PHAS0030: Computational Physics  
Session 4: Ordinary Differential Equations

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In this session, we will consider methods to solve ordinary differential equations of different orders. We will see that this is not a straight-forward problem, with stability problems being very common. We will discuss methods that are stable.

# Objectives

The objectives of this session are to:

* understand how we can create a numerical solution for a differential equation
* explore the stability of the simplest approach, Euler’s method
* examine more stable methods, such as predictor-corrector and Runge-Kutta
* consider boundary conditions and how different methods are applicable depending on the boundary conditions

# Review of Session 3

In the third session, we looked closely at the topic of discretization, covering:

* Grids that are commonly used in calculations
* The finite difference approach to differentiation on a grid
* Numerical integration in one dimension,  
  including the trapezoidal rule and Simpson’s rule

We will continue to use discrete space and time variables throughout the next three sessions, where we will consider the numerical solution of differential equations.

# Introduction

Ordinary differential equations (ODEs) are equations for a function, often written as , of one independent variable, , and its derivatives (including polynomials in ). The most common, and certainly most easily solved, class is that of linear ODEs, where the unknown function and its derivatives only appear to first order. We can write a general linear ODE as:

where indicates the differential , is an arbitrary function of and is a source term (again, an arbitrary function of ). Examples of this kind of equation include: classical mechanics (particle motion), LRC circuits and harmonic oscillators. These are all second order linear equations, for instance:

for the charge in an LRC circuit: here, and , the driving term , and the coefficients are independent of .

The equation of motion for the angle made by a pendulum relative to the vertical is linear for small angles, but becomes non-linear for large angles (this is defined by the point at which the approximation breaks down). A simple model of radioactive decay is a first-order linear ODE: , where is the number of particles in the sample at a given time and is related to the half-life.

We will consider *partial* differential equations (PDEs) in the next two sessions; these are differential equations that feature more than one independent variable. Examples include the wave equation, Maxwell’s equations for electromagnetism, heat flow and Schrödinger’s equation (in both time-dependent and time-independent forms).

The most common form of ODE is one where *time* is the independent variable; while we use commonly in these notes, it may well often be substituted by .

## Specifying conditions

In order to solve a differential equation, we must specify some conditions on the problem. The number of conditions depends on the order of the equation, so a first order equation will require one condition, and a second order equation will require two. For instance, the simple model of radioactive decay given above requires the number of atoms at . The LRC circuit might require an initial charge, , and current . Once we move beyond first order equations, there are *two* ways to specify the conditions:

* Initial value problems, where all conditions are specified at one point in space, , or time, ; for a second order ODE, we would specify and .
* Boundary value problems, where conditions are specified for only one variable, but at extremes; for a second order ODE, we could specify the value of at two points, and .

It is of course possible to solve in more than one dimension (this is particularly common for classical mechanics). Note that in this case we still have the same *order* of differential equation (second order for problems involving non-zero acceleration, say) but the equation is now a *vector* equation. The initial conditions that must be specified are now vector quantities.

