CSS.201.1 Algorithms

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Finding Closest Pair of Points

FIND CLOSEST

Input: Set $S = \{(x_i, y_i) \mid x_i, y_i \in \mathbb{R}, \forall i \in [n]\}$. We denote $P_i = (x_i, y_i)$.

Question: Given a set of points find the closest pair of points in \mathbb{R}^2 find P_i , P_j that are at minimum l_2 distance

i.e. minimize $\sqrt{(x_i - x_j)^2 + (y_i - y_j)^2}$.

1.1 Naive Algorithm

Now the naive algorithm for this will be checking all pairs of points and take their distance and output the minimum one. There are total $\binom{n}{2}$ possible choices of pairs of points. And calculating the distance of each pair takes O(1) time. So it will take $O(n^2)$ times to find the closest pair of points.

Idea: $\forall P_i, P_i \in S$ find distance $d(P_i, P_i)$ and return the minimum. Time taken is $O(n^2)$.

1.2 Divide and Conquer Algorithm

Below we will show a Divide and Conquer algorithm which gives a much faster algorithm.

Definition 1.2.1: Divide and Conquer

- Divide: Divide the problem into two parts (roughly equal)
- Conquer: Solve each part individually recursively. If the subproblem sizes are small enough, however, just solve the subproblems in a straightforward manner.
- Combine: Combine the solutions to the subproblems into the solution.

1.2.1 Divide

So to divide the problem into two roughly equal parts we need to divide the points into two equal sets. That we can do by sorting the points by their x-coordinate. Suppose S^x denote we get the new sorted array or points. And similarly we obtain S^y which denotes the array of points after sorting S by their y-coordinate.

Algorithm 1: Step 1 (Divide)

1 Function Divide:

Sort S by x-coordinate and y-coordinate

$$S^x \leftarrow S$$
 sorted by x -coordinate

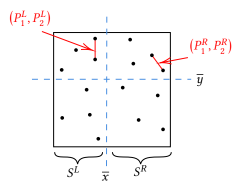
$$S^y \leftarrow S$$
 sorted by y-coordinate

$$\bar{x} \leftarrow \lfloor \frac{n}{2} \rfloor$$
 highest x -coordinate

$$\bar{y} \leftarrow \lfloor \frac{\bar{n}}{2} \rfloor$$
 highest y-coordinate

$$7 \quad S^L \longleftarrow \{P_i \mid x_i < \bar{x}, \ \forall \ i \in [n]\}$$

$$8 \mid S^R \longleftarrow \{P_i \mid x_i \geq \bar{x}, \ \forall \ i \in [n]\}$$



1.2.2 Conquer

Now we will recursively get pair of closest points in S_L and S_R . Suppose the (P_1^L, P_2^L) are the closest pair of points in S^L and (P_1^R, P_2^R) are the closest pair of points in S^R .

Algorithm 2: Step 1 (Solve Subproblems)

1 Function Conquer:

- Solve for S_L , S^R . 2
- (P_1^L, P_2^L) are closest pair of points in S_L .
- (P_1^R, P_2^R) are closest pair of points in S_R . $\delta^L = d(P_1^L, P_2^L), \, \delta^R = d(P_1^R, P_2^R)$ 4
- $\delta_{min} \longleftarrow \min{\{\delta^L, \delta^R\}}$

1.2.3 Combine

Now we want to combine these two solutions.

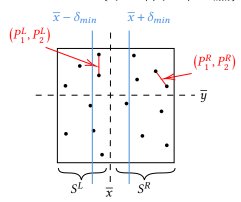
Question 1.1: We are not done

Is there a pair of points $P_i, P_i \in S$ such that $d(P_i, P_i) < \delta_{min}$

If Yes:

- One of them must be in S_L and the other is in S_R .
- x-coordinate $\in [\overline{x} \delta_{min}, \overline{x} + \delta_{min}].$
- $|y_i y_j| \leq \delta_{min}$

So we take the strip of radius δ_{min} around \overline{x} . Define $T=\{P_i\in S\mid |x_i-\overline{x}|\leq \delta_{min}\}$



