gravity.	9.81
h0	base still water depth	bathymetry[0]

In the case that v is a function, the wave number is the first argument and the peak wavenumber is the second argument, so the following will result in the same low-pass filter $v = 0.7$:

```
def v(k, peak):
    if k/peak <= 0.7:
        return 1
    else:
        return 0
```

2.1.2 Simulator1D: step()

```
sim.step("RK4")
```

Steps the simulation forward using the given method. Any arguments for the method can be specified as optional arguments or keyword arguments.

argument	description
method	The method to use. This can be a string or a function. When a string is passed the method in <code>integrator.py</code> of the same name is used. See section (2.2) for such methods.

Optional arguments specific to a method (e.g. tolerance for implicit methods) can be passed into `step()`, which will be transferred to the method when called. In particular, `P_atmos` is common to all integration methods, and is shown below.

argument	description	default
P_atmos	Atmospheric pressure at the surface. This can be either a function with arguments <code>eta, phiS, eta_x, phiS_x, w</code> or a numpy array of length <code>Nx</code> . See the beginning of section (2.2) for more specifics on the function.	array of zeros

2.1.3 Simulator1D: calculate_time_derivatives()

```
sim.calculate_time_derivatives(eta, phiS, zeta, zeta_x, zeta_t, P_a)
```

Returns the tuple `(eta_t, phiS_t)` of time derivatives of η and Φ^S respectively. Takes the following arguments:

argument	description
eta	free surface height at the given time step; pass simulator.eta if you want the current time derivative.
phiS	free surface velocity potential at the given time step; pass simulator.phiS if you want the current time derivative
zeta	bathymetry (ζ), with 0 corresponding with a depth of -h0
zeta_x	spatial derivative of ζ
zeta_t	time derivative of ζ
P_a	atmospheric pressure at every point, should have the same samples as bathymetry. Expected to be a numpy array or a function. See the beginning of section (2.2) for more specifics on the function.

2.1.4 Simulator1D: volume()

`sim.volume()`

Returns the integral

$$\int \eta \, dt$$

over the bounds of the simulation, which represents the volume of water in the simulation, offset by a constant that depends only on the bathymetry. This value should be invariant in the simulation, and can give a means of measuring the accuracy of the simulation.

2.1.5 Simulator1D: peak_location()

`sim.peak_location()`

Returns

$$\arg \max_x \eta(x),$$

the value of x that corresponds with the heighest point of the surface. This value is equivalent to the index of the highest eta, times `dx`, and provides a means of finding the approximate position of a solitary wave.

2.1.6 Simulator1D: zeta_at()

`sim.zeta_at(x)`

Returns $\zeta(x)$, using a linear interpolation scheme for non-discrete points. calling `zeta_at(i*dx)` is equivalent to evaluating `zeta[i]`.

2.1.7 Simulator1D: run_simulation()

`sim.run_simulation(plot_dt, data_dt, directory)`

Runs a simulation, time-stepping until a stop-condition is met, and saving plots and/or data at given intervals of time.

argument	description
saveplot_dt	the timestep between saved plots. This number is rounded to the nearest multiple of the simulation dt. If this value does not round to a positive number, plots are not saved. Plots are saved as PNG files with a name corresponding to the order it is saved in. A plot with number 'i' represents the data at time 'i*saveplot_dt'
savedata_dt	the timestep between saved data. This number is rounded to the nearest multiple of the simulation dt. If this value does not round to a positive number, data is not saved. Data is saved as a json file with name 'dat.json' which is created regardless if data should be saved or not. In the case that data is not saved, only the metadata of the simulation is stored.
directory	the directory to save the files to. This can also include a prefix to the file. If directory =" <code>~/sim/</code> ", then plots are saved as " <code>[number].png</code> " in the <code>~/sim/</code> directory. If directory =" <code>~/sim</code> ", then plots are saved as " <code>sim[number].png</code> " in the home directory. If directory =None, no files are saved.

`run_simulation` has the following optional arguments:

argument	description	default
should_continue	function that determines if a simulation should stop or not. This takes the simulation as an argument and returns a boolean. The simulation is run until <code>should_continue</code> returns false. By default, this is the lambda function <code>sim: sim.t > 500</code> , which stops the simulation when a time of 500 is reached	(see description)
integrator	function or string that timesteps the simulation. functions should only take the simulation as an argument and return nothing, modifying the passed simulation. Strings should be the name of a method in <code>Integrator1D</code> , which will be called by the simulation.	"RK4"
plot_kwargs	The keyword arguments to be passed into <code>matplotlib.pyplot.savefig()</code>	{}
save_eta	Parameters for how eta should be saved when data is saved. This should be generated using <code>Simulator1D.data_save_params()</code> . If None, then eta is not saved.	None
save_phi	Parameters for how phiS should be saved when data is saved. This should be generated using <code>Simulator1D.data_save_params()</code> . If None, then phiS is not saved.	None
loop_callback	this function is called after every simulation step. It should be a void function that takes the arguments sim , step , plot , data , where sim is the simulator at the step step is the integer multiple of dt that the simulation has run plot is a boolean representing if the plot was saved this step data is a boolean representing if the data was saved this step By default, <code>loop_callback</code> makes a print statement after every 100 time steps.	(see description)