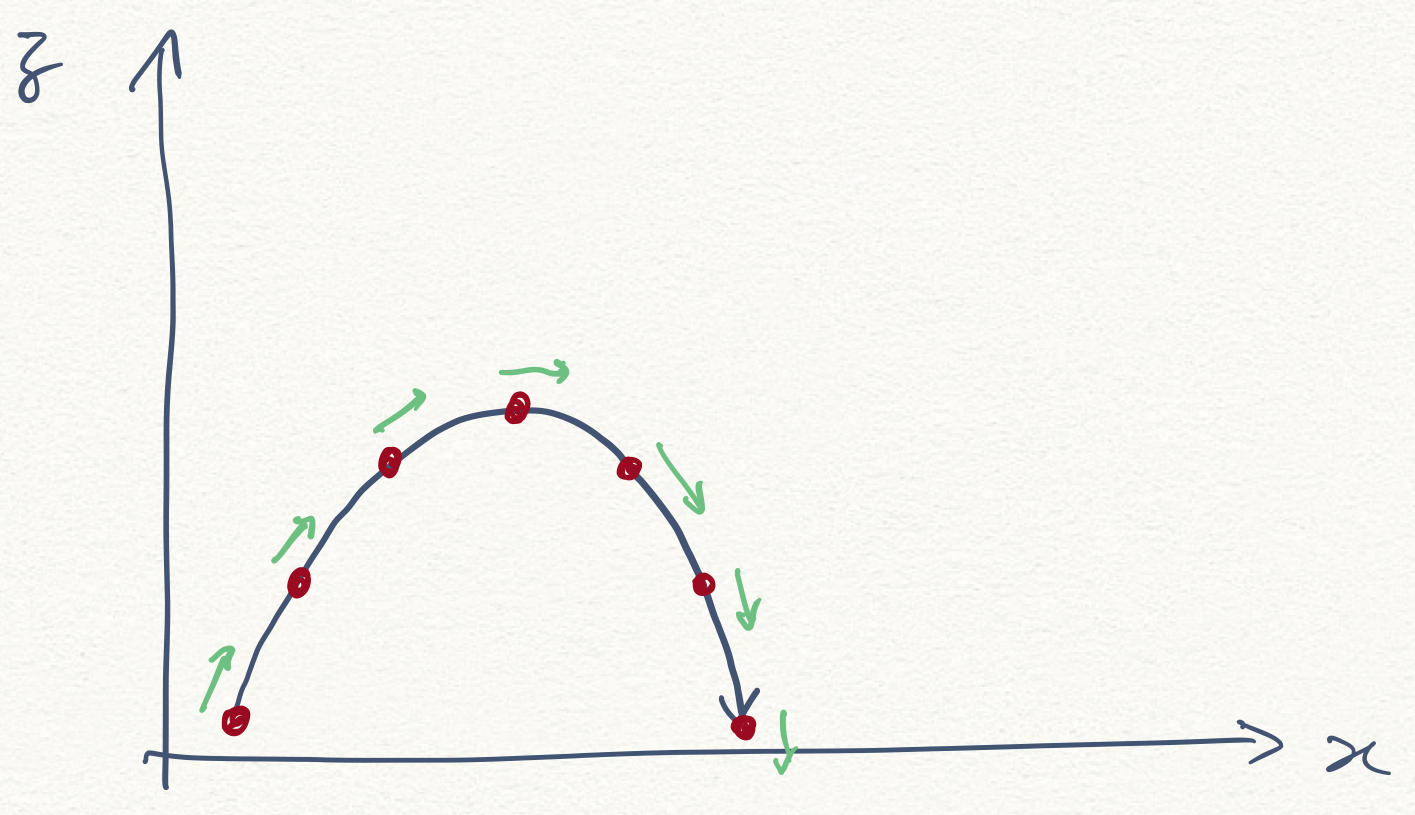
## Discretization again

The approach that we will take to solve differential equations will be based on the finite difference work that we did in the last session. We will substitute the finite difference form of the differential, and then perform numerical integration. If we are working with then you can think of the approach as stepping forward in time by a step ; at each point in time, we calculate the appropriate differential and use that to update .

It is useful to consider a Taylor expansion:

This will enable us to think about errors in our approach.

The overall aim of a numerical solution is to calculate for a series of points . The way that we do this is via step-wise numerical integration: from and we can evaluate at , and therefore find ; we then evaluate at , and find ; and so on. This is illustrated very roughly in the sketch below.



A sketch of a numerical integration for the trajectory of a ball, indicating the velocity at each point (green arrow) in the process. The gaps between points correspond to the timestep.

# Solving first-order ODEs: Euler’s method

Euler’s method is a very simple approach to solving ODEs, so we will start with it. But it is often a very poor choice, prone to instabilities and should not generally be used for practical calculations; after we introduce the basic concept, we will turn to more reliable and accurate approaches. If we return to the simple definition of a (finite) differential, we find:

In terms of array indexing, we can write this as:

This is equivalent to the first-order Taylor expansion, and indicates one way in which we could solve a first order equation: starting from some initial condition, , we could integrate with steps of size to get values of at successive values of .

