

# Optimisation Methods Notes

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# 1 What is Optimisation?

## 1.1 Combinatorial Optimisation

Most problems can be represented as  $(R, C)$  where:

**R** is a **finite** set of configurations, each of which is a possible solution to the problem. They effectively represent the search space of the problem, and in most cases won't be explicit. e.g.

$S = 1, 2, 3, \dots, 100$  and  $S = \{x \mid x \in \mathbb{N}, x \leq 100\}$  are the same space, but the second is much more compact

**C** is the cost function, which tells us the value of each configuration. It takes in a configuration and outputs a real number.

**An optimisation problem is to find the configuration with the lowest cost** (i.e.  $\arg \min(r)$ ). If we wanted to find the configuration with the highest cost, we could just multiply the costs by -1.

### 1.1.1 SAT Problem

Given a boolean CNF (a boolean consisting of 'AND' with 'OR' as sub-problems (e.g.  $(a \text{ OR } b) \text{ AND } (c \text{ OR } d)$ )), find a configuration of truth values for each variable that satisfies the CNF (i.e. the whole CNF results in true).

The cost function is simply 1 if the CNF is true, and 0 otherwise. X-SAT refers to a problem where the longest sub-problem's length is X. Any  $X > 2$  can be converted to a 3-SAT by adding extra variables. 2-SAT is P complexity, 3-SAT is NP-Complete (Explained in 1.2.2).

## 1.2 Time Complexity

### 1.2.1 Algorithms

Measuring the running time of algorithms is futile: it depends on the hardware used, the background processes running on the computer, etc. Instead it's assumed that each step takes a constant amount of time, and the growth in the number of steps based on the size of the input determines the time complexity of an algorithm.

Again, rather than comparing the exact number of steps, an algorithm is abstracted to the largest influence on the number of steps. e.g. If the algorithm takes  $4n^2 + 2n + 3$  steps ( $n$  is the input size), we represent only consider the  $n^2$ . This is because, for large enough input values, the lower powers have little impact on the final value. e.g. The difference between  $n^2$  and  $n^2 + n$  at  $n = 1000$  is 0.1%.

To represent the time complexity of an algorithm ( $f(x)$ , the growth in the number of steps with the growth in the input size), we use the following three notations:

**Big-O(O)** An algorithm is  $O(g(x))$  if there are some constants  $c$  and  $d$  such that  $c * g(x) \geq f(x)$  when  $x > d$ . This gives an upper bound on the performance of an algorithm, so the **Worst Case**, and is the notation mostly used.

**Big-Omega( $\Omega$ )** An algorithm is  $\Omega(g(x))$  if there are some constants  $c$  and  $d$  such that  $c * g(x) \leq f(x)$  when  $x > d$ . This gives an lower bound on the performance, so the **Best Case**.

**Big-Theta( $\Theta$ )** An algorithm is  $\Theta(g(x))$  if there are some constants  $c_1, c_2$  and  $d$  such that  $c_1 * g(x) \leq f(x) \leq c_2 * g(x)$  when  $x > d$ . This gives a tight bound on the performance, so the **Average Case**.

$O$  and  $\Omega$  could be very loose but still technically correct. e.g  $f(x) = n$  is  $O(n^{10})$ , but that doesn't tell us anything useful. Since we mostly expect large input values, we use  $d$  to remove problems at smaller values: e.g.  $5n > n^2$  for  $n < 5$ , but  $n^2$  grows quicker,

$$O(\log n) < O(n)(Linear) < O(n * \log n) < O(n^x)(Polynomial) < O(X^n)(Exponential) \\ O(n!) \approx O(n^5).$$

### 1.2.2 Problems

**P** denotes the class of problems for which there exists a **Polynomial-Time** algorithm to solve them. These are computationally 'easy'. e.g. Shortest Path with Positive Weights, Linear Programming

**NP** problems can't be solved, but a solution can be verified, in polynomial time. Alternatively, they can be solved in polynomial time by a **Non-deterministic Turing Machine** (a type of automaton with memory). Finding a polynomial solution to one of these is an ongoing problem. e.g. Travelling Salesman Problem, Boolean SAT

Chess is NP-Hard (as hard/harder than every other NP problem), but is EXP-Complete (an exponential-time algorithm), since verification is exponential. A problem is NP-Complete if it is harder than/as hard as every other NP problem, and is also an NP problem itself.

## 1.3 Travelling Salesman Problem

The input is a graph  $(V, E)$  with the cost of each edge  $\in E$  known. The objective is to find the path that visits every vertex  $\in V$  exactly once with a minimal cost. Some problems may require the path to start and end at the same node.

This can be converted to a decision problem: given a length  $X$ , is there a path with cost  $< X$ ? Alternatively, each vertex can be given as an  $(x, y)$  point in space, and the euclidean distance  $(\sqrt{(x_1 - x_2)^2 + (y_1 - y_2)^2})$  between points is used as the cost. The classic problem is NP-Hard, the decision problem is NP-Complete.

## 2 Branch and Bound

### 2.1 Partial Configurations

A configuration is a possible solution to a problem: so a partial configuration is a partial solution. For example, a partial configuration of a TSP is a path that doesn't contain every vertex. A partial configuration can be extended (e.g. adding another vertex) to make another configuration, either complete or partial.

Every complete configuration can only be obtained from **one** partial configuration in NP problems. In EXP problems like chess, this doesn't hold. The initial partial configuration (for TSP a path with no vertices) can be extended to every other configuration.

### 2.2 Branching

The same as extending, this procedure generates new configurations from a partial configuration. This depends on the problem, but it always takes one step: that is, it only generates the neighbourhood of the configuration. Again using TSP as an example, extending is only adding one vertex: if two were added at a time, some configurations might be missed.

By making every possible branch starting from the initial configuration (state), the entire search space of the problem can be traversed. However, blindly branching will take a very long time.

### 2.3 Bounding

Rather than branching every possibility, this uses the cost function to decide whether or not to follow a branch. The bounding procedure keeps a track of the best cost found so far, and when branching checks the best cost obtainable from the generated state or its descendents. If this cost is worse (higher) than the current best cost, the algorithm doesn't follow the branch, since it can't provide a better configuration.

When calculating the cost, the algorithm normally ignores validity to get a faster result. For example in TSP, the cost of a partial configuration is:

$$\text{bound}(PC) = \text{cost}(PC) + \text{min.cost edge from last node to node not in the PC} + \text{min. cost edge from all} \quad (1)$$

Obviously, this might not be the cost a valid path: but it represents a lower bound. For a complete configuration, this is just the cost of the path. Creating a minimum valid path would take as long as generating all the descendents from the state, so defeats the purpose.

## 2.4 Algorithm

A  $to\_visit = [initial\_state]$ ,  $best\_bound = \infty$

B  $curr = \text{Null}$ ,  $best = \text{Null}$

C while  $to\_visit$  is not empty:

1  $curr = \text{state from } to\_visit \text{ with lowest bound}$

2 remove  $curr$  from  $to\_visit$

3 if  $bound(curr) \geq best\_bound$  : *(If the best unvisited state is worse than the current best, the search is over)*

break

4 else:

i.  $children = \text{branch}(curr)$

ii. for  $child$  in  $children$ :

I. if  $bound(child) < best\_bound$  :

$\alpha$ . if  $child$  is a complete configuration: *(We've found a new best solution)*

$best = child$ ,  $best\_bound = bound(child)$

$\beta$ . else: *(We've found a branch that might lead to a best solution)*

$to\_visit.add(child)$

This algorithm is greedy, in that it always chooses the node with the lowest cost at each iteration. However, since it keeps a track of all possible nodes it won't miss an optimal solution. Until it finds a complete configuration it will add all branches to *to\_visit*, as the initial bound is infinite.



## 3 Shortest Path Algorithms

### 3.1 Graphs

A graph is normally represented as  $(V, E)$  for **V**ertices and **E**dges respectively. **Weighted graphs** have a cost/weight associated with each edge, and in **Directed graphs** each edge has a direction, a start node and an end node. In directed graphs, a **Cycle** is a set of edges that return to the same node (e.g.  $(a, b), (b, c), (c, a)$ ).

