Advanced Algorithms II

Todd Davies

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Overview

This course will explore certain classes of algorithms for modelling and analysing complex systems, as arising in nature and engineering. These examples include: flocking algorithms - e.g., how schools of fish or flocks of birds synchronised; optimisation algorithms; stability and accuracy in numerical algorithms.

Aims

By the end of the course, students should:

- Be able to apply what they learnt across different disciplines.
- Appreciate the emergence of complex behaviours in networks not present in the individual network elements.

Syllabus

- PART 1: NUMERICAL STABILITY AND ACCURACY OF COMPUTATIONS (10 hours)
 - Introduction to finite precision computation (2 hours)
 - Floating point arithmetic (examples that include error analysis insummation, evaluation of polynomials, recurrences, and basic linear algebra)(3 hours)
 - Mixed precision algorithms (basic concept of iterative refinement, different speed of execution on different architectures, linear algebraexamples) (1 hour)
 - Numerical solution of initial value problems (explicit/implicit methods, multistep methods, consistency, stability, convergence) (2 hours)
 - Find the equilibria of a dynamical system, and analyse the stability of said system (2 hours)

- PART 2: COMPLEX NETWORKS AND COLLECTIVE BEHAVIOUR (8 hours)
 - Complex networks are groups of systems (normally, a big number of them) interconnected in a non-trivial and non-regular way Introduction to complex networks (2 hours) Where are the networks and the complexity? Characterisation of complex networks Basic network properties and terminology. Topological analysis
 - Complex network models. The structure of the network (2 hours) Regular networks Random-graph networks Small-world networks Scale-free networks
 - Network dynamics and collective behaviour (3 hours)
 Distributed/local versus centralised/global The concept of self-organisation Synchronisation in complex dynamical networks Consensus over complex networks Swarm dynamics Consensus protocols Flocking algorithms

Attribution

These notes are based off of both the course notes (found on Blackboard). Thanks to the course staff (Eva Navarro-Lopez, Milan Mihajlovic and Pedro Mendes) for such a good course! If you find any errors, then I'd love to hear about them.

Contribution

 $Pull\ requests\ are\ very\ welcome:\ \verb|https://github.com/Todd-Davies/third-year-notes|$

Contents			1 Minimising univariate functions	
1	Initial Value Problems 1.1 IVP's: Linear Multistep Methods	5 6 6 3.	3.1.1 Bisection algorithm	1 1 1 1 1 1 1
2	Complex Networks	10	5 Evolutionary algorithms	
3	Optimisation and nature inspired algorithms: DEPRECATED	3.	7 Cramer & Koza's Genetic Programming	19

1 Initial Value Problems

An initial value problem is an ordinary differential equation with a given value called the initial condition, which is the value of the unknown function (that we're trying to model) at a given point in the domain of the solution.

In the notes, the example of a chef leaving a large pot of soup to cool is given. If the pot starts at $T_0 = 100$, and the desired temperature of the soup is $T_e = 20$, then how long should the chef leave it cooling?

We can't know this until we have the initial conditions, which is that the ambient temperature is $T_{\alpha} = 10$, and the given point in the solution domain; the chef left the soup for ten minutes and the temperature was $T_i = 60$.

The cooling of the soup is modelled by $\frac{dT}{dt}$ (change in temperature over time), and we can use Newton's law of cooling and heating to change this to: $\frac{dT}{dt} = f(T - T_{\alpha})$. Since we know that the ambient temperature is larger than the starting temperature $(T > T_{\alpha})$ and that the soup is actually cooling (as evidenced by $T_i < T_0$), we know that k < 0.

We can work out the value of k from the analytical solution:

$$T(t) = T_{\alpha} + (T_0 - T_{\alpha})e^{-kt}$$

By subbing in the values $t = 600, T(600) = 60, T_{\alpha} = 10$ we can get k:

$$60 = 10 + (100 - 10)e^{-600 \times k}$$

$$50 = 90e^{-600 \times k}$$

$$\frac{5}{9} = e^{-600 \times k}$$

$$log(\frac{5}{9}) = -600k$$

$$k = 9.796 \times 10^{-4}$$

To see how long the cook should leave the soup for, we can simply sub in k and sub in 20 for the desired ending temperature:

$$T(t) = T_{\alpha} + (T_0 - T_{\alpha})e^{-kt}$$

$$20 = 10 + 90e^{-9.796 \times 10^{-4}t}$$

$$\frac{1}{9} = e^{-9.796 \times 10^{-4}t}$$

$$t = \frac{\log(\frac{1}{9})}{-9.796 \times 10^{-4}}$$

$$t = 2242s$$

This was an easy problem though; often, an analytical solution (the equation for T(t)) will not exist and we will need to solve the problem numerically.

1.1 IVP's: Linear Multistep Methods

Basically we will have a function that takes in the current x and y values and output the next y value:

$$y' = f(x, y)$$

Where a < x < b, and we know the 'first' y value $y(a) = \alpha$.

Linear multistep methods aim to emulate a target function using y'. Given y' (a function that produces the next value of a target function y' = f(x, y) for all a < x < b) and $y(a) = \alpha$, we can use Taylor expansion to find the value of any point where b is greater than a:

$$y(x) \approx y(a) + (b - a)y'(a)$$
$$\approx y(a) + (b - a)f(a, y(a))$$

We want to move from a to b in small increments, otherwise any inaccuracies in the Taylor expansion will become quickly apparent, so we define h to be the step size between x_n and x_{n+1} (where x_k are the intermediate steps between a and b), such that $h = \frac{b-a}{N}$, where N is the number of steps we want to take.