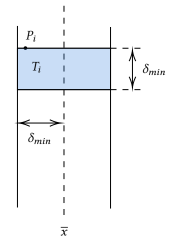
We now sort all the points in the T by their decreasing y-coordinate. Let T_y be the array of points. For each $P_i \in T_y$ define the region

$$T_i = \{ P_j \in T_y \mid 0 \le y_j - y_i \le \delta_{min}, j > i \}$$

Lemma 1.2.1

Number of points (other than P_i) that lie inside the box is at most 8

Proof: Suppose there are more than 8 points that lie inside the box apart from P_i . The box has a left square part and a right square part. So one of the squares contains at least 5 points. WLOG suppose the left square has at least 5 points. Divide each square into 4 parts by a middle vertical and a middle horizontal line. Now since there are 5 points there is one part which contains 2 points but that is not possible as those two points are in S_L and their distance will be less than δ_{min} which is not possible. Hence contradiction. Therefore there are at most 8 points inside the box.



Hence by the above lemma for each $P_i \in T_y$ there are at most 8 points in T_i . So for each $P_j \in T_i$ we find the $d(P_i, P_j)$ and if it is less than δ_{min} we update the points and the distance

1.2.4 Pseudocode and Time Complexity

Assumption. We will assume for now that for all $P_i.P_j \in S$ we have $x_i \neq x_j$ and $y_i \neq y_j$. Later we will modify the pseudocode to remove this assumption

```
Algorithm 3: FIND-CLOSEST(S)
```

```
Input: Set of n points, S = \{(x_i, y_i) \mid x_i, y_i \in \mathbb{R}, \ \forall \ i \in [n]\}. We denote P_i = (x_i, y_i).
    Output: Closest pair of ponts, (P_i, P_i, \delta) where \delta = d(P_i, P_i)
 1 begin
 2
           if |S| \leq 10 then
             Solve by Brute Force (Consider every pair of points)
 3
           S^x \leftarrow S sorted by x-coordinate,
                                                                          S^y \leftarrow S sorted by y-coordinate
 4
           \overline{x} \leftarrow \lfloor \frac{n}{2} \rfloor highest x-coordinate, \overline{y} \leftarrow \lfloor \frac{n}{2} \rfloor highest y-coordinate
           S^{L} \longleftarrow \{P_{i} \mid x_{i} < \bar{x}, \ \forall \ i \in [n]\}, \qquad S^{R} \longleftarrow \{P_{i} \mid x_{i} \geq \bar{x}, \ \forall \ i \in [n]\}
(P_{1}^{L}, P_{2}^{L}, \delta^{L}) \longleftarrow \text{Find-Closest}(S^{L}), \qquad (P_{1}^{R}, P_{2}^{R}, \delta^{R}) \longleftarrow \text{Find-Closest}(S^{R})
           \delta_{min} \longleftarrow \min{\{\delta^L, \delta^R\}}
 8
           10
11
            P_1 \longleftarrow P_1^L, P_2 \longleftarrow P_2^L
12
           T \longleftarrow \{P_i \mid |x_i - \overline{x}| \le \delta_{min}\}
13
           T_y \leftarrow T sorted by decreasing y-coordinate
14
           for P \in T_y do
15
                  U \leftarrow Next 8 points
16
                  for \hat{P} \in U do
17
                        if d(P, \hat{P}) < \delta_{min} then
18
                              \delta_{min} \longleftarrow d(P, \hat{P})
(P_1, P_2) \longleftarrow (P, \hat{P})
19
20
           return (P_1, P_2, \delta_{min})
```

Notice we used the assumption in the line 5 for finding the medians. So the line 4 takes $O(n \log n)$ times. Lines 5,6 takes O(n) time. Since \overline{x} is the median, we have $|S^L| = \lfloor \frac{n}{2} \rfloor$ and $|S^R| = \lceil \frac{n}{2} \rceil$. Hence FIND-CLOSEST(S^L) and FIND-CLOSEST(S^R) takes $T\left(\frac{n}{2}\right)$ time. Now lines 8 – 12 takes constant time. Line 13 takes O(n) time. And line 14 takes $O(n \log n)$ time. Since U has 8 points i.e. constant number of points the lines 16 – 20 takes constant time for each $P \in T_U$. Hence the for loop at

line 15 takes O(n) time. Hence total time taken

$$T(n) = O(n) + O(n \log n) + 2T\left(\frac{n}{2}\right) \implies T(n) = O(n \log^2 n)$$