A **Fully Connected graph** has a direct connection between every node. In these graphs, there are  $\frac{v(v-1)}{2}$  edges, where  $v$  is the number of vertices. In **Connected graphs**, where all the nodes are connected directly or indirectly, there are at least  $v - 1$  edges, where  $v$  is the number of vertices.

### 3.2 Paths

In a directed graph, a path is an ordered list of edges: the **source** of a path is the start node of the first edge, and the **sink/destination** is the end node of the last edge. In a weighted directed graph, where the weight of an edge  $e$  is given by  $w(e)$ , the weight of a path  $P$  is:

$$\sum_{e \in P} w(e) \quad (2)$$

For two points  $a$  and  $b$ , we define the **Shortest Path Weight** as:

$$\delta(a, b) = \begin{cases} \text{Weight of Path with Lowest Weight} & , \text{if there is a path(s) between } a \text{ and } b \\ \infty & , \text{otherwise} \end{cases} \quad (3)$$

The shortest path is the path with this weight. Note that by this definition there can be **multiple shortest paths**, as long as each path has the lowest weight.

By definition, the weight of a path with a negative cycle is  $-\infty$ , since any path can just repeat the cycle infinitely. Most shortest path algorithms stop or break when they encounter a negative cycle, since the path is 'longer' in that it takes more edges, and the concept of  $-\infty$ .

### 3.3 Shortest Path Properties

- Any subpath (subset) of a shortest path is also a shortest path
- The concatenation of two shortest paths is not necessarily a shortest path
- A shortest path can have at most  $v-1$  edges, where  $v$  is the number of vertices
  - If there are more than  $v-1$  edges, this means there's a vertex has been visited more than once, so there's a cycle
  - negative-cost cycles aren't allowed by most algorithms, positive-cost cycles won't exist in a shortest path since they add weight, zero-cost cycles can be removed for no change
  - Therefore there are no cycles in a shortest path, so there can't be more than  $v-1$  edges.

### 3.4 Multiplicative Path Weights

For certain applications, rather than a sum of edge costs the product is used (e.g converting between various currencies). This can be converted to a summation by converting each edge weight to a logarithm. i.e if  $w(a,b)$  is  $x$ , set it to  $\ln x$ . This converts the product to a sum, since  $\sum_{i=0}^n \ln(x_i) = \ln(\prod_{i=0}^n x_i)$ , so standard techniques can be used.

### 3.5 Single Source

Any path has a source and end (sink). A single-source is a graph where one node  $s$  is designated as the source, and the shortest path to each node is calculated from it. Each node  $v$  in such a graph has the following two attributes:

$v.d$  is the shortest path estimate, the current upper bound on the shortest path from  $s$  to  $v$ . This is initialised to  $\infty$  for all nodes, then  $s.d$  is set to 0.

$v.\pi$  is the node before  $v$  on the current shortest path from  $s$  to  $v$ . This is initialised to Nil for all nodes, since a path hasn't been defined yet.

At the end of an algorithm,  $v.d = \delta(s, v)$  for all  $v$ , and  $v.\pi$  is the predecessor on the shortest path for each vertex.

#### 3.5.1 Relaxation

The process of reducing the upper bound of a node, this is used by the following algorithms to generate the shortest paths. Where  $u$  and  $v$  are nodes,  $(u,v)$  is the edge from  $u$  to  $v$  and  $w$  is the cost function for an edge:

$$\begin{aligned} \text{If } v.d > u.d + w(u,v): \\ v.d &= u.d + w(u,v) \\ v.\pi &= u \end{aligned} \tag{4}$$

For this,  $u.d \neq \infty$ . So in the first step of an algorithm, only the nodes connected from  $s$  can be initialised. Also, if the bound on source node of an edge ( $u.d$  in the example) is reduced, that means the edges from it can be further reduced: so care has to be taken to re-relax edges if required.

### 3.6 Shortest Paths Tree

A tree based on a graph  $G$  and rooted at  $s$  such that, for every other node  $v$ , the path from  $s$  to  $v$  in the tree is the shortest path from  $s$  to  $v$  in  $G$ . Unlike the traditional definition, the tree only contains one path from  $s$  to  $v$ .

If a node doesn't appear in the tree then there's no path from  $s$  to  $v$  in  $G$ . Using this approach to store shortest paths is  $O(v)$  space, while using a list would be  $O(v^2)$ .

### 3.7 Bellman-Ford Algorithm

We've seen in previous sections that for a graph  $(V,E)$ , the shortest path between any two nodes/vertices has at most  $V-1$  edges. We've also seen that relaxing an edge reduces the bound on the destination node, and this relaxation spreads out from the source node. Therefore, if we relax every edge in the graph  $V-1$  times, we guarantee that the bound on every node is equal to the shortest path weight.

**Bellman\_Ford(V,E):**

```
A   $v.d = \infty, v.\pi = \text{Nil} \ \forall v \in V$ 
B   $s.d = 0$ 
C  for  $i$  in  $\text{range}(0, V-1)$ : (range(a,b) takes values a to b-1)
    for  $(a,b)$  in  $E$ :
        relax( $a,b$ )
D  for  $(a,b)$  in  $E$ :
    if  $(a,b)$  can be relaxed
        return False
E  return  $(V,E)$ 
```

Since every shortest path is less than  $V-1$ , if an edge can still be relaxed after step C that means there's a negative cycle: this would give us a weight of  $-\infty$ , so the algorithm just rejects it by returning false.

This algorithm is  $O(VE)$  - it loops through all the edges  $V$  times in total ( $V-1$  times in step C, 1 time in step D). It returns the graph, with  $d$  and  $\pi$  set for each vertex.

#### 3.7.1 Improvements

- Edges only need to be updated when their source nodes are updated: the current algorithm updates every edge every time, this can be reduced by keeping a track of which nodes get updated.
- Most shortest paths aren't going to be  $V-1$  edges long: the loop can terminate early if there are no edges to relax at a generation. A shortest path of length  $k$  will finish relaxing every node in it in generation  $k$ .
- The negative cycle check can be done in the first loop: by keeping a track of the updated nodes, if there are any nodes to relax at iteration  $V$  then there's a negative cycle

## 3.8 Dijkstra's Algorithm

An alternate single-source shortest-paths algorithm that is faster than the Bellman-Ford, **but can't handle negative edge weights**. If there are negative edge weights/cycles, the algorithm doesn't raise a flag: it finishes executing, but the answers produced will be incorrect.

### 3.8.1 Lemmas Used

**Triangle Inequality:**  $\delta(u, v) \leq \delta(u, x) + w(x, v)$ . i.e. the shortest path between two points can't be further relaxed

**Upper Bound:**  $v.d \geq \delta(s, v)$  at every point of the algorithm

**No Path:** If there is no path between S and V,  $v.d = \delta(s, v) = \infty$

**Convergence:** Once  $v.d = \delta(s, v)$ , there is no change in v.d

### 3.8.2 Algorithm

In the below, N.adj is the list of nodes that are connected to N by an outgoing edge (i.e.  $N \rightarrow n$ ), and `extract_min(Q)` returns the node with the lowest upper bound in Q.

**Dijkstra(V,E):**

A  $v.d = \infty, v.\pi = \text{Nil} \ \forall v \in V, s.d = 0$

B  $S = \emptyset, Q = V$

C while  $Q \neq \emptyset$ :

1  $N = \text{extract\_min}(Q)$

2  $S.\text{add}(N)$

3 for n in N.adj:

$\text{relax}(N, n)$

D return (V,E)

Each edge is only relaxed once, since each node is only considered once. As the nodes are reached in a greedy order and there are no negative edge weights, this means that the nodes are reached as fast as possible and thus there shouldn't be a shorter path. The algorithm returns the graph, with with d and  $\pi$  set for each vertex.

### 3.8.3 Proof of Correctness

If we can prove that for any  $u \in S$ ,  $u.d = \delta(s, u)$ , then at the end all nodes are in  $S$  so all have the shortest path.