This means that $x_n = a + nh$.

For Euler's method, we can describe:

$$y_{n+1} = y_n + (x_{n+1} - x_n)f(x_n, y_n)$$

= $y_n + h f_n$

The Euler method only uses the previous point to generate the next point, but in general, linear multistep methods use the previous k steps to calculate the next value. As such, they are described by a list of values $\alpha = (\alpha_1, \dots, \alpha_k), \beta = (\beta_1, \dots, \beta_k)$ (α_k is always 1), such that:

$$\sum_{i=0}^{k} \alpha_i y_{n+i} = h \sum_{i=0}^{k} \beta_i f_{n+i}$$

The designer of the method determines the coefficients $\alpha_{0,...,k-1}$ (alpha_k is 1 remember) and $\beta_{0,k}$, and they will balance the accuracy of the approximation with the need to make the method easy to apply.

If $\beta_k = 0$, then the method is said to be explicit, since the formula can directly compute y_{n+k} , whereas if $\beta_k \neq 0$, then the y_{n+k} depends on $f(x_{n+x}, y_{n+k})$, which means an equation must be solved to find y_{n+k} :

$$y_{n+k} = h\beta_k f(x_{n+k}, y_{n+k}) + h \sum_{i=0}^{k-1} \beta_i f_{n+i} - \sum_{i=0}^{k-1} \alpha_i y_{n+i}$$

Lets use the Euler method applied to the soup problems as an example. Since $\frac{dT}{dt} = -k(T - T_a)^1$, $f(x_{n+1}, y_{n+1}) = -k(T_n - T_a)$.

Forward Euler
$$T_{n+1} = T_n + h(-k(T_n - T_a))$$

Backward Euler $T_{n+1} = T_n + h(-k(T_{n+1} - T_a))$

Since we know the starting temperature of the soup $(T_0 = 100)$, we can simply iterate towards the end value with the forward Euler method. Lets say h = 20 and $k = 9.796 \times 10^{-4}$ as we worked out earlier:

¹The change in temperature over time is equal to the temperature minus the ambient temperature multiplied by the cooling coefficient.

$$T_0 = 100$$

 $T_1 = 100 + 20(-9.796 \times 10^{-4}(100 - 10)) = 98.237$
 $T_2 = 98.237 + 20(-9.796 \times 10^{-4}(98.237 - 10)) = 96.508$
 \vdots
 $T_{111} = \dots = 20.009$
 $T_{112} = \dots = 19.813$

Since we did 112 iterations before we dropped below the target temperature of 20, and h = 20, we know the time is between 2220 and 2240 seconds. Since we previously calculated the actual time it would take (2242 seconds), we're pretty close to the right answer. We probably got a bit of inaccuracy because the forwards Euler method is unstable (we'll get to that later). If we decreased h, then we would get a better answer (in fact, when h = 1, we get 2242 seconds).

We can also use the backward Euler method. To do this we need to rearrange the method to give T_{n+1} so that it is not a recursive definition:

$$T_{n+1} = T_n + h(-k(T_{n+1} - T_a))$$

$$= T_n - hk(T_{n+1} - T_a)$$

$$\frac{T_{n+1}}{-hk} = \frac{T_n}{-hk} + T_{n+1} - T_a$$

$$\frac{T_{n+1}}{-hk} - T_{n+1} = \frac{T_n}{-hk} - T_a$$

$$T_{n+1} + khT_{n+1} = T_n + hkT_a$$

$$T_{n+1}(1 + hk) = T_n + hkT_a$$

$$T_{n+1} = \frac{1}{1 + hk}(T_n + hkT_a)$$

So lets apply the backwards Euler method with h = 20:

$$T_0 = 100$$

$$T_1 = \frac{1}{1 + 20 \times 9.796 \times 10^{-4}} (100 + 20 \times 9.796 \times 10^{-4} \times 10) = 98.271$$

$$T_2 = \frac{1}{1 + 20 \times 9.796 \times 10^{-4}} (98.271 + 20 \times 9.796 \times 10^{-4} \times 10) = 96.574$$

$$\vdots$$

$$T_{112} = 20.047$$

$$T_{113} = 19.854$$

Therefore we can see that the backwards Euler method says that the temperature reaches 20 degrees between 2240 and 2260 seconds (112 and 113 times h). If we decreased the step size h, we could get a more accurate calculation (when h=1 we get 2244-2245 seconds). The benefit of the backwards method is that it is unconditionally stable (i.e. errors in the input between successive iterations will diminish as the algorithm executes)².

1.1.1 Consistency of Linear Multistep Methods

We have seen how to construct linear multistep methods, and had a look at how to make them stable. Now we must examine their consistency, which is the property that as h decreases, the

²http://mathworld.wolfram.com/NumericalStability.html

truncation error can be made to approach zero; i,e, as h approaches $\frac{1}{\infty}$, we approach the true solution of the equation.