2.1.8 Simulator1D: data_save_params()

`Simulator1D.data_save_params()`

Returns a dictionary of parameters for how to save data from a simulation. The output of `data_save_params()` should be used for arguments `save_eta` and `save_phi` in `run_simulation()`.

argument	description	default
<code>dx</code>	The spatial resolution to save with. If None, then the resolution is the same as the simulation. This value will always be rounded to a whole number multiple of the simulation dx.	None
<code>point_conversion</code>	A boolean that represents if data should be coded as a vector (array), or if the vector should be converted into a list of (x,y) points. If true, then the conversion is made.	False
<code>eps</code>	The tolerance of the save data. The data is rounded to the nearest multiple of eps. That is, with <code>eps=0.001</code> , the data is saved up to the 3rd decimal place. 0 corresponds with no rounding.	0
<code>lin_tol</code>	Only used when <code>point_conversion</code> is true. Specifies a tolerance for which points should not be saved when they are close enough to a linear interpolation of the data. If the points are $\{(0,0),(0.5,0.5),(1,1)\}$, any nonnegative tolerance will discard (0.5,0.5). If no points should be discarded, a negative value should be given.	-1
<code>zero_trunc</code>	If a value is less than this distance from 0, the value is truncated to 0 before saving.	0

2.1.9 Simulator1D: soliton()

`Simulator1D.soliton(x0,a0,h0,Nx,dx)`

Returns a tuple (`eta`,`phiS`) corresponding to the initial conditions of η and Φ^S of a soliton at a given point in space.

argument	description
<code>x0</code>	The x coordinate of the soliton, where <code>x=0</code> corresponds with an index of 0 in the vectorization of η and Φ^S
<code>a0</code>	The amplitude of the soliton
<code>h0</code>	The water depth beneath the soliton
<code>Nx</code>	The number of points in the vectorization of η and Φ^S
<code>dx</code>	The spatial resolution (distance between points)

argument	description	default
<code>g</code>	acceleration due to gravity	9.81

This can be used in the constructor of `Simulator1D` through unpacking:

```
sim = Simulator1D(bathymetry, dt, dx, *Simulator1D.soliton(x0,a0,h0,Nx,dx))
```

2.1.10 Simulator1D: KY_bathym()

Simulator1D.KY_bathym()

Produces a bathymetry profile similar to Knowles and Yeh's paper. Expects $h_0 = 1$, but the result can be multiplied by the desired h_0 if not 1.

argument	description	default
Nx	number of points	2^{14}
dx	spatial resolution (distance between each point)	0.04
s0	nominal slope of the bathymetry	0.002
d0	height of the beach plateau	0.9
gamma	smoothing parameter	0.1
X1	position where the bathymetry should start sloping up	4

2.1.11 Simulator1D: KY_sim()

sim = Simulator1D.KY_sim()

Returns a new simulator similar to Knowles and Yeh's initial conditions. The bathymetry is produced by KY_bathym() and the initial η and Φ^S values are produced by soliton().

argument	description	default
Nx	Number of points (nodes) in the discreteized simulation	2^{14}
dx	Spatial resolution	0.04
dt	Temporal resolution (time step)	0.01
s0	Slope of the bathymetry	1/500
x0	location of the center of the starting soliton	30
a0	amplitude of the soliton	0.1
h0	depth of the water	1

2.1.12 Simulator1D: fields

In an instance of Simulator1D, the following fields may be of importance:

argument	description
dt	Temporal resolution (time step) of the simulation. This variable is used by an integrator when stepping the simulation. Most methods will use this time step, but an adaptive method may use a timestep that is smaller. This can be modified externally.
dx	Spatial resolution. This is the distance between two points of a vectorized function of x . This should not be modified externally.
eta	Vectorized η with a spatial resolution \mathbf{dx} at the current time step. This can be modified externally, but must have the same length (Nx).
phiS	Vectorized Φ^S with a spatial resolution \mathbf{dx} at the current time step. This can be modified externally, but must have the same length (Nx).
M	Terms in the perturbation expansion of Φ . Calculating the vertical velocity Φ_x scales approximately $O(M^2)$. This can be modified externally.
g	The acceleration due to gravity in this simulation. This can be modified externally.
h0	base still water depth. We approximate the bathymetry as h0 in many calculations. This can be modified externally.
zeta	Vectorized bathymetry (ζ), offset so $\zeta = 0$ corresponds with $z = -h0$. This can be modified externally, but must have the same length (Nx). Additionally, zeta_x should also be changed to the gradient of the new bathymetry.
zeta_x	Vectorized bathymetry gradient ($\nabla\zeta$). This can be modified externally, but must have the same length (Nx). Additionally, zeta should also be changed to match the new bathymetry.
Nx	Number of points used in the discretization (vectorization) of the simulation along the x -axis. This should not be modified.
sim_length	distance in x that the simulation uses. The vectorizations of η , Φ^S , and ζ have the domain $[0, \text{sim_length})$. This should not be modified.
x	Vectorized domain. It holds that $\mathbf{x}[\mathbf{i}] = \mathbf{dx} * \mathbf{i}$. This should not be modified.
kxdb	<p>The double-domain of wave number. When performing an FFT on a function f with spacing dx, the values of the output correspond to the wavenumber by index. If V is such a vectorization of the function f, then</p> $(\mathcal{F}(f))(\mathbf{kxdb}[\mathbf{i}]) \approx \text{FFT}(V)[\mathbf{i}]$ <p>where \mathcal{F} is the continuous fourier transform. This should not be modified.</p>
kappadb	Normalized wavenumber of kxdb. This is equivalent to $\mathbf{abs}(\mathbf{kxdb})$. This should not be modified
chi	Vectorized low-pass filter. Any function in the wavenumber domain can apply the filter by pointwise multiplication. This can be modified externally, but kxdb_im must also be modified accordingly.
kxdb_im	The precomputed value $ik\chi(k)$, which is computed from $\mathbf{complex}(0,1) * \mathbf{kxdb} * \mathbf{chi}$. This should only be modified when chi is modified.
t	The time the simulation has run. This is only ever incremented by an integration method inside a <code>step()</code> call, and can be freely modified and accessed externally.

2.2 integrator.py

This python file has the class `Integrator1D` which contains only static members. Each of which is a function that takes a `Simulator1D` instance, an atmospheric pressure argument `P_atmos`, and potential optional arguments.

`P_atmos` can either be a numpy array or a function that takes in the arguments `eta, phiS, eta_x, phiS_x, w` and returns a numpy array. The array should have a length equal to the number of nodes `Nx` used in the simulator, which matches the length of the `bathymetry` array passed into the constructor. For example, when measuring the wind effect, one may consider

$$P_a = P \frac{d\eta}{dx}$$

for some constant P . Such a function can be expressed as

```
def P_atmos(eta, phiS, eta_x, phiS_x, w):
    return P * eta_x
```

2.2.1 Integrator1D: euler

Referenced by string "euler". Makes one derivative calculation per step, using the method:

$$y_{n+1} = y_n + hf'(y_n)$$

2.2.2 Integrator1D: RK4

Referenced by string "RK4". Makes 4 derivative calculation per step, using the classic 4 step, 4th order, Runge-Kutta method with the Butcher tableau:

0	0	0	0	0
$\frac{1}{2}$	$\frac{1}{2}$	0	0	0
$\frac{1}{2}$	0	$\frac{1}{2}$	0	0
1	0	0	1	0
	$\frac{1}{6}$	$\frac{1}{3}$	$\frac{1}{3}$	$\frac{1}{6}$

2.2.3 Integrator1D: implicit_midpoint

Referenced by string "implicit_midpoint". Uses the implicit midpoint rule:

$$y_{n+1} = y_n + hf\left(\frac{y_n + y_{n+1}}{2}\right)$$

This equation is solved using fixed point iteration after an initial guess from euler's method.

Takes additional arguments:

argument	description	default
max_iters	The most iterations used to achieve the desired tolerance, after which, RK4 is defaulted to.	100
tol	The tolerance allowed for the iteration to stop.	10^{-10}

2.2.4 Integrator1D: AM1

Referenced by string "AM1". Uses the one step Adams-Moulton method, the implicit trapezoidal rule:

$$y_{n+1} = y_n + h \frac{f(y_n) + f(y_{n+1})}{2}$$

This equation is solved using fixed point iteration after an initial guess from euler's method.

Takes additional arguments:

argument	description	default
max_iters	The most iterations used to achieve the desired tolerance, after which, RK4 is defaulted to.	100
tol	The tolerance allowed for the iteration to stop.	10^{-10}

2.2.5 Integrator1D: DIRK3

Referenced by string "DIRK3". Uses Nørsett's 3 stage, 4th order diagonally implicit Runge-Kutta method with the Butcher tableau:

x	x	0	0
$\frac{1}{2}$	$\frac{1}{2} - x$	x	0
$1 - x$	$2x$	$1 - 4x$	x
	$\frac{1}{6(1-2x)^2}$	$\frac{3(1-2x)^2-1}{3(1-2x)^2}$	$\frac{1}{6(1-2x)^2}$

where $x = 1.06858$. This equation is solved using fixed point iteration for each stage, where each stage has an initial guess of the former stage, with the first stage's initial guess as the derivative at y_n . For example, if k_1 and k_2 are the results of the first and second stage respectively, the second stage solves

$$k_2 = f \left(y_n + \left(\frac{1}{2} - x \right) h k_1 + x h k_2 \right)$$

with an initial guess $k_2 = k_1$.

Takes additional arguments:

argument	description	default
max_iters	The most iterations used to achieve the desired tolerance, after which, RK4 is defaulted to.	100
tol	The tolerance allowed for the iteration to stop.	10^{-10}