- We can prove this by induction:
- At  $S=\emptyset$ , trivially true.
- At  $S=\{s\}$ ,  $s.d = \delta(s, s) = 0$

**Hypothesis** We assume that for  $S=X$ , all nodes in  $S$  and the edges from them have been relaxed (this is what the algorithm does at each step)

- At  $S = X + \{u\}$ , a proof by contradiction:
  - Assume the contradiction, that  $u.d \neq \delta(s, u)$ 
    - \*  $u$  can't be  $s$ , since  $s.d = \delta(s, s) = 0$
    - \* There must be a path  $s \rightarrow u$ , otherwise  $s.d = \delta(s, s) = \infty$
  - Let  $y$  be the first node on  $s \rightarrow u$  that's NOT in  $S$  (this might be  $u$ ), and  $x$  be  $y$ 's predecessor (so  $x \in S$ )
    - \* Since  $y$  is before or equal to  $u$ ,  $\delta(s, y) \leq \delta(s, u)$
    - \* We know that  $x.d$  and  $(x, y)$  have been relaxed, by the inductive hypothesis
    - \* By convergence, since  $x.d$  and  $(x, y)$  have been relaxed,  $y.d$  is relaxed as well, so  $y.d = \delta(s, y)$
    - \* **Since the algorithm chooses the lowest bound each time**,  $u.d \leq y.d$
  - Combining the above, this gives us  $u.d \leq y.d = \delta(s, y) \leq \delta(s, u)$
  - $u.d$ , by definition, can't be  $< \delta(s, u)$ . Therefore,  $u.d = \delta(s, u)$
  - This contradicts our assumption, so it can't hold. Therefore we've proved that  $u.d = \delta(s, u)$  if  $u$  is the node with the lowest bound

### 3.8.4 Implementation and Performance

The algorithm's performance depends on the implementation of  $Q$ :

- If  $Q$  is a regular list, `extract_min` is  $O(V)$  every time ( $V$  times across the algorithm), and updating the bound on a vertex is  $O(1)$  ( $E$  times across the algorithm) so the algorithm is  $O(V^2 + E) \approx O(V^2)$
- If  $Q$  is a binary min-heap, `extract_min` is  $O(\log V)$  every time ( $V$  times across the algorithm), and updating the bound on a vertex is  $O(\log V)$  ( $E$  times across the algorithm) so the algorithm is  $O((V + E) \log V) \approx O(E \log V)$

Therefore, this depends on if the graph has more edges or vertices. A dense graph has more edges so a regular list works best, a sparse one has more vertices so the min-heap is better. Dense/Sparse can be determined by  $\frac{v^2}{\log v} : E$ .  $E$  is usually lower, so the min-heap is more commonly used.

## 3.9 Johnson's Algorithm

Both of the above algorithms give the shortest path to each node from a single source node. To get this for each node, we could run Dijkstra/Bellman-Ford once for each node as the source. But the former can't handle negative weights, and the latter would be  $O(V^2E)$ .

### 3.9.1 Re-Weighting

An easy fix to the problem is removing all the negative edge weights. Adding a constant  $M$  to each edge to make all the weights positive would fix that problem, but then it would add  $M$  to the cost of a path for each edge in it: since negative weights give a shorter path weight with more edges, this will change the results.

The new weight function ( $\hat{w}$ ) needs to preserve the shortest paths, and be positive for every edge. This can be achieved by giving each vertex  $v$  a value  $h(v)$ , and defining  $\hat{w}$  as:

$$\hat{w}(a, b) = w(a, b) + h(a) - h(b) \quad (5)$$

Using this equation, the weight of a path  $P$  would be:

$$\hat{w}(s, v) = \sum_{(a,b) \in P} w(a, b) + h(s) - h(v) \quad (6)$$

This is because, other than the source/sink, every node  $n$  on the path will have an incoming edge (so  $-h(n)$ ) and an outgoing edge ( $+h(n)$ ). Therefore, for any path between the nodes, the added values will be the same: this preserves the shortest paths. Calculating  $h(v)$  to remove negativity is shown in the next section.

### 3.9.2 Calculating $h(v)$

Since the aim is to remove negative weights, we need to show that for an edge  $(a,b)$  that  $h(a) - h(b) \geq 0$ .  $h(v)$  is computed by creating a new node, say  $s$ , connecting it to every other node with a zero-cost edge, then setting  $h(v) = \delta(s, v)$ .

Briefly, since  $s$  has access to every other node (and thus every possible path in the graph),  $\delta(s, v)$  is the shortest path into  $v$  from any other node. Since shortest is by path weight, we can say this is the most negative path weight leading into  $v$ . If this is subtracted from any incoming path weight, it'll end up as positive. A more mathematical proof:

$$\begin{aligned} \text{By the triangle inequality, } \delta(s, v) &\leq \delta(s, u) + w(u, v) \\ \text{Therefore, } \delta(s, v) - \delta(s, u) &\leq w(u, v) \\ \hat{w}(u, v) &= w(u, v) + h(u) - h(v) \\ &= w(u, v) + \delta(s, v) - \delta(s, u) \\ &\geq 0 \quad \forall u, v \end{aligned}$$

Since we compute  $h(v)$  using Bellman-Ford,  $s$  has access to every path so it can also check if there's a negative cycle and terminate. Using Dijkstra wouldn't work, since there would be negative edges.

### 3.10 Algorithm

Since the algorithm has to store the shortest path for each algorithm, it uses a  $V \times V$  matrix. Normally the algorithm stores either  $d$  or  $\pi$  for each vertex for each source, this pseudocode stores both. In the below pseudocode,  $G.V$  and  $G.E$  are the vertices/edges for a graph  $G$ .

**Johnson( $G, W$ ):**

A  $G' = (G.V, G.E)$  ( $G'$ 's is a copy of  $G$ , which we use to generate node weights)

B  $G'.V.add(s)$  (*Add the extra node*)

C for  $v$  in  $V$ : (*The original list of vertices*)

1  $G'.E.add((s, v))$

2  $w(s, v) = 0$  (*Adding zero-cost edges*)

D  $G'' = \text{Bellman\_Ford}(G'.V, G'.E)$  ( $G''$  is the graph with the shortest paths from  $s$ )

E If  $\text{Bellman\_Ford}$  returned false:

return False

F for  $(a, b)$  in  $E$ : (*Setting the new weights*)

$w(a, b) = w(a, b) + G''.V[a].d - G''.V[b].d$  ( $G''.V[v].d = h(v)$ )

G Results = []

H for  $v$  in  $V$ : (*Run Dijkstra for each vertex in the original list*)

1 Set  $v$  as source

2  $G_v = \text{Dijkstra}(G.V, G.E)$

3 Results.append( $G_v.V$ ) (*Add the vertices from the solved graph to the list of results*)

I return Results

All the weights will be off by the node values: they can be fixed before step H.2. The weight of the shortest path to node  $v$  from node  $u$  can be fixed by adding  $h(u) - h(v)$ : essentially reversing the effects of the re-weighting.

## 4 Flow Networks

Flow networks are directed graphs with two additions:

- Each edge  $(u,v)$  has a capacity  $c(u,v)$
- Two distinct nodes are marked as the source and sink

Usually, optimising a flow network is finding the maximum flow: the amount of units that can travel from the source to the sink simultaneously. Units travel along the edges, but an edge can only support up to its capacity simultaneously. Other problems may be the minimum flow that hits a certain target, or having multiple types of flow at the same time.

Most of the below algorithms assume that if an edge  $(a,b)$  exists, then the edge  $(b,a)$  also exists. If it's not present in the given graph, it's given a default capacity of zero.

### 4.1 Flow

The flow  $f(u,v)$  of an edge  $(u,v)$  is the amount of units passing through the edge: it's abstracted to a function that maps edges to real numbers. Flow is subject to the following:

1.  $f(a,b) \leq c(a,b) \forall (a,b)$  - Flow for an edge is always less than/equal to capacity
2. For all nodes except source/sink, incoming flow = outgoing flow (net flow on a node = 0)
3. If  $f(a,b) > 0$ , then  $f(b,a) = 0$

A **saturated edge** is one where  $f(a,b) = c(a,b)$ . The **value of a flow** in a network is defined as the net flow from the source/net flow into the sink.