Improved Algorithm for $O(n \log n)$ **Runtime** 1.3

Notice once we sort the points by x-coordinate and y-coordinate we don't need to sort the points anymore. We can just pass the sorted array of points into the arguments for solving the smaller problems. Their is another time where we need to sort which is in line 14 of the above algorithm. This we can get actually from S^y without sorting just checking one by one backwards direction if the x-coordinate of the points satisfy $|x_i - \overline{x}| \le \delta_{min}$. So

$$T_y = \text{Reverse}(\{P_i \in S^y \mid |x_i - \overline{x}| \le \delta_{min}\})$$

So we form a new algorithm which takes the input S^x and S^y and then finds the closest pair of points. Then we will use that subroutine to find closest pair of points in any given set of points.

```
Algorithm 4: FIND-CLOSEST-SORTED(S^x, S^y)
    Input: Set of n points, S = \{(x_i, y_i) \mid x_i, y_i \in \mathbb{R}, \forall i \in [n]\}.
               S^x and S^y are the sorted array of points with
               respect to x-coordinate and y-coordinate
               respectively
    Output: Closest pair of ponts, (P_i, P_i, \delta) where
                  \delta = d(P_i, P_i)
 1 begin
          if |S| \leq 10 then
 2
           Solve by Brute Force
         \overline{x} \leftarrow \lfloor \frac{n}{2} \rfloor highest x-coordinate
                                                                                                         Algorithm 5: FIND-CLOSEST(S)
          \overline{y} \leftarrow \lfloor \frac{n}{2} \rfloor highest y-coordinate
 5
          S^L \longleftarrow \{P_i \in S^x \mid x_i < \bar{x}, \ \forall \ i \in [n]\}
                                                                                                            Input: Set of n points,
         S_y^L \longleftarrow \{P_i \in S^y \mid x_i < \overline{x}\}
                                                                                                                        S = \{(x_i, y_i) \mid x_i, y_i \in \mathbb{R}, \ \forall \ i \in [n]\}.
         S^{R} \longleftarrow \{P_i \in S^x \mid x_i \geq \bar{x}, \ \forall \ i \in [n]\}
                                                                                                                        We denote P_i = (x_i, y_i).
 8
                                                                                                            Output: Closest pair of ponts, (P_i, P_i, \delta)
         S_u^R \longleftarrow \{P_i \in S^y \mid x_i \ge \overline{x}\}
                                                                                                                          where \delta = d(P_i, P_i)
          (P_1^L, P_2^L, \delta^L) \leftarrow \text{Find-Closest-Sorted}(S^L, S_y^L)
10
                                                                                                         1 begin
          (P_1^R, P_2^R, \delta^R) \leftarrow \text{Find-Closest-Sorted}(S^R, S_u^R)
11
                                                                                                                  if |S| \leq 10 then
                                                                                                         2
          \delta_{min} \longleftarrow \min{\{\delta^L, \delta^R\}}
12
                                                                                                                    Solve by Brute Force
         if \delta_{min} < \delta^L then
13
                                                                                                                  S^x \leftarrow S sorted by x-coordinate
           14
                                                                                                                  S^y \leftarrow S sorted by y-coordinate
                                                                                                         5
15
                                                                                                                  return FIND-CLOSEST-SORTED(S^x, S^y)
           P_1 \longleftarrow P_1^L, P_2 \longleftarrow P_2^L
16
          T \longleftarrow \{P_i \mid |x_i - \overline{x}| \le \delta_{min}\}
17
          T_y \leftarrow \text{Reverse}(\{P_i \in S^y \mid |x_i - \overline{x}| \leq \delta_{min}\})
18
          for P \in T_y do
19
               U \leftarrow Next 8 points
20
               for \hat{P} \in U do
21
                     if d(P, \hat{P}) < \delta_{min} then
22
                        \delta_{min} \longleftarrow d(P, \hat{P})(P_1, P_2) \longleftarrow (P, \hat{P})
23
24
          return (P_1, P_2, \delta_{min})
```