To find if a method is consistent, we must first find its order. This is done by defining the constants C_0, \ldots, C_j :

$$C_{0} = \alpha_{0} + \dots + \alpha_{k}$$

$$C_{1} = \alpha_{1} + 2\alpha_{2} + \dots + k\alpha_{k} - (\beta_{0} + \dots + \beta_{k})$$

$$C_{j} = \frac{1}{j!}(\alpha_{1} + 2^{j}\alpha_{2} + \dots + k^{j}\alpha_{k}) - \frac{1}{(j-1)!}(\beta_{1} + 2^{j-1}\beta_{2} + \dots + k^{j-1}\beta_{k})$$

A linear multistep method is of order p if $C_{p+1} \neq 0$. In other words $C_0 = C_1 = \cdots = C_p = 0, C_{p+1} \neq 0$. If the method is of order $p \geq 1$ then it is consistent. This can also be expressed as the sum of the alphas being 0, or the sum of the product of the index and the alpha being equal to the sum of the betas:

$$\sum_{i=0}^{k} \alpha_i = 0, \sum_{i=0}^{k} i\alpha_i = \sum_{i=0}^{k} \beta_i$$

1.1.2 Stability of Linear Multistep Methods

A method is stable if its first characteristic polynomial does not have zeros with a modulus greater than one, and if all the zeros with a modulus equal to one are simple. This sounded very abstract to me, but after talking to Milan about it, I'll give my best shot at what it means:

We must define the first characteristic polynomial to be:

$$p(\zeta) = \alpha_k \zeta^k + \alpha_{k-1} \zeta^{k-1} + \dots + \alpha_0 \zeta^0$$

= $\alpha_k \zeta^k + \alpha_{k-1} \zeta^{k-1} + \dots + \alpha_1 \zeta + \alpha_0$

Now, we need to find the roots of this polynomial (i.e. where $p(\zeta) = 0$), which will give us a set of values ζ_1, \ldots, ζ_n (usually there are only a few values). Some of these values can be complex numbers (e.g. if $\zeta = a + ib$ is a solution, then $\zeta = a - ib$ is too). To calculate the modulus of one of these, you get the actual value if the number is real, or for complex numbers, you can do $|\zeta_n| = \sqrt{a^2 + b^2}$.

1.1.3 Optimal Methods

An optimal method is one that is both stable, and has an order of k+2 (where k is the number of terms it has, e.g. the Euler method has k=1, therefore the Euler method is not optimal).

The First Dahlquist Barrier states that a method cannot attain an order of convergence greater than k + 1 is k is odd or k + 2 if k is even.

1.1.4 Examples of Analysing Linear Multistep Methods

Example One:

Lets examine the Euler method first, defined as $y_{n+1} = y_n + hf_n$. We first need to get it into the form where all the y's are on one side, and all the f's are on the other side (this is easy):

$$y_{n+1} - y_n = hf_n$$

Now we can simply read off the values of α and β :

$$\alpha_0 = -1$$

$$\alpha_1 = 1$$

$$\beta_0 = 1$$

$$\beta_1 = 0$$

Using these, we can calculate the values of the constants:

$$C_0 = \alpha_0 + \alpha_1 = -1 + 1 = 0$$

$$C_1 = \alpha_1 - \beta_0 - \beta_1 = 1 - 1 - 0 = 0$$

$$C_2 = \frac{1}{2}\alpha_1 - \beta_1 = \frac{1}{2} - 0 = \frac{1}{2} \neq 0$$

Since C_2 is not 0, then the method is consistent and of the first order. To calculate the stability, then we must find the first characteristic polynomial:

$$p(\zeta) = \alpha_1 \zeta^1 - (\alpha_0 \zeta^0) = \zeta - 1$$

Now we can solve for ζ :

$$p(\zeta) = 0$$
$$\zeta - 1 = 0$$
$$\zeta = 1$$

Since $\zeta = 1$ (which satisfies the conditions we said before), we know that it is stable. Since it is stable and it is consistent, then it is convergent.

Example Two:

For the next example, lets look at the equation:

$$y_{n+2} - y_{n+1} = \frac{h}{3}(3f_{n+1} - 2f_n)$$

Lets see if it is consistent. To do this we must compute the constants:

$$C_0 = 0 - 1 + 1 = 0$$

$$C_1 = -1 + 2(2) - (0 + \frac{3}{3} - 23)$$

$$= 3 - \frac{1}{3}$$

$$= 2\frac{2}{3}$$

Since $C_1 \neq 0$, the method is not consistent (and therefore cannot be convergent either). Next we can see if it is stable, but getting the first characteristic polynomial:

$$p(\zeta) = \alpha_2 \zeta^2 + \alpha_1 \zeta + \alpha_0$$
$$= \zeta^2 - \zeta$$
$$p(\zeta) = 0$$
$$\zeta = 0, \zeta = 1$$

Therefore we can see that the method is stable (since the roots are between 0 and 1).

Example Three:

We have:

$$y_{n+2} + 4y_{n+1} - 5y_n = \frac{2}{h}(8f_{n+1} + 4f_n)$$

The consistency would be:

$$C_0 = 1 + 4 - 5 + 0 = 0$$

$$C_1 = 2(1) + 4 - (4 + 2) = 0$$

$$C_2 = \frac{1}{2!}(-5 + 2^2(4) + 3^2(1)) - \frac{1}{1!}(2 + 2^1(4))$$

$$= \frac{1}{2}(4(1) + 4) - 4$$

$$= 0$$

$$C_3 = \frac{1}{3!}(8(1) + 4) - \frac{1}{2!}(4)$$

$$= \frac{1}{6}(12) - 2$$

$$= 0$$

$$C_4 = \frac{1}{4!}(16(1) + 4) - \frac{1}{3!}(4)$$

$$= \frac{1}{24}(20) - 1.5$$

$$= -\frac{2}{3}$$

Since it is of order 3, the method is consistent.

To find the stability, we must find the first characteristic polynomial:

$$p(\zeta) = \zeta^{2} + 4\zeta - 5$$
$$p(\zeta) = 0$$
$$\zeta = 5, \zeta = 1$$

Since $\zeta = 5$, the method is not stable, and is therefore not convergent.