A flow can be decomposed into **paths or cycles**, which start from the source and end at the sink. There can be at most  $m$  paths/cycles, where  $m$  is the number of edges.



## 4.2 Max Flow

More of a general approach than an algorithm, this uses the concept of residual capacity to iteratively build the optimal flow. The following algorithms use the ideas here.

## 4.3 Residual Capacity

The residual capacity for an edge (a,b) is defined as:

$$c_r(a, b) = \begin{cases} c(a, b) - f(a, b) & , \text{ if } f(a, b) > 0 \\ c(a, b) + f(b, a) & , \text{ if } f(b, a) > 0 \end{cases} \quad (7)$$

By the definition of a flow, if  $f(u, v) > 0$ , then  $f(v, u) = 0$ . The residual capacity of an edge with a flow is how much more flow it can take (capacity-flow), and the backward edge has how much flow can be reversed (capacity + reverse edge flow).

A **Residual Edge** is an edge with residual capacity  $> 0$ . The **Residual Network** of a flow  $f$  is the flow network made of residual edges after applying  $f$ . This network can be used to find flows not included in  $f$ , and these can be 'added' to  $f$ .

## 4.4 'Adding' Flows

By using the residual network, there might be a flow which improves the current one: this flow is 'added' to the current flow, altering it and increasing the overall flow. Assuming there are two flows  $F_1$  and  $F_2$ , adding them together is done edge-by-edge to produce the flow  $F'$ . Assuming that  $F_1(u, v) > 0$  (and therefore  $f(v, u) = 0$ ):

- If  $F_2(u, v) > 0$ , then  $F'(u, v) = F_1(u, v) + F_2(u, v)$ 
  - Both flows are in the same direction for this edge, just add them together
- If  $F_2(v, u) > 0$ , then  $F'(u, v) = \max(0, F_1(u, v) - F_2(v, u))$  and  $F'(v, u) = \max(0, F_2(v, u) - F_1(u, v))$ 
  - The flows run in opposite direction, so think of them as colliding
  - The larger flow direction is set to the difference of the flows
  - The smaller flow direction is set to 0

## 4.5 Algorithm

A  $F = \text{Zero Flow } (f(a, b) = 0 \text{ for all } (a, b))$

B while True:

1  $N = \text{Residual Network of } F$

2  $F' = \text{Flow in } N$

3 If  $F'$  does not exist:

    break

4  $F = F.\text{add}(F')$

C return  $F$

## 4.6 Ford\_Fulkerson Method

### 4.6.1 Augmenting Path

Building on the definition of a residual network, a **Residual Path** is a path (collection of consecutive edges) in the residual network. An **Augmenting Path** is a residual path from the source to the sink: essentially, it's a part of a flow in the residual network. The residual capacity of this augmenting path is the lowest residual capacity of its edges. i.e for an augmenting path  $p$ :

$$c_r(p) = \min_{(a,b) \in p} c_r(a,b) \quad (8)$$

### 4.6.2 Path Flow

Given a residual network  $R$  and an augmenting path  $P$ , the path flow for any edge  $(u,v)$  is:

$$f_P(u,v) = \begin{cases} c_r(p) & , (u,v) \text{ is on } P \\ 0 & , otherwise \end{cases} \quad (9)$$

For the path flow, the only flow in the network is on the residual path.

### 4.6.3 Augmentation

When adding the path flow  $F_p$  from an augmenting path  $P$ , the flow value is constant for every edge: its the capacity of the augmenting path, say  $q$ . This can be used to speed up addition. If an edge  $(a,b) \in P$  ( $a \rightarrow b$  is from the source to the sink),  $f$  is the current flow and  $F$  is the resulting flow then:

- $F(a,b) = f(a,b) + \max\{q-f(b,a), 0\}$ 
  - If  $f(a,b) > 0$ , then the flows were in the same direction and  $f(b,a) = 0$ : so they're just added
  - if  $f(a,b) = 0$ , then the flows were in the opposite direction, and  $f(b,a) \geq 0$ :  $f(a,b)$  is set to the result of removing the original flow from the augment
- $F(b,a) = \max\{f(b,a)-q, 0\}$ 
  - The flow in the augmenting path is in the opposite direction: if they collided  $f(b,a)$  is set to the difference, if  $f(b,a)$  was 0 it stays at 0

#### 4.6.4 Algorithm

A  $F = \text{Zero Flow } (f(a,b) = 0 \text{ for all } (a,b))$

B while True:

1  $N = \text{Residual Network of } F$

2  $F_p = \text{Augmenting path in } P$

3 If  $F_p$  does not exist:

    break

4  $F = F.\text{augment}(F_p)$

C return  $F$

The advantage over the basic method is its easier to find augmenting paths than full flows.

#### 4.6.5 Performance

For a graph  $(V,E)$ : constructing the residual network is  $\Theta(E)$  (checking edge by edge), finding an augmenting path is  $O(E)$  (For example, a breadth-first search), and augmenting is  $O(V)$  (each vertex appears max. once).

This means each iteration is normally  $O(E)$  (most graphs are sparse,  $\frac{v^2}{\log v}:E$ ). The number of iterations depends on which augmenting path is selected.

The bound on the number of iterations is the size of the maximum flow  $(|f * |)$ . This is obtained when the flow is comprised entirely of distinct paths (i.e size of flow = number of paths in flow). The Ford\_Fulkerson method discovers one path at a time, so the overall runtime will be  $O(|f * | * E)$ .

#### 4.6.6 Edmonds\_Karp Method

A variation of the Ford\_Fulkerson method: select the shortest (lowest number of edges) augmenting path each time. This can be found using a Breadth-First Search or similar. The performance is  $O(VE^2)$ , with the number of iterations being  $O(VE)$ . This is obtained by there being at most  $V-1$  paths in any network (every time a path is augmented, one edge is saturated), and any path length repeating at most  $E$  times. The intuition behind these: every time an augmenting  $V-1$

## 4.7 Cuts in a Flow Network

A **Cut** is splitting the nodes in a network into two groups: group S contains the source, group T contains the sink.

- The **Capacity** of a cut is  $\sum c(u, v)$  where  $u \in S, v \in T, (u, v) \in E$  : the capacity of the edges going from S to T
- The **Net Flow** of a cut is  $\sum f(a, b) - \sum f(c, d)$  where  $a, c \in S, b, d \in T, (a, b), (c, d) \in E$  : the flow from S to T minus the flow from T to S

For any cut,  $\text{flow}(\text{cut}) = \text{flow}(\text{sink}) \leq \text{capacity}(\text{cut})$  (equal when there is no flow from S to T). Therefore, **max. flow in a network  $\leq$  min. capacity of all cuts.**

### 4.7.1 Max-Flow Min-Cut Theorem

1. Max. flow in a network  $\leq$  min. capacity of any cut in the network
2. The following three statements are equivalent:
  - (a) F is a max. flow in G
  - (b) There's no augmenting path in the residual network  $G_F$
  - (c) For some cut (S,T) in G,  $c(S,T) = f(S,T) = F$

Proof:

**a→b** If there was an augmenting path, it could be used to increase the flow. The flow can't be increased, so there's no augmenting path

**b→c** No augmenting path means all the paths from source to sink are saturated, which means there are no reverse edges. All edges from S to T in a cut will be at capacity, all edges from T to S will be empty.

**c→a** When  $c(S,T) = F(S,T)$ , we've shown that this is a max flow. Since  $f(S,T) = F$ , F is a max flow

This is also a proof for the Ford-Fulkerson algorithm: **when there are no augmenting paths left, a max. flow has been found.**

## 4.8 Flow-Feasibility Problems

A variation on flow networks, rather than a single source/sink each vertex has a value  $d(v)$ . Usually  $\sum_{v \in V} d(v) = 0$ .