Why might this approximation be a poor choice? Our experience with the forward difference formula from last session gives us a hint—that formula was only accurate to *first order* in the difference. To examine the accuracy of integration, we need to find the order of the error. At each step, we know from the Taylor expansion that the leading term in the error in will be:

(where indicates the order of the term). However, this is just the error in a *single* step. The global (or cumulative) error when we reach a particular point will be the sum of the errors for the number of steps required to reach from : , so that the global error ( times the error in each step) will be of order .

We could have guessed that the method would be a bad approach from our analysis of finite differences. Last week, we saw that the centred difference approach was much more accurate than a forward difference (second order in step size), and in Session 7 we will see that we can use the same ideas to make a stable equivalent for differential equations (these are known as Verlet approaches). In this session, we will look at alternative routes to improve the integration.

## Stability analysis

In the exercises, we will see that the Euler method is unstable, particularly for step sizes that pass a certain threshold. Why might that be? Let’s consider a very simple differential equation, and analyse it[[1]](#footnote-1).

This clearly has the analytic solution , with found from the initial condition . (Notice that this is a very common equation in physics and elsewhere, for instance giving a simple model of radioactive decay.)

What would the Euler method give us for an update step?

So when we apply this to steps, we find:

This very simple example immediately indicates some of the issues that will occur if we choose unwisely. We have a stable solution if , while if , then while at each step the solution will decay, it will *alternate* in sign (clearly wrong given the analytic solution). If , then the overall magnitude will *increase* with . Clearly the choice of step size depends on key parameters in the problem, and can have significant effects on the solution. We will return to the question of stability in later sections.

## Exercises

### In class

1. Using a simple loop, solve the differential equation using Euler’s method. For flexibility, set up as a variable, calculate the possible values of and store the results in an array. You will also need to store in an array. Use and solve for , with . (You might find the basic set-up below useful.)

* *# Specify step size and simulation length*  
  dx = 0.5  
  total\_x = 20  
  N = int(total\_x/dx)  
  x = np.linspace(0,total\_x,N+1)  
  *# Initial condition*  
  y0 = 1.0  
  k = 1.2  
  y = np.zeros(N+1)  
  y0 = 1.0  
  y[0] = y0

Plot the approximate and exact solutions.

1. Now write a function to calculate N steps of the Euler solver. Use the specification below:

def euler\_solver(fun\_rhs,y0,dt,N):  
 *"""Solve dy/dt = fun(y,t) using Euler's method.  
 Inputs ...  
 Returns: array of length N+1 with values of y  
 """*

At the start of the function, you will need to create an array to store results, which you will return (note that it should be length N+1 if you store the initial conditions for t=0). Use this Euler function for the problem in question 1 (you will need to define and pass the function fun\_rhs though in this case it is extremely simple!), and explore how the step size affects agreement. Look at the effect of increasing .

### Further work

1. Create a set of four sub-plots for values of that give divergence, oscillation, stability but poor agreement, and good agreement respectively. Use the fig = plt.figure, ax = fig.add\_subplot, ax.plot protocol that we discussed in Session 1.
2. Solve the equation with using Euler’s method. What values of give stable solutions? How well do these match the exact solution[[2]](#footnote-2)?

## Beyond first order and one dimension

There are many situations where we will have a vector ODE to solve; in terms of classical dynamics we could write:

where is the position vector. This can be solved using Euler’s method in exactly the same was as we did for the scalar, by resolving the components of and . It will give *coupled* first order ODEs (one for the -component, one for the -component, etc):

To implement the equation, we would need to work on an N-component array (for N dimensions) but otherwise there would be no change.

But we can also use this approach to solve *second order* equations, separating them into *two coupled* first order equations. Consider simple dynamics, where we have the basic equation:

If we work in terms of position and velocity , then we can rewrite this equation as two *coupled* first-order equations (compare to the form above if you like):

To make this clearer, consider simple harmonic motion, where . We could write this as a matrix equation (using :

Notice that we will now need *two* initial conditions: one for and one for .