This algorithm only sorts one time. So time complexity for FIND-CLOSEST-SORTED(S^x, S^y) is

$$T(n) = 2T\left(\frac{n}{2}\right) + O(n) \implies T(n) = O(n\log n)$$

and therefore times complexity for FIND-CLOSEST(S) is $O(n \log n)$.

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1.4 Removing the Assumption

For this there nothing much to do. For finding the median \overline{x} if we have more than one points with same x-coordinate which appears as the $\left\lfloor \frac{n}{2} \right\rfloor$ highest x-coordinate we sort only those points with respect to their y-coordinate update the S^x like that and then take $\left\lfloor \frac{n}{2} \right\rfloor$ highest point in S^x . We do the same for S^y and update accordingly. All this we do so that S^L and S^R has the size $\frac{n}{2}$.

Median Finding in Linear Time

Median Find

Input: Set S of n distinct integers

Question: Find the $\left\lfloor \frac{n}{2} \right\rfloor^{th}$ smallest integer in *S*

2.1 Naive Algorithm

The naive algorithm for this will be to sort the array in $O(n \log n)$ time then return the $\left\lfloor \frac{n}{2} \right\rfloor^{th}$ element. This will take $O(n \log n)$ time. But in the next section we will show a linear time algorithm.

2.2 Linear Time Algorithm

In this section we will show an algorithm to find the median of a given set of distinct integers in O(n) time complexity. Consider the following two problems:

Rank-Find (S, k)

Input: Set *S* of *n* distinct integers and an integer $k \le n$

Question: Find the k^{th} smallest integer in S

Approximate-Split(S)

Input: Set *S* of *n* distinct integers

Question: Given S, return an integer $z \in S$ such that z where $rank(z) \in \left[\frac{n}{4}, \frac{3n}{4}\right]$

2.2.1 Solve Rank-Find using Approximate-Split

```
Algorithm 6: RANK-FIND(S,k)

Input: Set S of n distinct integer and k \in [n]
Output: k^{th} smallest integer in S

1 begin

2 | if |S| \le 100 then

3 | Sort S, return k^{th} smallest element in S

4 | z \leftarrow \text{Approximate-Split}(S) | (z \text{ is the } r^{th} \text{ smallest element for some } r \in \left[\frac{n}{4}, \frac{3n}{4}\right])

5 | S_L \leftarrow \{x \in S \mid x \le z\}, S_R \leftarrow \{x \in S \mid x > z\}

6 | if k \le |S_L| then

7 | return RANK-FIND(S_L, k)

8 | return RANK-FIND(S_R, k - |S_L|)
```

Certainly if we can solve Rank-Find (S, k) for all $k \in [n]$ we can also solve Median-Find. We will try to use both the problems and recurse to solve Rank-Find in linear time.

In the above algorithm $rank(z) \in \left[\frac{n}{4}, \frac{3n}{4}\right]$. So $\frac{n}{4} \leq |S_L|, |S_R| \leq \frac{3n}{4}$. For now suppose Rank-Find(S, k) takes $T_{RF}(n)$ time and Approximate-Split(S) takes $T_{AS}(n)$ time. Then the time taken by the algorithm is

$$T_{RF}(n) \le O(n) + T_{AS}(n) + T_{RF}\left(\frac{3n}{4}\right)$$

2.2.2 Solve Approximate-Split using Rank-Find

We first divide *S* into groups of 5 elements. So take $t = \lceil \frac{n}{5} \rceil$. Now we sort each group. Since each group have constant size this can be done in O(n) time. So now consider the scenario:



After sorting each of the groups we takes the medians of each group. Let z be the median of the medians. We claim that $rank(z) \in \left[\frac{n}{4}, \frac{3n}{4}\right]$.

```
Algorithm 7: Approximate-Split(S)
   Input: Set S of n distinct integers
   Output: An integer z \in S such that z where rank(z) \in \left[\frac{n}{4}, \frac{3n}{4}\right]
1 begin
        if |S| \le 100 then
2
         Sort, return Exact median
4
        S_i \leftarrow i^{th} block of 5 elements in S for i \in [t-1]
5
        S_t \leftarrow Whatever is left in S
        for i \in [t] do
         Sort S_i, Let h_i be the median of S_i
        T \longleftarrow \{h_i \mid i \in [t]\}
        return RANK-FIND (T, \lfloor \frac{t}{2} \rfloor)
10
```

So in the picture among elements in upper left the highest element is z and among the elements in lower right the lowest element is z. We will show that the number of elements smaller than z is between $\frac{n}{4}$ and $\frac{3n}{4}$. Lets call the set of elements in upper left box is S_u and the set of elements in lower right box is S_d .

2.2 Linear Time Algorithm Page 8

```
Lemma 2.2.1 |S_u|, |S_d| \ge \frac{n}{4}
```

Proof: $|S_u| \ge 3 \times \left\lfloor \frac{t}{2} \right\rfloor$. For $n \ge 100$, $3 \left\lfloor \frac{t}{2} \right\rfloor > \frac{n}{4}$. Hence $|S_u| \ge \frac{n}{4}$. Now similarly $|S_d| \ge 3 \left\lfloor \frac{t}{2} - 1 \right\rfloor \ge \frac{n}{4}$.

```
Lemma 2.2.2
```

Number of elements in *S* smaller than *z* lies between $\frac{n}{4}$ and $\frac{3n}{4}$.

Proof: Now number of elements in S smaller than $z \ge |S_u| \ge \frac{n}{4}$. The number of elements greater than $z \ge |S_d| \ge \frac{n}{4}$. So number of elements in S smaller than $z \le n - n$ number of elements greater than $z \le n - \frac{n}{4} = \frac{3n}{4}$.

Hence the Approximate-Split(S) takes time

$$T_{AS}(n) = O(n) + T_{RF}\left(\frac{n}{5}\right)$$

2.2.3 Pseudocode and Time Complexity

Hence using Approximate-Split the final algorithm for Rank-Find is the following:

```
Algorithm 8: RANK-FIND(S,k)
```

```
Input: Set S of n distinct integer and k \in [n]
   Output: k^{th} smallest integer in S
 1 begin
        if |S| \leq 100 then
         Sort S, return k^{th} smallest element in S
        S_i \leftarrow i^{th} block of 5 elements in S for i \in [t-1]
        S_t \leftarrow Whatever is left in S
 6
        for i \in [t] do
         Sort S_i, Let h_i be the median of S_i
        T \longleftarrow \{h_i \mid i \in [t]\}
        z \leftarrow \text{Rank-Find}\left(T, \left\lfloor \frac{t}{2} \right\rfloor\right)
10
        S_L \longleftarrow \{x \in S \mid x \leq z\}, S_R \longleftarrow \{x \in S \mid x > z\}
11
        if k \leq |S_L| then
12
          return RANK-FIND(S_L, k)
13
        return RANK-FIND(S_R, k - |S_L|)
14
```

Replacing $T_{AS}(n)$ in the time complexity equation of $T_{RF}(n)$ we get the equation:

$$T_{RF}(n) \le O(n) + T_{RF}\left(\frac{n}{5}\right) + T_{RF}\left(\frac{3n}{4}\right)$$

Let $T_{RF}(n) \le kn + +T_{RF}\left(\frac{n}{5}\right) + T_{RF}\left(\frac{3n}{4}\right)$. We claim that $T_{RF}(n) \le cn$ for some $c \in \mathbb{N}$ for all $n \ge n_0$ where $n_0 \in \mathbb{N}$. By induction we have

$$T_{RF}(n) \le kn + \frac{cn}{5} + \frac{3cn}{4} = \left(k + \frac{19c}{20}\right)n$$

To have $k + \frac{19c}{20} \le c$ we have to have $k + \frac{19c}{20} \le c \iff c \ge 20k$. So take $c \ge 20k$ and our claim follows. Hence $T_{RF}(n) = O(n)$. Since we can find any k^{th} smallest number in a given set of distinct integers in linear time we can also find the median in linear time.