- If  $d(v) > 0$ , it's a **Supply Node**. Supply nodes have an outgoing flow (negative).
- If  $d(v) < 0$ , it's a **Demand Node**, with a demand of  $\text{abs}(d(v))$ . Demand nodes have an incoming flow (positive).
- If  $d(v) = 0$ , it's a **Transitional Node**.

These problems try to find a flow such that  $d(v) = 0$  for all nodes, and can be reduced to max. flow problems.

### 4.8.1 Reduction to Max. Flow Problems

1. Make a source node and connect it to each supply node  $s$  with an edge of capacity  $d(s)$ .
2. Make a sink node and connect it to each demand node  $x$  with an edge of capacity  $d(x)$ .
3. If the max. flow saturates the edges from the source/into the sink, that's a feasible flow.

## 4.9 Minimum Cost Flows

A common type of problem that builds off Flow-Feasibility: each edge is given a cost per flow, and the objective is to find the flow that satisfies the demands (or as close as possible) while minimising the total cost. Edges are usually labelled as  $(c,a)$ , where  $c$  is the capacity and  $a$  is the cost.

### 4.9.1 Flow Network Tweaks

- If the edge  $(a,b)$  exists and  $c(a,b) > 0$ , then  $c(b,a)$  is 0. Therefore in the residual network,  $c_r(a,b) = c(a,b) - f(a,b)$  and  $c_r(b,a) = f(a,b)$
- The cost of a reverse edge is the negative of the existing edge. i.e if edge  $(u,v)$  exists, then  $a(v,u) = -1 * a(u,v)$ .
- $d(v)$  in the residual network is  $d(v) + \text{incoming flow} - \text{outgoing flow}$  from the original network

### 4.9.2 Successive Shortest Paths Algorithm

This algorithm is very similar to the Edmonds\_Karp algorithm, with some key differences:

- Since there are multiple start/end points for shortest augmenting paths, a supply node is chosen (arbitrarily) and a shortest path tree to every demand node is created
  - The tree could also be created to every node then checked for demand
  - A list of supply nodes is kept: when a  $d(\text{node}) = 0$ , it can be removed from the list
  - The tree can be made with Dijkstra's if the costs are re-weighted, so  $O(E \log V)$
- The capacity of the chosen path is  $\min\{d(\text{supply}), \text{abs}(d(\text{demand})), c_r(p)\}$ 
  - This is because previously the source/sink could release/accept infinite flow, but now the supply/demand also constrain how much flow can go through the path
- Unfortunately, the total number of iterations could be **exponential** with regards to  $V$

**Algorithm:**

A  $F = \text{Zero Flow}$  ( $f(a,b) = 0$  for all  $(a,b)$ )

B  $S = \text{list of supply nodes}$  (*Nodes where  $d(v) \neq 0$* )

C while True:

1  $N = \text{Residual Network of } F$

2  $v = \text{Arbitrarily chosen Supply node/vertex}$

3  $T = \text{Shortest augmenting path tree to demand nodes in } N \text{ reachable from } v$

4  $F_p = \text{Shortest augmenting path in } T \text{ from } v \text{ to } x$

5  $c_r(F_p) = \min \{d(v), d(x), c_r(F_p)\}$

6 If  $F_p$  does not exist:

    break

7  $F = F.\text{augment}(F_p)$

D return  $F$

As mentioned previously, each iteration is  $O(E \log V)$  to generate the tree, and the total number of iterations is exponentially-bounded.

The best known running time of a minimum cost flow algorithm is  $O(E^2 \log V)$ , and there are multiple polynomial-time algorithms.

## 4.10 Multi-Commodity Flows

These problems have a single flow network with **multiple types of flows** running simultaneously. Each type of flow (commodity) has its own source and sink (some networks might have multiple sources/sinks in a single node), and a demand to be fulfilled. Each commodity is represented as (**source (node), sink (node), demand (real value)**).

Problems are to find a simultaneous flow of multiple commodities to solve a global objective. The overall flow is composed of multiple sub-flows, each of which represents one commodity. All other flow rules are followed, though some problems don't account for capacity/have separate capacities for each commodity.

### 4.10.1 Types of Problems

1. Feasibility: find a flow that satisfies edge capacities and commodity demands.
2. Minimum-cost: find a flow that satisfies edge capacities, gets as close to commodity demands as possible, while having a minimum cost
3. Minimum-Congestion: Find a flow that satisfies demands while minimising the maximum edge congestion across the graph
  - $\text{Congestion}(a,b) = \frac{\text{flow}(a,b)}{\text{capacity}(a,b)}$
  - The solution might break feasibility. i.e max. congestion might be  $> 1$



## 5 Linear Programming

A method of minimising a given function according to various constraints. Effectively, if the constraints are plotted on a graph (by bounding the areas of the graph that don't break them), the minimum value of the function lies on one of the corners of the bounded area (intersections between constraints). Even if the area is unbounded, the corners still provide the minimum values. **If the function is linear and the variables accept real values, then this method is polynomial in the number of variables.**

**Objective Function:** The function to minimise.

- If this isn't present, then the program acts as a feasibility problem
- To Maximize a function  $f(x)$ , minimize  $-f(x)$

**Constraint Functions:** These are functions that the solution must satisfy.

- If a constraint is non-linear, they must be split into linear constraints.
- The 'standard form' for constraints is  $ax_1 + b_x + \dots \leq c$ . The exception is the non-negativity constraints, which are of the form  $x_i \geq 0$ .

### 5.1 Shortest Path

When converting problems to linear programs, care has to be taken to make sure the conversion doesn't remove necessary features from the problem. An example of this is the single-pair shortest path problem, to find the shortest path weight from a source node to a destination node.

#### 5.1.1 Program

**Notation:**  $v.d$  is written as  $v_d$ ,  $s$  is the source,  $t$  is the destination,  $E$  is the list of edges

**Objective Function:** Maximise  $d_t$

**Constraints :**

- $d_v \leq d_u + w(u, v) \quad \forall (u, v) \in E$
- $d_s = 0$

#### 5.1.2 Explanation

Considering we're trying to find the shortest path, maximizing is counter-intuitive. However if the problem was  $\min d_t$ , then it could be satisfied by setting  $d_v$  to 0  $\forall V$ . The first constraint forces the program to set  $d_v = \delta(s, v)$  or less, so only the shortest valid path and shorter paths will meet this constraint. Therefore the maximum path that meets this constraint is the required shortest path.

## 5.2 Max. Flow in a Flow Network

**Notation:**  $f(u,v) \rightarrow f_{uv}$ ,  $s$  is the source,  $t$  is the destination,  $E$  is the list of edges

**Objective Function:** Maximise net. flow from source

$$\text{Maximise } \sum_{(s,v) \in E} f_{sv} - \sum_{(v,s) \in E} f_{vs}$$

**Constraints :**

- $f_{uv} \leq c(u, v) \quad \forall (u, v) \in E$   
Flow < capacity for each edge
- $f_{uv} \geq 0 \quad \forall (u, v) \in E$   
Flow  $\geq 0$  for each edge
- $\sum_{(x,v) \in E} f_{xv} - \sum_{(v,x) \in E} f_{vx} = 0 \quad \forall x \in V - \{s, t\}$   
Net flow = 0 for all nodes except the source/sink

The constraints on the sink's flow aren't explicitly mentioned to minimise the number of constraints, but are still implied by the other constraints.

## 5.3 Min. Flows

Since LPs are polynomial, this general approach is better than the specialised approach shown earlier. However, this approach is for a single source/sink, rather than the supply/demand approach from earlier.

One constraint is that there are no 1-cycles (edges  $(a,b)$  and  $(b,a)$ ) allowed, but these can be easily fixed: add a node  $x$ , remove  $(b,a)$  and add the edges  $(b,x)$  and  $(x,a)$ , both with the capacity of  $(b,a)$  and one with 0 cost, the other with the cost of  $(b,a)$ .