To implement this, we start by defining a NumPy array which contains the two variables, say y = np.array([x,v]) The Python function for the right-hand side of the equation, , would need to return a two-component array containing the derivatives of and : (v,-k\*x/m) in the case of harmonic motion. This approach is actually a powerful general way to solve second order differential equations on a computer. [[3]](#footnote-3)

If you write your Euler method carefully, it should extend to this type of problem trivially. Consider the following code for the right-hand side of the equation:

def rhs\_sho\_function(x,v,t):  
 *"""Implement RHS of SHO equation for x and v  
 Defines k and m within function"""* k = 1.0  
 m = 1.0  
 *# Note that we make no assumption about what v and x   
 # are: they can be scalars or arrays*  
 dx = v  
 dv = -k\*x/m  
 return dx, dv  
  
*# Sample Euler update step  
# Assume x and v are already defined*dx, dv = rhs\_sho\_function(x,v,t)  
x\_next = x + dt\*dx  
v\_next = v + dt\*dv

This is fine, but does not treat the two components together. The alternative, which is more flexible and efficient computationally, is to pass a single array to the Euler update scheme, and have the update function, , split up and increment the individual components:

def rhs\_sho\_function(y,t):  
 *"""Implement RHS of SHO equation for y (array   
 containing x and v). Note that t is unused here but   
 is passed for compatibility with general  
 solvers. Defines k and m within function"""*  
 k = 1.0  
 m = 1.0  
 *# Separate out for clarity*  
 x = y[0]  
 v = y[1]  
 *# Calculate update*  
 dx = v  
 dv = -k\*x/m  
 return [dx, dv]  
  
*# Sample Euler update step  
# Assume x and v are already defined*  
y = [x, v]  
dy = rhs\_sho\_function(y,t)  
y\_next = y + dt\*dy

In this case, if x and v are N-dimensional arrays themselves then y will be a (2,N) dimensional array, and we can again work with vector quantities.

Finally, we can consider the stability of coupled ODEs, extending our analysis in Section [4.1](#sec:stability-analysis). We will take an equation that can be written as:

If we integrate numerically, we will be applying the matrix repeatedly to our solution, which will accumulate powers of the matrix: . If there is a large ratio between the largest and smallest eigenvalues of , it can be shown that the ODE will be challenging to solve numerically: this ratio is known as the *stiffness*[[4]](#footnote-4). A stable solution for this type of equation can require *very* small steps, or a different kind of solution (known as implicit methods, which we will consider in the next session).

## Exercises

### In class

1. Update your Euler function above to solve more generally for arrays. You can allow the results array y to be of arbitrary dimension (in which case you will need to use the NumPy function to find the right size) or assume a simple two-component problem as seen for the SHO.
2. Apply this function to the simple harmonic oscillator above, and plot your result (you should know what it looks like!). Check the effect of step size.

### Further work

1. Add a damping term[[5]](#footnote-5) to the SHO solver you have created above in question 4 above and explore the effect of the damping coefficient, . Plot your solutions using sub-plots.
2. You could explore adding a driving term on the RHS of the SHO equation if you have time.

# Beyond Euler

There are many methods that improve on the stability and accuracy of Euler. We can gain some insight into their development by considering the question: why is Euler so poor? Part of the answer lies in the use of the gradient at the start of the interval, which means that when we step forwards, we *extrapolate*[[6]](#footnote-6). Various of the common approaches to ODEs attempt to improve on this poor gradient estimate. Note that throughout these notes (not just this section) the label for the independent variable could just as easily have been as and that the examples generally use .

The simplest thing that we have already seen is simply to reduce the timestep in the Euler method. We can see that this is equivalent to taking a larger step but using the average of the gradients at the start and middle of the step:

The predictor-corrector method takes a step (the prediction) and calculates the gradient at the end of the step, and then averages this with the gradient at the *start* (the correction):

There are more sophisticated versions of this approach (often associated with the names Adams, Moulton and Bashforth) which can be extremely accurate and stable.