Example Four:

Sometimes, Milan will ask for the roots of a complicated polynomial like he did in 2014. Here is the question:

c) Consider the following linear multistep method:

$$\frac{1}{3}y_{n+3} + \frac{1}{2}y_{n+2} - y_{n+1} + \frac{1}{6}y_n = hf_{n+2}, \quad n = 0, 1, \dots,$$
 (2)

where h is the discrete step size and f is the right hand side of (1).

i) Determine the order of method (2). Is the method consistent? Write the first characteristic polynomial for method (2) and, based on its zeros, discuss the stability of the method. (6 marks)

Here is my answer:

In order to determine the order of method two, we first must find the values of α and β :

$$\alpha_0 = 0, \alpha_1 = \frac{1}{6}, \alpha_2 = -1$$

$$\alpha_3 = \frac{1}{2}, \alpha_4 = \frac{1}{3}$$

$$\beta_0 = 0, \beta_1 = 0, \beta_2 = 1$$

And now using those, we can find the constants:

$$C_0 = \sum_{i=0}^k \alpha_i = \frac{1}{6} - 1 + \frac{1}{2} + \frac{1}{3} = 0$$

$$C_1 = \sum_{i=1}^k i\alpha_i - \sum_{i=0}^k \beta_i = \frac{1}{6} - 2(-1) + \frac{3}{2} + \frac{3}{3} - 1 = -\frac{1}{3}$$

Therefore we can see that the method is not consistent as it is of order 0.

The first characteristic polynomial is:

$$p(\zeta) = \frac{1}{3}\zeta^3 + \frac{1}{2}\zeta^2 - \zeta + \frac{1}{6}$$

Since we can (relatively easily) see that $\zeta = 1$ is a root, we can use polynomial division to get the rest:

$$\begin{array}{r}
\frac{\frac{1}{3}x^2 + \frac{5}{6}x - \frac{1}{6}}{\frac{1}{3}x^3 + \frac{1}{2}x^2 - x + \frac{1}{6}} \\
-\frac{\frac{1}{3}x^3 + \frac{1}{3}x^2}{\frac{5}{6}x^2 - x} \\
-\frac{\frac{5}{6}x^2 + \frac{5}{6}x}{-\frac{1}{6}x + \frac{1}{6}} \\
-\frac{\frac{1}{6}x - \frac{1}{6}}{0}
\end{array}$$

Now we have $(\zeta - 1)\left(\frac{\zeta^2}{3} + \frac{5\zeta}{6} - \frac{1}{6}\right)$, we can use the quadratic formula to find the rest of the roots:

9

$$a = \frac{1}{3}, b = \frac{5}{6}, c = \frac{-1}{6}$$

$$\zeta = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a} = \frac{15 \pm \sqrt{209}}{12}$$

$$\zeta = 2.556, \zeta = 0.0453$$

Since 2.556 is greater than 1, the method is not stable.

2 Complex Networks

Eva's notes and lectures are really very good, use her notes, not mine:)

3 Optimisation and nature inspired algorithms: DEPRE-CATED

This section is no longer examinable as of the 2017-2018 syllabus.

To start, lets recap some stuff you should already know; a minima is a point of a function where all points in its vicinity have a higher value than itself, and a maxima is the opposite; a point where all points in its vicinity have a lower value than itself.

Since we can invert a function by putting a minus in front of it, we don't really need to differentiate between maxima and minima, since we can always express one in terms of the other. We can however differentiate between *global optimums*, which are the best values for the entire domain, and *local optimums* are best in some bounded region.

One dimensional functions are easiest to visualise; they are just a graph, where the y value is the value of the function and the x value is the input. As the dimensionality increases, the functions get harder to visualise; 2d functions are visualised on a 2d graph, where the intensity of the colour in each square represents the value of the function at that coordinate.

An optimisation problem is one where we attempt to maximise or minimise a function. This involves finding the input that will produce the largest or smallest value. Optimisation problems are really search problems; given a domain, find an input that produces the most desirable output.

If we know exactly what the function we're trying to optimise is (i.e. we have an equation), then it's an explicit problem, but if we just have some inputs to an unknown function, and their corresponding output values, then it's a black box problem.

Similarly, there are two different approaches to solving these problems:

Single solution:

This is where there is a single candidate solution that is incrementally improved by the algorithm throughout the procedure.

Population based solution:

This is where there is a set of candidate solutions (the population) and an iterative operation combines and mutates the best ones, or increments the whole population to improve the quality of the population.

All of the optimisation methods follow a cycle:

- Guess some parameters for the initial solution
- While we're not satisfied:
 - Evaluate how the current parameters perform

- If we're satisfied, then we've finished
- Otherwise, determine new parameters based on the current ones and their evaluation.

We can use derivatives to try and find maxima and minima in a function. If we do not have an explicit function for the derivative, then we can calculate them using **finite differences**, of which two approaches are presented here:

Forward difference:

$$f'(x) \approx \frac{f(x+h) - f(x)}{h}$$

Central difference:

$$f'(x) \approx \frac{f(x+h) - f(x-h)}{2h}$$

Here, the h parameter is the range over which we're calculating the differential.

3.1 Minimising univariate functions

For one dimensional functions, we can use two approaches:

- Interval reduction, where we iteratively reduce the range of values that we think the optimum is inside.
- Interpolation, where we try to find an approximate function and use the optimum of that function to iterate to find a better approximate function.

3.1.1 Bisection algorithm

This is the easiest algorithm, we start with the full range in the domain, and then cut it in half based on the value of the differential at the midpoint at the range.

```
bisection(f'(x), a, b, e) {
  while (b - a >= e) {
    c = (a + b) / 2;
    if (f'(c) < 0) a = c;
    else if(f'(c) > 0) b = c
    else return c;
}
```

3.1.2 Quadratic interpolation algorithm

Here, we make a quadratic function that goes through three points a, b and c, then we either drop a or b depending on which way we should move the quadratic function. Here, we do not need the differential which is good since we won't always have it!