### 5.3.1 Program

**Notation:**  $f(u,v) \rightarrow f_{uv}$ ,  $s$  is the source,  $t$  is the destination,  $E$  is the list of edges,  $d$  is the demand we place on the system

**Objective Function:** Minimise total cost

$$\text{Minimise } \sum_{(u,v) \in E} a(u, v) * f_{uv}$$

**Constraints :**

- $f_{uv} \leq c(u, v) \quad \forall (u, v) \in E$   
Flow < capacity for each edge
- $f_{uv} \geq 0 \quad \forall (u, v) \in E$   
Flow  $\geq 0$  for each edge
- $\sum_{(x,v) \in E} f_{xv} - \sum_{(v,x) \in E} f_{vx} = 0 \quad \forall x \in V - \{s, t\}$   
Net flow = 0 for all nodes except the source/sink
- $\sum_{(s,v) \in E} f_{sv} - \sum_{(v,s) \in E} f_{vs} = d$   
The flow from the source is  $d$

## 5.4 Multi-Commodity Feasibility

LPs are currently the only known polynomial solver for multi-commodity flows. Looking at the problem (see 4.10.1), its to see if the source/demand can be met for every flow while meeting capacity constraints. This means there's nothing to optimise.

### 5.4.1 Program

**Notation:**  $s_q/t_q/d_q$  is the source/destination/demand for commodity  $q$ ,  $E$  is the list of edges,  $f_{quv}$  is the flow for commodity  $q$  through edge  $(u,v)$  and  $f_{uv}$  is the net flow through an edge ( $f_{uv} = \sum_q f_{quv}$ )

**Objective Function:** None

**Constraints :**

- $f_{uv} \leq c(u, v) \quad \forall (u, v) \in E$   
Flow < capacity for each edge
- $f_{uv} \geq 0 \quad \forall (u, v) \in E$   
Flow  $\geq 0$  for each edge
- $\sum_{(x,v) \in E} f_{qxv} - \sum_{(v,x) \in E} f_{qvx} = 0 \quad \forall x \in V - \{s_q, t_q\} \quad \forall q$   
Net commodity flow = 0 for all nodes except the commodity source/sink for all commodities/edges
- $\sum_{(s,v) \in E} f_{qsv} - \sum_{(v,s) \in E} f_{qvs} = d_q$   
The flow from the source is  $d_q$  for each commodity

Once again, rather than adding separate constraints that  $\text{flow}(\text{sink}) = -d_q$ , since they're implied by the others these constraints can be omitted to shrink the problem.

## 5.5 Min. Congestion Multi-Commodity

Again recalling the problem (section 4.10.1), this is to minimise the maximum congestion  $\lambda$ : alternatively, that the congestion for every edge  $\leq \lambda$ . Another note is that a solution might break feasibility, so the upper capacity constraints need to be removed.

### 5.5.1 Program

**Notation:**  $\lambda$  is the target minimum maximum flow,  $s_q/t_q/d_q$  is the source/destination/demand for commodity  $q$ ,  $E$  is the list of edges,  $f_{quv}$  is the flow for commodity  $q$  through edge  $(u,v)$  and  $f_{uv}$  is the net flow through an edge ( $f_{uv} = \sum_q f_{quv}$ )

**Objective Function:** Minimize  $\lambda$

**Constraints :**

- $f_{uv} - \lambda * c(u, v) \leq 0 \quad \forall (u, v) \in E$   
Congestion  $\leq$  max. congestion for each edge
- $f_{uv} \geq 0 \quad \forall (u, v) \in E$   
Flow  $\geq 0$  for each edge
- $\sum_{(x,v) \in E} f_{qxv} - \sum_{(v,x) \in E} f_{qvx} = 0 \quad \forall x \in V - \{s_q, t_q\} \quad \forall q$   
Net commodity flow = 0 for all nodes except the commodity source/sink for all commodities/edges
- $\sum_{(s,v) \in E} f_{qsv} - \sum_{(v,s) \in E} f_{qvs} = d_q \quad \forall q$   
The flow from the source is  $d_q$  for each commodity

Once again, rather than adding separate constraints that  $\text{flow}(\text{sink}) = -d_q$ , since they're implied by the others these constraints can be omitted to shrink the problem. The first constraints are converted into the standard form by:

$$\begin{aligned} \frac{f_{uv}}{c(u, v)} &\leq \lambda \\ f_{uv} &\leq \lambda * c(u, v) \\ f_{uv} - \lambda * c(u, v) &\leq 0 \end{aligned}$$

## 6 Searching a Graph

There are multiple ways to search graphs (especially weighted graphs), but the two simplest are **Breadth-First Search** and **Depth-First Search**. Both are very similar: DFS can be faster, but BFS is less reliant on the graph structure.

### 6.1 Breadth-First Search

In short: select a node, add all its children to the back of a list, then select the front node of the list and repeat. This particular variant also marks  $v.d$  and  $v.\pi$  for each vertex. Vertices that haven't been added to the list are undiscovered (white), vertices on the list are discovered (grey), vertices that were on the list are visited (black). Discovered is used to prevent the same vertex being added to the list multiple times, which is just needless computation.

#### 6.1.1 Algorithm

**BFS(V,E,s):**

```
A  $v.V = \text{False}$ ,  $v.d = -1$ ,  $v.\pi = \text{Nil} \ \forall v \in V$ 
B  $s.V = \text{True}$ ,  $s.d = 0$  ( $s$  is the node to start the search from)
C  $\text{to\_visit} = s$ 
D while  $\text{to\_visit}$  is not empty:
    1  $\text{curr} = s[0]$ 
    2 for child in  $V.\text{Adj}[\text{curr}]$  ( $V.\text{Adj}(v)$  is the list of nodes next to  $v$  (children of  $v$  if directed graph))
        i. if  $\text{child}.V = \text{False}$ : (To prevent nodes being added multiple times)
            A.  $\text{child}.V = \text{True}$ 
            B.  $\text{child}.d = \text{curr}.d + 1$ 
            C.  $\text{child}.\pi = \text{curr}$ 
            D.  $\text{to\_visit}.\text{append}(\text{child})$ 
    3 remove  $\text{curr}$  from  $\text{to\_visit}$ 
```

Any vertices that still have  $v.d = -1$  aren't reachable from  $s$ : in non-directed graphs this means they're not connected to any other node.  $v.V$  stands for  $v.\text{Visited}$ : rather than tracking 3 states, its sufficient to just track 2: a visual display would need all 3.

#### 6.1.2 Predecessor Graph

A graph built from the BFS results, that shows the edges and order taken by the search. For a graph  $G$ , the predecessor graph  $G_\pi$  consists of:

- $V_\pi = (v \text{ for } v \in G.V \text{ where } v.\pi \text{ is not Nil}) + s$
- $E_\pi = (v.\pi, v) \text{ for } v \text{ in } V_\pi - S$

## 6.2 Performance

$O(v)$  nodes are visited,  $O(E)$  edges are traversed across the program to check if the adjacent nodes have been visited, so in total its  $O(V+E)$ . It's not  $\Theta(V+E)$  because not every node/edge is visited, some might not be reachable.

## 6.3 Depth-First Search

Normally almost identical to BFS, the difference is that new nodes are added to the front of the list: this causes the algorithm to follow the currently chosen path as far as it can before backtracking and trying another one. Since BFS has the current node 'teleporting' around the graph, DFS is more-commonly used for things like maze traversals.

This version of the DFS uses a recursive method, and keeps track of when each vertex is first discovered ( $v.d$ ) and when it's finished ( $v.f$ ). These values allow us to track some further statistics and classify edges.