A different approach, known as the mid-point method, uses an estimate of the gradient half-way along the step:

Notice that this differs from the updated Euler method above, in that we calculate and use the gradient at the mid-point rather averaging the initial and mid-point gradients. It can be shown that this formula is accurate to second-order in .

The mid-point method is an example of a class of methods called Runge-Kutta methods. These use intermediate gradients to improve the accuracy in sub-divisions, and are named by the order of accuracy that they achieve. The most commonly used is RK4: it is fourth order in accuracy () *but* it requires *four* function evaluations. The update scheme can be written as:

We can make a closer link to the arrays used in a typical Python implementation if we write the update as follows:

It is important to realise that, computationally, the function is a *continuous* function (i.e. it is a standard function that you define which takes any values of and ) while and are entries in arrays.

Note that the first quantity, , is the same as the Euler update, while the next two are mid-point evaluations, and the final is an end-point. By combining these, we can eliminate errors below fourth order.

Note that all these methods allow us to solve most ODEs, with a reasonably large step size (though this must always be tested). There are certain classes of equation which require alternative methods, but we will not consider them here.

## Exercises

### In class

1. Write a function to implement a fourth-order RK solver, using the equations above, using your Euler function as a basis (assume that you will solve a second-order equation which will be split into two coupled first-order equations).
2. Apply it to a pendulum, for the moment working simply with the linear solution:

where , the acceleration due to gravity, and , the length of the pendulum. You will need to:

* 1. break this into two coupled equations;
  2. write an appropriate function for the right-hand sides;
  3. write a loop calling your RK4 solver to propagate the motion of the pendulum.
* How large can the step size be while still giving a stable solution? (If you have time, try the same solution with an Euler solver, and compare the necessary step sizes). Be sure to make the initial angle (**in radians!**) small enough that the approximation is good.

### Further work

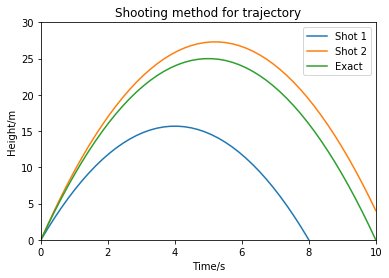
1. Write either a predictor-corrector function or a mid-point function
2. Now model the non-linear pendulum, comparing the RK4 and the function that you coded in question 1 of this further work:

You will need to write a new function for the right-hand side. Start with an angle of , and compare the two methods for the same step size.

1. You should now extend the model further: test an initial angle of just less than (to observe strongly non-linear effects); add a damping term () and test the effect of .

# Boundary Value Problems

Instead of specifying all conditions at the *start* of the simulation, as is done for initial value problems, we can also specify conditions at the *boundaries* of the problem. A simple example is the motion of a projectile, which as a second order ODE requires two pieces of information: we can either specify the position and the velocity at ; or we can specify the position at two different times[[7]](#footnote-7). This gives us a condition to meet *at the end of the simulation*, but does not give enough information to start the solver; we have to guess an initial condition on the velocity, solve to find the position at the end of the simulation, and then update the initial velocity until the boundary condition is met. On a computer, we would use one of the root finding approaches (bisection, secant) we discussed in Session 2.



An illustration of how the shooting method works for a projectile which is required to have height of zero at t=10s. The initial guess (blue line) falls short, while the second guess (orange line) over-shoots. The root finder is able to find the correct initial condition to match the specified boundary conditions with the exact trajectory (green line).

The shooting method starts with a guess for the initial value of the gradient as well as the position at (or ), and then uses a root finder to adjust the initial gradient until the solution matches the specified boundary condition at or The method is sketched in the figure above.

Let’s use the example of the projectile which starts with height 0 and is specified to have height after time . To solve this using a simple root-finding approach we would need to find solutions for the height of the projectile based on two different initial velocities, such that the final heights at bracketed . We would then use (say) the secant method to search for the value of the initial velocity which gave the final height at equal to zero. In terms of the secant method, which usually seeks roots of a function , the independent variable would be the initial velocity, while the function whose roots we are seeking would be the final height.