Please look at the lecture notes for psudo code here.

3.1.3 Stopping criteria

Eventually, we will have to stop our optimisation algorithms, but some of them will run indefinitely. We need criteria to make them stop after a sane amount of time. The simple approach is to stop after n iterations, and is applicable to all iterative functions. The more clever approach is to only stop when we reach a certain threshold, i.e. when a method converges asymptotically on an optimum.

3.2 Minimisation of multivariate functions

There are two different classes of minimisation methods for multivariate functions:

Based on derivatives:

Moves are determined based on information from derivatives of the multivariate function.

A directional derivative is a derivative indicating the rate of change in a specific dimension:

$$\Phi_r' = \lim_{h \to 0} \frac{f(x+hr) - f(x)}{h}$$

Where r is the direction. If Φ'_r is positive, then the direction is ascending, if it's negative, then the direction is descending and if it's equal to zero, then there is no slope.

Finding the gradient of a vector of a multidimensional function f(x) (and hence the vector of all partial derivatives) is:

$$\frac{\delta f}{\delta x_i} = \lim_{h \to 0} \frac{f(x + he_j) - f(x)}{h}$$

 e_j is the unit vector in the direction of axis j, so to construct the whole gradient for the vector, we need to do:

$$\Delta f(x) = \begin{bmatrix} \frac{\delta f}{\delta x_1}(x) & \frac{\delta f}{\delta x_2}(x) & \dots, \frac{\delta f}{\delta x_n}(x) \end{bmatrix}^T$$

The **Hessian** is the matrix containing the second order partial derivatives, and is always of size $N \times N$:

$$Hf(x) = \begin{bmatrix} \frac{\delta^2 f}{\delta x_1 \delta x_1}(x) & \dots & \frac{\delta^2 f}{\delta x_1 \delta x_n}(x) \\ \vdots & \ddots & \vdots \\ \frac{\delta^2 f}{\delta x_n \delta x_1}(x) & \dots & \frac{\delta^2 f}{\delta x_n \delta x_n}(x) \end{bmatrix}$$

Most algorithms have some kind of parameter update scheme, where they will move in a given direction r for a length of α every iteration, and r, α change based on the current position:

$$x_{i+1} = x_i + r_i \alpha_i$$

3.2.1 The alternating variable method

A conceptually easy method, you start from an arbitrary position x, and then for each dimension d:

- Find the differential for that dimension
- Find the minimum according to that differential
- Move the value of x_d to be equal to that minimum.

Then repeat for all dimensions until no progress is made.

3.2.2Newton's method

As soon as Newton invented calculus, he also started inventing methods for finding minima. This method requires a function to be twice differentiable:

$$x_{i+1} = x_i - \alpha \frac{\Delta f(x_i)}{Hf(x_i)}$$

The trouble with Newton's method is that it will only converge if the initial point is close to the solution, and the memory requirements for the Hessian matrix is n^2 .

Direct search:

Moves are determined using methods other than derivatives, for example, using geometric concepts.

3.3 Simplex methods

If you did Advanced Algorithms 1, you're either groaning now or you're jumping for joy at being able to skip a section. However, these aren't the simplex methods you learned about last semester.

For this, you construct a simplex (a shape of n+1 vertices in n dimensions), and then observe that you can always form a new simplex from an existing one by adding a new point.

The Spendley, Hext and Himsworth method is like so:

- 1. Create n+1 vectors for a regular simplex (where all sides are the same length).
- 2. For each vector, the value of the objected point is calculated for that point.
- 3. The vector with the highest point is removed and substituted by a new one.
- 4. If the worst vector is also the most recently introduced one, then the next worst one is used.
- 5. The newly substituted vector is the mirror image of the old one along the axis of the remaining vectors.
- 6. When the simplex simply rotates around a point, then the vertex in the middle is the minimum.

The maximum age of a vector is $1.65n + 0.05n^2$, and if a vertex exceeds age a, then the search stops, or is restarted using a smaller simplex. If the search was restarted, then the initial (smaller) simplex is started from the site of the rotating.

The Nelder and Mead method is similar, though the simplices are no longer regular (though they can be). It involves having rules that either expand, contract or reflect the simplex (or any combination of them), which leads to quicker results and better approximations of the solution.

Where $y_0^{(r)}$ is the best value in the previous simplex, and $y_{n-1}^{(r)}$ is the worst, the new simplex results from:

$$y_0^{(r)} < y_i^{(r+1)} < y_{n-1}^{(r)} \colon$$

 $\begin{aligned} y_0^{(r)} &< y_i^{(r+1)} < y_{n-1}^{(r)} \\ &\text{The simplex is reflected like in the SHH method.} \end{aligned}$

$$y_i^{(r+1)} < y_0^{(r)}$$
:

The simplex is reflected and expanded.

$$y_i^{(r+1)} > y_{n-1}^{(r)}$$

 $\begin{aligned} y_i^{(r+1)} &> y_{n-1}^{(r)} \\ &\text{The simplex is reflected and contracted.} \end{aligned}$

3.4 Optimisation and nature inspired algorithms

As we have seen, a global optimum is the best set of admissible conditions to achieve an objective (i.e. minimising the function) under a set of constraints.