### 6.3.1 Algorithm

**Traversal(V,E):**

1.  $v.V = \text{False}$ ,  $v.d = -1$ ,  $v.f = -1$ ,  $v.\pi = \text{Nil} \ \forall v \in V$
2.  $\text{time} = 0$  (*time is a global variable*)
3. for  $v$  in  $V$ :  
    if  $v.V = \text{False}$ :  
        DFS( $V, E, v$ )

**DFS(V,E,v)**

1.  $\text{time} += 1$ ,  $v.d = \text{time}$
2. for  $u$  in  $V.\text{adj}[v]$ :  
    if  $u.V = \text{False}$ :  
        i.  $u.\pi = v$   
        ii. DFS( $V, E, u$ )
3.  $\text{time} += 1$ ,  $v.f = \text{time}$

Since every node is visited thanks to the loop in traversal,  $v.d$  and  $v.f$  will be  $> 0$  for every node. Since not all nodes are connected, this may create multiple trees within the same graph. Nodes with  $v.\pi = \text{Nil}$  are the roots of these trees.

### 6.3.2 Predecessor Graph

In addition to the definitions for BFS, the edges of the trees are also classified as one of the following:

**Tree Edge** An edge that is part of the search

**Forward Edge** An edge from a node  $n$  to a descendant  $d$  in the same tree ( $n.d \leq d.d$ , the node has already been finished)

**Backward Edge** An edge from a node  $n$  to an ancestor  $a$  in the same tree ( $n.d < a.d$ , the ancestor is currently on the stack) or a self-looping edge

**Cross Edge** An edge between two different trees/vertices of the same tree that aren't ancestor/descendant: any edge that isn't one of the above three

### 6.3.3 Performance

DFS is called on every vertex ( $O(V)$ ), and the loop in DFS runs  $\Theta(E)$  times in total across the whole program, once for each edge, so in total its  $\Theta(V + E)$ .

## 7 Topological Sort

A **Directed Acyclic Graph** is a directed graph that doesn't have cycles. By this definition, it's technically a tree, and there's a strict ordering on the nodes in it. Given a set of nodes from a DAG, a topological sort places them in an array such that if there's an edge from  $a$  to  $b$ , then  $\text{idx}(a) < \text{idx}(b)$ . Effectively, all edges go from left to right.

### 7.1 Algorithm

The algorithm is identical to DFS, with the exception that **each node is added to the front of the result list as it finishes**. This is the same as sorting the list of vertices by v.f.

The idea behind this is that if a node is finished, either there are no outgoing edges or all of its descendants are already on the list: therefore any edges will go to nodes with later indices.

Inserting to the front of a list is  $O(1)$ , so this algorithm is also  $\Theta(V + E)$ .

### 7.2 Shortest Paths in a DAG

Using this sorted array, a single-source shortest path graph can easily be calculated. Any node can be designated as the source and this method can handle negative edges: there are no cycles in a DAG, so there's no issue with negative cycles.

#### 7.2.1 Algorithm

**DAG\_Shortest(V,E,s):**

A  $D = \text{Topological\_sort}(V,E)$

B  $v.d = \infty, v.\pi = \text{Nil} \ \forall v \in V$

C  $s.d = 0$

D for  $v$  in  $D$ : (*traverses vertices in the sorted order*)

    for  $(a,b)$  in  $V.\text{adj}[v]$ :

$\text{relax}(a,b)$

Since any path between two nodes is from left to right in the sorted array, the edges can be relaxed in this order without worrying about the effects on previous nodes.



## 8 Minimum Spanning Tree

For an undirected weighted graph, the **Minimum Spanning Tree** is the tree (set of edges with no cycles) with the lowest weight that reach every vertex. By necessity, this has  $|V| - 1$  edges: any less won't reach every vertex, more would have a cycle.

### 8.1 Kruskal's Algorithm

**Kruskal(V,E)**

1. Create  $|V|$  sets, each with one vertex
2.  $T = []$
3. Sort  $E$  by ascending cost
4. for  $(a,b)$  in  $E$ :
  - If  $a$  and  $b$  aren't in the same set
  - i.  $T.add(a,b)$
  - ii. Merge the sets containing  $a$  and  $b$
5. return  $T$

Since each edge is added in order of ascending cost, this is a fairly greedy approach. The condition that the nodes aren't in the same set prevents a cycle: nodes in the same set are connected, so adding a new connection would make a cycle.

This algorithm can stop as soon as a spanning tree is created: this happens when there's a single vertex, since the algorithm makes sure no cycles are introduced. This is why the algorithm can't automatically take the first  $|V| - 1$  edges, since they may contain a cycle. The algorithm maintains a single tree throughout.

## 8.2 Prim's Algorithm

**Prim(V,E,r)**

1.  $v.d = 0, v.\pi = \text{Nil} \ \forall v \in V$
2.  $r.d = 0$  (*r is an arbitrary root node*)
3.  $Q = V$
4. while  $Q \neq \emptyset$ 
  - (a)  $u = \text{vertex with min. } d \text{ in } Q$  (*d represents the distance from the tree*)
  - (b) Relax all edges from  $u$
  - (c) Remove  $u$  from  $Q$
5. return  $(V,E)$

Starting from  $r$ , a tree can be constructed using  $v.\pi$  for each vertex. Like Dijkstra, the running time depends on  $Q$ : it could be a priority queue, a binary min-heap or something else. Since this method uses the shortest distance from the tree rather than the shortest edges, there's no chance of a cycle: this means  $\Theta(V)$  iterations in total.

## 9 Genetic Algorithms

Genetic Algorithms work by evolving a pool of candidate solutions (a population-based method), optimising them in the process till it eventually converges to an optimal enough solution. This is a **meta-heuristic**: its a way to find a way to find an optimal solution. Each configuration is represented as an **individual**, which contains values for the variables of the given problem. **Binary Genetic Algorithms** represent individuals as a string of binary characters of length  $n$ . BGAs assume that **all** binary strings of length  $n$  represent a valid configuration: if this isn't true then there's an overhead in checking validity.

One change for this version of the Genetic Algorithm is to define the **fitness function** of a configuration: how good a solution is. This version tries to find the individual with the highest fitness function: this can be obtained by multiplying the cost function by -1.

This form of the genetic algorithm has 4 main parts:

1. Natural Selection
2. Pair Selection
3. Crossover
4. Mutation

Combining all of these together, the overall flow of the algorithm is:

- A Create a random initial population
- B While stopping criteria are unmet:
  - 1 Create pairs via Pair Selection
  - 2 Make new individuals via Crossover
  - 3 Filter population with Natural Selection
  - 4 Apply mutation to the population
- C Select the best performing individual as the solution

For problems that have constraints or multiple objectives to be optimised, having the cost function be a weighted sum of each individual requirement is sufficient.

### 9.1 Stopping Criteria

Any/all of the below can be used: it depends on the situation:

1. An acceptable (high enough fitness) solution has been found
2. The mean/standard deviation/range of the individuals is within a target value
3. No change in individuals best/cost/worst cost after  $X$  iterations
4.  $N$  iterations have elapsed

## 9.2 Pair Selection

This part of the algorithm determines how individuals are paired up for crossovers (creating new individuals). This pairing up is important as it can influence how genes spread throughout the population, and can prevent good genes being overshadowed by poor ones. All of the below are independent of the cost function. i.e they don't require it to be of a certain form, such as differentiable.

The first two are **Roulette Selections**. These give each individual a probability, and to generate a pair they randomly sample from the entire population twice using the probabilities. One way to do this is to give each individual a cumulative probability, then generate a random number and see which probability that corresponds to. e.g.

$$\begin{aligned} P &= (010, 110, 011, 101) \\ cProb(P) &= (0.3, 0.4, 0.6, 1.0) \\ Pair &= (0.37, 0.81) \\ &= (110, 101) \end{aligned}$$

### 9.2.1 Cost-Weighted Roulette

The probability for each individual is dependent on its **fitness function**, so the probability of  $i_n$  in the population  $P$  is calculated as:

$$prob(i_n) = \frac{Fitness(i_n)}{\sum_{x \in P} Fitness(x)} \quad (10)$$

### 9.2.2 Rank-Weighted Roulette

The probability for each individual is dependent on its rank in the population. Higher fitness functions are given a higher rank, so for a population  $P$  of size  $k$  the individual with the highest fitness function is given the rank  $k$ . The probability of  $i_n$  is:

$$prob(i_n) = \frac{Rank(i_n)}{\sum_{x=0}^k x} \quad (11)$$

This gives probability  $\frac{1}{\sum_{x=0}^k x}$  to the worst individual and  $\frac{k}{\sum_{x=0}^k x}$  to the first. For small populations, this gives a very skewed probability, and sorting/ranking is an overhead.