There is another class of solutions for boundary value problems that we will meet in the next session, called relaxation methods (which include the Gauss-Seidel method that we mentioned in Session 2).

## Exercises

### In class

1. Write a simple piece of code (either as a function or just a loop) to find the initial vertical velocity at which a ball moving vertically under the influence of gravity needs to be launched from the ground at so that it reaches the ground again at . You should use the RK4 solver that you have written, and implement a basic bisection or secant solver, as described in the text above (you could copy this from your answers to Session 2). Plot the trajectory versus time.
2. If you have time, try storing and plotting all the trajectories that the solver works through.

### Further work

1. Try extending the solver so that you are now working in two spatial dimensions ( and , say). You will need to be careful about how you specify your equations—you will need position and velocity for both dimensions (four variables in total). You will need to specify both and at and .
2. If you want to extend this model, you could add an air-resistance term (proportional to the velocity) in one or two dimensions.

# SciPy Functions

There are of course implementations of ODE solvers in SciPy, which generally assume that we have rather than as written above[[8]](#footnote-8). These are all part of the integrate module in SciPy, so you will need to import it as usual: from scipy import integrate.

For initial value problems, there is a general solver which implements various methods:

integrate.solve\_ivp(fun, t\_span, y0, t\_eval = array, args=())

where fun is an implementation of which can take extra arguments (optionally specified by the parameter args=(), though this is only available in SciPy 1.4 or more recent), t\_span is a tuple[[9]](#footnote-9) of two times, the start and end, and y0 gives the initial value. The optional argument t\_eval allows the user to provide an array of times at which to return the solution (otherwise the solver chooses the times). The solver returns a single object; if you call it with a = integrate.solve\_ivp, you can then extract two arrays, a.t and a.y, which give values at times . Remember that if you pass a two-dimensional array for y, then it will return a two-dimensional array. If you want to specify the points at which the result should be returned, you can pass an optional parameter which is an array of times, which should of course lie between the start and end times, t\_eval=your\_array. You can select the method used to solve the problem with the optional parameter method; for a fourth-order Runge-Kutta solver, you would pass the parameter method=’RK45’.

An older function which is still useful, and provides a link to a FORTRAN library, is:

integrate.odeint(func, y0, t, args=())

where func is an implementation of which can take extra arguments (optionally specified by the parameter args=() as we have seen before), y0 is the initial value, and t is an array of times at which to solve for ; the first value should correspond to y0. The function returns an array y with the same number of entries as t. **Be careful:** the order of and differs between these two solvers!

There is also a boundary-value problem solver, solve\_bvp, but its use is somewhat complex (you have to create a function to evaluate the error in the boundary conditions, amongst other things) so we will not consider it here.

## Exercises

### In class

1. Use solve\_ivp and odeint to solve for the linear pendulum model from section [5](#sec:beyond-euler).

### Further work

1. Write a non-linear pendulum function that takes parameters (g, L and a damping term) and pass it to solve\_ivp or odeint to make sure that you understand how optional arguments are passed.

# Assignment

This week, you will investigate two simple gravitational orbit problems, applying two different integrators for first order differential equations (Euler’s method and the velocity Verlet method). You will first consider a small mass in orbit around a large mass with Euler’s method, fixing the large mass (an approximation). You will then consider two masses which are closer in magnitude, in orbit around their centre of mass, with the velocity Verlet method. You can work with a two dimensional coordinate system for this problem. This work will form the opening part of the mini-project, which you will build on independently for the rest of term.

The gravitational force on a mass, *m*1, at a point **r**1 due to another mass, *m*2, located at a point **r**2, is given by:

where **r**12 = **r**2 – **r**1, *r*12 = |**r**12| and you should be careful with signs and directions of forces. For these problems, total energy and angular momentum should be conserved exactly.