Finding such an optimum is non-trivial, mainly because the search space is often continuous, so there are an infinite number of parameters to test. The following algorithms try to find global optimums:

Grid Search:

We overlay a grid onto the search space (in as many dimensions as we need to), and test each point on the grid. We can change the resolution of the grid by making the space between the grid lines larger or smaller. The algorithm scales exponentially in the number of dimensions, but linearly in the number of points tested (dependent on the grid resolution), therefore the runtime is $O(n^d)$.

The quality of the solution depends on the density of the grid, but getting a high quality solution is usually impractical, since the search space grows so quickly (even in two dimensions, it is quadratic).

Random search:

Here, we simply generate n random points and sample the objective function at these points. The point that gives the best value for the objective function is remembered for possible later use (since that's the optimum). Obviously this algorithm runs in O(n) time, and its quality depends on the number of random points chosen. It is possible that the random search method could outperform all the other methods if it got lucky, though this is unlikely.

Due to its ease of implementation, random search is often used as a reference for other methods.

Multistart:

Here, we guess n random points again, but for each one, we find the local minimum. This improves the quality (since there's more chance of finding the global minimum, since it's definitely a local minimum too), but it can degrade performance.

This method often visits the same local minima multiple times (if optimisations aren't made to avoid doing this) so it can be inefficient.

Random walk:

We can generate a single random start point, but then generate subsequent points by choosing a random direction and length for a vector and adding that to the current point.

Using nature to help minimise:

If a crystal is being formed, and it cools slowly, then the atoms in the crystal will vibrate around until they are in a local minima, which results in some cool shapes. We can use techniques to emulate this to find minima in our functions.

Furthermore in genetics, non-optimal genes are 'selected out' of the population by natural selection and evolution, leading to a genotype that is optimum for the environment.

The remaining algorithms use nature as inspiration:

Metropolis algorithm:

This is an algorithm that emulates an ensemble of particles in equilibrium at a certain temperature. It uses the Boltzmann probability density function:

$$p.d.f = e^{\frac{-E}{k_B T}}$$

To give the probability that a certain particle configuration with an energy E has a certain temperature T.

In nature, perfect crystals are formed by the cooling down from being molten very slowly so that the material can reach an equilibrium at each temperature. If the cooling is too fast, then an equilibrium will not be reached at each temperature, and the crystal will have imperfections.

Local optimisation algorithms have parallels with letting crystals cool too fast!

The algorithm is as follows:

- Start with a random position for an atom x^0
- Create a small random displacement to obtain x^1 and calculate the difference in energy $\Delta E = E^1 E^0$.
- If $\Delta E<0$ accept the new position, otherwise accept it with a probability of $P(\Delta E)=e^{\frac{-\Delta E}{k_BT}}$
- Iterate a lot of times, simulating the thermal motion of particles in a heat bath of temperature T.

The Metropolis algorithm works because it lets the energy of the particle increase even though that's a 'step back' in terms of optimisation. Sometimes the algorithm needs to get out of a local minima in order to find the global optimum.

Simulated Annealing:

This takes advantage of the fact that we can see the energy of a particle was similar to the value of an objective function we're trying to maximise, and the coordinates that it is at as similar to the parameters of the function we're finding an optimum for.

The only other variable is the temperature, which acts as a control parameter with the same units as the objective function. Simulated annealing starts by melting the objective function at a high temperature, then using the Metropolis algorithm to calculate the equilibrium of the objective function at temperature decreases. Here's the algorithm:

- Start with a high temperature T^0 at some random position $x^{(0,T^0)}$.
- Apply the Metropolis algorithm to determine the average equilibrium value of the objective function and parameter values $x^{(Ex,T^0)}$.
- \bullet keep reducing T and repeating the previous step, and only stop when the function freezes; you've reached the global minimum.

Note that the start position is irrelevant, since there are many iterations required for each temperature, and an equilibrium is found for each. The start temperature, T^0 is very important though; if its too low, then a local minimum will be found, and if its too high, then the algorithm will take too long.

Simulated annealing can be reformulated computationally as a Markov Chain, and a proof has been given that states the algorithm will converge to a global optimum in infinite time, which makes simulated annealing *asymptotically complete*.

Random and grid searches are also asymptotically complete, but they converge at much

Press' modification:

In 1989, Press et al. suggested that each temperature cycle should last for a predetermined number of iterations N. After each cycle, the temperature should be reduced by a constant factor p:

$$T^{n+1} = pT^n \quad < 0 \le p \le 1$$

If after a cycle has finished, there have been no successful moves, then the algorithm stops. If we increase N, then the accuracy increases, but the execution time increases faster too. Similarly if we increase p, the solution will improve (and the reliability too), but the rate of cooling drops so it takes longer.

There is also a slight modification to the displacement step; a single parameter is changed to a random value within the boundaries of the parameter.

Corana's modification:

Corana et al. suggested that the displacement step should be based off a vector; each parameter is changed according to the same coordinate of a vector v_i . After N_s rounds, v_i is changed so half the new candidate parameters are changed. Just like Press' modification, after N_r rounds, the temperature is reduced.

As a result, the annealing will adapt to the shape of the function, but it is also slower; N in Press' algorithm is equal to $N_s \times N_r$ in Corana's algorithm.

3.5 Evolutionary algorithms

Natural populations evolve by having variation amongst the members of the population and having selection (or unfit members of the population dying before they get chance to reproduce). Evolutionary algorithms are a class of optimisation algorithms that evolve candidate solutions as a group (ensemble) rather than looking at one solution at a time.