### 9.2.3 Tournament Selection

A small subset of individuals is randomly chosen, and the best performing individual from this subset is selected. This is done twice to get a pair, and the size of each subset is normally 2-3. Since in each instance only a trivial subset has to be sorted, it reduces computation on sorting and thus works well for problems with large populations.

### 9.3 Crossover

New individuals are primarily created using the crossover operation. To generate offspring we take parts from the selected two individuals and combine them to make new individuals. The new individuals are added to the population, and then natural selection is applied. Repeatedly doing just crossovers only covers a limited section of the search space, so isn't sufficient to find an optimal solution.

As previously mentioned, individuals are represented by a binary string (0s and 1s) of length  $n$ : this length  $n$  is constant for all configurations of a program.

#### 9.3.1 Single Point

This breaks each parent into two parts, and creates offspring by swapping the second part for each. A random number  $0 \leq i < n$  is generated, and used to split the chromosomes ( $n$  is the length of the binary string). Using slice notation, this can be shown as:

$$\begin{aligned} Child_1 &= Parent_1[0 : i] + Parent_2[i : end] \\ Child_2 &= Parent_2[0 : i] + Parent_1[i : end] \end{aligned}$$

#### 9.3.2 Double Point

This breaks each parent into 3 parts, and swaps the middle parts for each. This middle section is denoted by two crossover points, which are the start and end of the section respectively. These follow  $0 \leq i_1 < i_2 \leq n$ . Using slice notation, this can be shown as:

$$\begin{aligned} Child_1 &= Parent_1[0 : i_1] + Parent_2[i_1 : i_2] + Parent_1[i_2 : end] \\ Child_2 &= Parent_2[0 : i_1] + Parent_1[i_1 : i_2] + Parent_2[i_2 : end] \end{aligned}$$

### 9.4 Natural Selection

In this version, natural selection is used to bring the population back to its normal size after adding the crossover offspring. If there were originally  $P$  individuals, the algorithm generates probabilities with rank/cost-roulette then uses them to select  $P$  individuals from the new population.

### 9.5 Mutation

Mutation is the process that actually drives evolution, by making small changes to the individuals. Crossovers move the genes into potentially better solutions, mutations discover new solutions.

Mutation is determined using a mutation rate,  $q$ . To prevent good solutions from being mutated, we can designate the top  $e$  chromosomes as *elite*, and not consider them for mutation. In this version, for a population of size  $P$ ,  $qP$  individuals are randomly selected then mutated: this could be flipping some of their bits ( $0 \rightarrow 1$ ,  $1 \rightarrow 0$ ), re-arranging them or other small changes.

## 9.6 Travelling Salesman Problem

Representing configurations/tours as a binary string is difficult, so instead we can use a sequence of vertices: each individual will have the same vertices, but in a different order.

### 9.6.1 Fitness Function

Say for each tour  $t$  the cost is  $\text{cost}(t)$ , and the individual representing that tour is  $t_i$ . If for a population  $P$  the worst cost is  $\text{cost}_{\max}(P)$ , then the fitness for  $t_i$  is:

$$\text{fitness}(t_i) = \text{cost}_{\max}(P) - \text{cost}(t) + r \quad (12)$$

$r$  is a constant is used to make sure the fitness for all individuals is positive: having a fitness of zero or less breaks the cost-weighting roulette. Having  $r$  too high weakens the cost-weighting roulette, as the fractional difference between each individual becomes too small to effectively provide probabilities.

### 9.6.2 Crossover

Given two individuals  $a$  and  $b$ , children are the first half of one individual + the remaining cities in the order they appear in the other individual. e.g.

$$\begin{aligned} a &= [1, 3, 5, 7, 4, 2, 6, 8] \\ b &= [2, 3, 4, 8, 6, 1, 5, 7] \\ \text{child}_a &= [1, 3, 5, 7 | 2, 4, 8, 6] \\ \text{child}_b &= [2, 3, 4, 8 | 1, 5, 7, 6] \end{aligned} \quad (13)$$

For  $\text{child}_a$ :  $[1, 3, 5, 7]$  are copied directly, then 4, 2, 6 and 8 are added in the order they appear in  $b$ . For  $\text{child}_b$ :  $[2, 3, 4, 8]$  is copied and 1, 5, 7 and 6 are added.

This crossover can be improved by making sure that new configurations follow the triangle inequality/etc. . This requires extra computation, but prevents needless iterations with obviously inferior configurations.

## 10 Simulated Annealing

A general heuristic (technique for solving problems that provides an acceptable, but not necessarily optimal) for solving optimisation problems.

As mentioned in 1.1, a optimisation problem can be represented as  $(R, C)$ , which is (set of configurations, cost function). For simulated annealing the **neighbourhood** is also defined: for a config  $c$ ,  $\text{neighbourhood}(c)$  is a set of configurations that can be obtained by a simple modification to  $c$ . The algorithm uses this in its **Generation Mechanism**: given a configuration, it returns a random neighbour.

### 10.1 Generation Mechanism

#### 10.1.1 TSP

A configuration in the travelling-salesman problem consists of a list of vertices, which gives us an order of cities to visit. A generation mechanism could be to swap two non-consecutive edges: this is still a valid tour since all the cities are present. This is called the **2-opt heuristic**, and is often used in practice. The 3-opt and 4-opt versions are also used.

#### 10.1.2 Weighted Graph Bisection Problem

Given a weighted graph  $(V, E)$ , split  $V$  into two parts with an equal number of vertices such that the sum edge weights between the two halves is minimum: effectively, make a cut on the graph into equal halves with minimum edge weights between the halves.

Since the sizes of the halves ( $A$  and  $B$ ) stay the same, a generation mechanism is to transfer a node from  $A$  to  $B$  and vice versa. A variation is swapping two nodes from each, rather than one.

### 10.2 Why Not Local Search?

Local search is a simple optimisation method: given a configuration, search its neighbourhood for a new configuration with a better (lower) cost, move to this, then repeat. If there aren't any better neighbours, then stop. This works fine on some functions, but it may get stuck in **Local Minima**: a configuration that has a better cost than all of its neighbours, but not the most optimal configuration.

### 10.3 Temperature

Simulated annealing has a chance to select a neighbour with a worse cost than the current configuration, and uses a parameter known as the **Temperature** to control this: the higher the temperature, the more likely a worse configuration is allowed.

By allowing worse configurations the algorithm is less likely to be stuck in local minima, as it can break out of them if the temperature allows it. The temperature starts high then is slowly decreased as the algorithm progresses, so in later stages the algorithm reverts to a local search. The temperature changes every  $L$  iterations, and  $L$  usually increases as the algorithm progresses.

## 10.4 Probability

The exact probability equation to accept a worse configuration is:

$$p(worse) = \exp\left(\frac{C(config) - C(new)}{T}\right) \quad (14)$$

Where config is the current configuration ( $c(config) = a$ ), new is the worse configuration ( $c(new) = x$ ), and T is the temperature. By definition,  $p(worse)$  is  $\leq 1$ :  $e^{\frac{a-x}{T}} = 1$  when  $a = x$ , and is less otherwise as  $x > a$ . An increase in T increases the probability, an increase in x (a worse neighbour) decreases the probability.

## 10.5 Algorithm

**Simulated Annealing(R, C, initial\_config, T, L):**

```
A config = initial_config

B Do:
    for j in range(L):
        i. new = generate(config)
        ii. if  $C(new) < C(config)$ 
            config = new
        iii. elif  $\exp\left(\frac{C(config) - C(new)}{T}\right) > rand(0, 1)$ 
            config = new
    1 Update T and L

While (Stopping Criteria Unmet)

C Return config
```

## 10.6 Stopping Criteria

- T hits a defined minimum value
- There weren't any/enough transitions in the last temperature phase