1. Write a function to evaluate the force given above (your function should take the positions and masses as inputs). You should assume that G is defined outside the function for now.
2. Now set up the first problem: set G=1 and the masses *m*1 = 0.0001, *m*2 = 1.0 (do not worry about the units) and define a time step and total number of steps, along with a radius of orbit *r*=1. Create arrays to store the positions and velocities of the two bodies, as well as the kinetic and potential energy at each step. Calculate, using pen and paper or equivalent, the initial velocity for the small mass (use simple mechanics: *mv*2*/r = GmM/r*2 but be sure to explain your reasoning). You should fix the large mass (set velocity to zero and do not update its position). You should check that your timestep is suitably small for accuracy, using the magnitude of the velocity as a guide.
3. Propagate this system forward in time using Euler’s method (you may choose how long you propagate, but it should be several periods of the orbit); you should either calculate and store the total energy at each time step, or at the end. The potential energy is given by *U = -Gm*1*m*2*/r*12 and the kinetic energy by the usual form.
4. Plot the resulting positions for the small mass, and (on separate graphs) the total energy and angular momentum against time. (You should be able to work out the angular momentum rather easily – describe this in a text cell.) Comment in a text cell on how well the conservation laws are obeyed, and the orbit that results (you might like to investigate the effect of timestep if you have time).
5. We now turn to a more complicated problem (two bodies, both moving) and a more reliable integrator: the velocity Verlet. The equations for updating position and velocity for a given body are:

You should set up new arrays for the masses, positions and velocities of the bodies in this system, making the masses much closer in magnitude (no more than a factor of ten different). In this case, both bodies will move, in orbit around their centre of mass, **r**CoM = (*m*1**r**1 + *m*2**r**2)/(*m*1+*m*2). It is easiest if you set the centre of mass to be the origin, and align the masses along the *x*-axis initially. Then set the distance between the bodies to 1 (so that *x*1 + *x*2 = 1) and use this equation and *x*CoM=0 to find the initial positions. The initial velocities follow from simple mechanics exactly as in the first case (though be careful: in this case, for the first body, you need *m*1*v*12*/r*1 *= Gm*1*m*2*/r12*2 with *r*1 the distance from the first body to the origin and *r*12 the distance between the two bodies, with an equivalent formula for the second body).

1. Propagate this system forward in time using the velocity Verlet method, ensuring that you cover several periods of the orbit. Plot the orbit, energy and angular momentum as before (though in this case you will need to add the energy and angular momentum for both bodies), and comment in a text cell on the conservation of these quantities with this method. How much better is this method, and what might better mean?

Note that I have also provided the assessment from last year as a separate file (this is *entirely optional*, and for interest/deeper understanding, particularly of boundary value problems; it carries no marks).

# Progress Review

Once you have finished *all the material* associated with this session (both in-class and extra material), you should be able to:

* write a simple Euler solver for first-order differential equations
* understand how mid-point and Runge-Kutta solvers work, and write a function to implement them given the formulae
* use SciPy solvers for ODEs.

You should also check that you understand all the concepts and skills practised in Sessions 1–3.

1. If you find yourself using a computer to solve this equation, then you need to spend some time working on basic mathematics! [↑](#footnote-ref-1)
2. You should be able to work this out yourself; a useful tip is to put the constant of integration with . [↑](#footnote-ref-2)
3. We can also extend it so that and are vectors: each of the arrays would become larger (for three dimensions, they would each be of length six) but no other change would occur. [↑](#footnote-ref-3)
4. We discussed the idea of a characteristic matrix whose eigenvalues determine the behaviour of a system already, when considering conjugate gradients. [↑](#footnote-ref-4)
5. Now you have the equation . [↑](#footnote-ref-5)
6. Extrapolation is almost always dangerous; interpolation is generally much safer. [↑](#footnote-ref-6)
7. We might specify that the height is zero at and some later time, [↑](#footnote-ref-7)
8. These are of course completely equivalent. [↑](#footnote-ref-8)
9. In Python, a tuple is a set of numbers in round brackets, e.g. (0,10). [↑](#footnote-ref-9)