If you took a biology course at some point, then you may be able to figure this out, but there is a mapping from biology terminology to CS terminology:

- Gene Parameter
- Chromosome All parameters
- Individual Candidate solution
- Generation One iteration
- Fitness Value of the objective function

Evolutionary algorithms evolve their populations. This means there will be n points in the parameter space, and each iteration will drop some of the points and create new ones (in hopefully better locations). Each individual in the population is a single candidate solution, and is stored as a *chromosome*. This is a sequence of bits, split into m sections called *genes*. As described above, each gene maps onto a parameter.

Chromosome:

1100110 1010101 1010111 0101101 1011010 Genes (each one is a parameter) 102 85 87 45 90 Alternately, floating point parameters 4.48 2.20 8.80 3.0 1.11 4.12

Fogel's algorithm for evolutionary programming is as such:

- 1. Generate a random initial population of n individuals
- 2. Calculate the fitness of each individual in the population (using the objective function)
- 3. Each individual from the current population generates and offspring by copying its own genes
- 4. Mutate each gene for each chromosome in the offspring by a small variance.
- 5. Put the offspring in a new population, 2n large
- 6. Probabilistically select n individuals as a function of fitness to be removed (now we have a population of n again).
- 7. Go back to step 2, or stop.

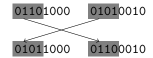
In this algorithm, we described mutation (in step 4) and selection (in step 6). For mutation, we add a small random number to the original value of a gene.

For selection, we want to select n individuals, where better individuals have a greater chance of remaining in the population. A roulette wheel is a manual way of doing this (each individual has a probability of being picked according to their fitness), but stochastic ranking (e.g. a sort that has a chance of being wrong) or a tournament where each individual is compared against a small number of others and receives a score based off of its performance; the n lowest scoring individuals are pruned.

Holland and DeJong's Genetic algorithm is as follows. It encodes parameters in binary, and genes are therefore strings of binary digits.

- 1. Generate n random individuals for the population
- 2. Calculate the fitness of each individual in the population
- 3. Choose two parent individuals from the current population as a probability of their fitness
- 4. Cross them over (see below) at a random locus to produce two offspring
- 5. Mutate each locus (gene) in the offspring with a small probability
- 6. Put the offspring in the new population
- 7. Go to step two, or stop.

The mutation operation flips bits in a gene with a small probability. The cross over operator swaps the latter halves of two chromosomes at a random index.



The differences in the two algorithms are as follows:

Evolutionary programming	Genetic algorithms
Genes are encoded as floating point numbers	Genes are encoded in binary.
Mutation is addition of a random value	Mutation is bit flipping.
Asexual reproduction (two parent	Sexual reproduction (two parents
makes one child)	make two children.)
No cross over	Cross over by swapping subsections of the chromosome.

Rechenberger and Schwefel made another algorithm. In this algorithm, each gene has an associated standard deviation which affects how it mutates. This is an innovation because in the standard approach, the size of each mutation is the same for all genes and is constant throughout the execution of the algorithm. Here, the standard deviation for each gene is updated throughout the algorithm, and henceforth mutations are different sizes for each gene and changes as the algorithm progresses:

- 1. Generate n random individuals
- 2. Calculate the fitness of each
- 3. Randomly pair up individuals
- 4. Recombine them to create two offspring
- 5. Mutate each gene locus with a small probability based of the gene's standard deviation.
- 6. Put the offspring in the new population (2n large)
- 7. Select the best n
- 8. Go back to step 2, or stop.

In this algorithm, the mutation happens in two stages: first the standard deviation is mutated first, and then the gene is added to by a random number based off the standard deviation.

The recombination operator is also slightly different; since each gene in the child is the average of the parent gene:

$$(12, 5, 34, 2), (10, 7, 30, 4) \rightarrow (11, 6, 32, 3)$$

It is important to note the differences between selection and variation so we can get the balance right:

- Selection is responsible for keeping improvements in the population and not discarding them
- Very strong selection limits the variation and all individuals will be the same
- Mutation and cross over are responsible for introducing variation and will make the parameters/gene pool change
- Very strong variation results in good solutions, but they will be replaced by random ones and not retained.

Many genetic algorithms progress at very slow rates, but then have short bursts of very rapid progress. In general, it is not common for these algorithms to have proven convergence properties, but some of them do when applied to simple problems.

Deciding when to stop a genetic algorithm isn't always simple; stopping after a predetermined number of generations requires lots of trial and error to get the number right, waiting until a desired fitness is reached is okay if you know what the optimal fitness is, but it may take an (infinitely) long amount of time, and waiting for the fitness variations from one population to the next to become stable isn't a good idea either because of the propensity for genetic algorithms to progress very slowly then have a burst of progress.

3.6 Genetic Programming

With evolutionary programming, we optimised the parameters of the input function in order to find the ones that produce the best solution. On the other hand, genetic programming optimises the function until it fits our purpose. This may seem like a strange approach, but when you consider that any computer program is just a big mathematical function, then genetic programming just becomes automatic computer programming.

In this sense, programs evolve to become better at solving the problem they are being evaluated against. If you think about it, programmers have two 'phrases' when to producing code; developing new code, and editing existing code. This maps cleanly onto genetic programming, since the algorithms first generate entirely new pseudo-random code, and then mutate/cross it over with other code.

We can represent programs as trees, where internal nodes are functions (which includes arithmetic/logical operations) with arguments, and where leaf nodes correspond to constant values, functions with no arguments, or inputs to the program.

The genetic algorithm must be given a set of functions and a set of terminals that it can use to generate the tree. Example functions include:

- Boolean functions (AND, OR,...)
- Arithmetic functions (+,%,...)
- Transcendental³ functions (sin, cos, log,...)
- Conditional statements (if, switch, etc)
- Loops (while, for, etc)
- Variable manipulation (set, etc)

³A transcendental function is one that cannot be described with a finite sequence of algebraic operations; i.e. it transcends algebra.

• Subroutines⁴ (http-request, read-sensor, etc)

The set of terminal nodes is smaller, and can include constants⁵, functions such as rand(), variables etc.

In order to create valid programs, the genetic algorithm should implement some type checking to ensure that only valid inputs are given to functions. This may include automatic type conversions to increase the number of valid trees that can be generated.

The algorithm must also attempt to stop things like RuntimeException occuring; i.e. the program must produce some usable output. This can be done by having functions return a default value if they fail with invalid inputs⁶, or by making functions that fail score very low when they are evaluated (so that they are selected out of the population).

3.7 Cramer & Koza's Genetic Programming

As previously stated, the program is encoded as a tree, using functions and terminals. Cramer and Koza's algorithm is as follows:

- 1. Initialise a random population of n individuals
- 2. Calculate the fitness of each individual
- 3. Select two parents at random proportionally to their fitness and reproduce them together, producing two children
- 4. Apply cross over with a certain probability
- 5. Apply mutation with a certain probability
- 6. Remove the parents
- 7. Loop back to step 2 unless satisfied.

While the algorithm may seem well specified, there are a few additional things we need to clear up; namely initialisation and mutation/cross over.

Initialisation is done using one of two methods:

Full method:

Choose a function node at random, and either fill its children with terminal nodes if we have reached the required depth, or loop and choose more function nodes.

Grow method:

Choose a function or a terminal node at random, and fill its children with child nodes if the maximum depth has been reached, or loop back and choose child or function nodes.

Both of these methods generate trees of up to a specified depth, though the full method always generates a tree where all branches are that depth, while the grow method generates branches up to that depth.

Mutation is applied to single individuals; a node is chosen at random and is substituted with a randomly generated sub-tree. Different implementations generate the sub-tree differently, some will maintain the depth of the tree to avoid it growing too big or generate a tree that can be used as input for the parent node (to avoid creating an invalid tree post-mutation), while others have no such constraints.

 $^{^4}$ Think of these like library calls. The evolutionary algorithm is really just trying to string these together.

⁵Random constants are determined at the start of the program (i.e. once per run).

⁶Type checking can't catch everything (though some functional languages have type systems that integrate proof systems and can check all kinds of funky stuff; Google for 'dependent types'), divide by zero is an example of this.

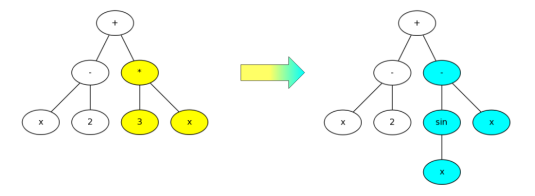


Figure 1: Mutation of tree.

Cross over is when children inherit information from both their parents. For our purposes, this involves making a tree with a sub-tree from one parent and a sub-tree from another parent. The node within each parent to cross over at is chosen at random (meaning that there could be a case when most of the tree is inherited from one parent and little of it is inherited from another.).

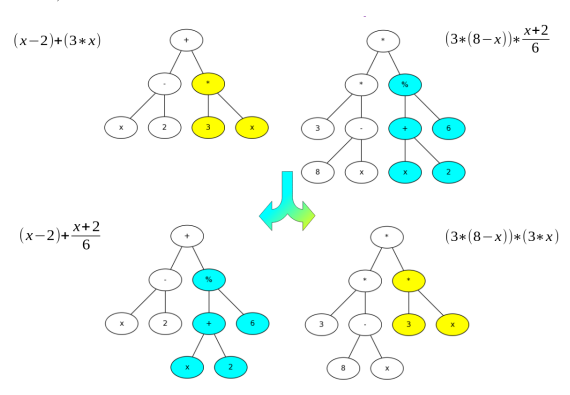


Figure 2: Cross over between two trees.

3.8 Bloat

Bloat is when programs created by genetic algorithms get larger and larger trees without significant improvement to the fitness of the program for the function it is trying to perform. This is undesirable, since more memory is used, the expressions are evaluated slower and the population evolves more slowly.

Bloat is caused by a variety of sources, such as when cross over is very unbalanced, and through random mutations. Bloat often 'survives' and accumulates because it is neutral to fitness; many bloated sub-trees will have no effect on the fitness (these are celled introns).

Examples of introns include expressions like x - x, x && true etc (basically the identity functions).

Controlling bloat is hard; but we can do so in a few different ways:

- Enforce tree size and depth limits, maybe by returning a parent of a tree if its child violates a limit.
- Anti bloat operations could be implemented such as making sure that the second subtree in cross over is small enough to not violate size constraints, or by making sure that a mutated tree is of the same size as the original one was.
- Selection can be used to reduce bloat; the fitness of trees can be reduced proportional to their size (parsimony pressure), or the fitness of trees violating the size constraints could simply be set to zero (the tarpeian method).

All evolutionary algorithms have a requirement for diversity and variability in their population in order to evolve. When all individuals in a population are similar, cross-over won't effect much change and mutation is unlikely to move individuals far in the 'solution space'. Unfortunately, due to selection, populations tend to evolve to be relatively uniform.

There are attempts to make evolutionary algorithms more similar to nature in the sense that there can be multiple populations of the same species in different areas with little mixing between them. Algorithms mimicking this effect are called *Distributed EA's*, and evolve multiple populations in near-isolation (some mixing will occur) in order to try and get heterogeneous sets of effective genes. Some swarm algorithms are an example of this (multiple swarms can be present in the solution space).