Polynomial Multiplication

POLYNOMIAL MULTIPLICATION

Given 2 univariate polynomials of degree n-1 by 2 arrays of their coefficients (a_0,\ldots,a_{n-1}) and (b_0,\ldots,b_{n-1}) such that $A(x)=a_0+a_1x+\cdots+a_{n-1}x^{n-1}$ and $B(x)=b_0+b_1x+\cdots+b_{n-1}x^{n-1}$

respectively

Given 2 polynomials of degree n-1 find their product polynomial C(x) = A(x)B(x) of degree 2n-2Question:

by returning the array of their coefficients.

Naive Algorithm 3.1

We can do this naively by calculating each coefficient of C in O(n) time since for any $i \in \{0, \dots, 2n-2\}$

$$c_i = \sum_{j=0}^i a_j b_{i-j}$$

Since there are 2n-1 = O(n) total coefficient of C it takes total $O(n^2)$ time. In the following section we will do this in $O(n \log n)$ time.

3.2 Strassen-Schönhage Algorithm

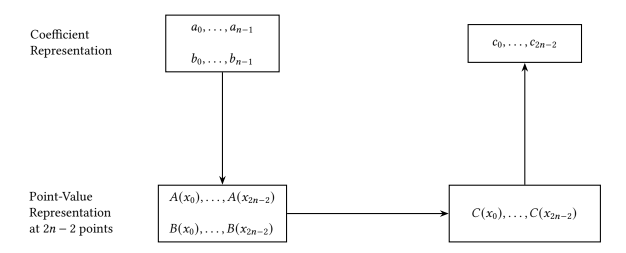
Before diving into the algorithm first let's consider how many ways we can represent a polynomial. Often changing the representation helps solving the problem in less time.

- Coefficients: We can represent a polynomial by giving the array of all its coefficient.
- Point-Value Pairs: We can evaluate the polynomial in distinct *n* points and give all the point-value pairs. This also uniquely represents a polynomial since there is exactly one polynomial of degree n-1 which passes through all these points.

Theorem 3.2.1

Given *n* distinct points $(x_0, y_0), \dots, (x_{n-1}, y_{n-1})$ in \mathbb{R}^2 there is an unique (n-1)-degree polynomial P(x) such that $P(x_i) = y_i \text{ for all } i \in \llbracket n-1 \rrbracket$

Since we want to find the polynomial C(x) = A(x)B(x) and C(x) has degree 2n-2, we will evaluate the polynomials A(x) and B(x) in 2n-1 distinct points. So we will have the algorithm like this:



3.2.1 Finding Evaluations of Multiplied Polynomial

Suppose we were given A(x) and B(x) evaluated at 2n-1 distinct points x_0, \ldots, x_{2n-2} . Then we can get C(x) evaluated at x_0, \ldots, x_{2n-2} by

$$C(x_i) = A(x_i)B(x_i) \ \forall \ i \in [2n-2]$$

Since there are O(n) many points and for each point it takes constant time to multiply we can find evaluations of C at x_0, \ldots, x_{2n-2} in O(n) time.

3.2.2 Evaluation of a Polynomial at Points

Question 3.1

Suppose there is only one point, x_0 . Can we evaluate a n-1 degree polynomial $A(x) = \sum_{i=0}^{n-1} a_i x^i$ at x_0 efficiently?

We can rewrite A(x) as

$$A(x) = a_0 + x(a_1 + x(a_2 + x(a_3 + \cdots + x(a_{n-1} + x(a_n)) \cdots)))$$

In this represent it is clear that we have to do n additions and n multiplications to find $A(x_0)$. Hence we can evaluate a n-1 degree polynomial at a point in O(n) time

But we have O(n) points. And if each point takes O(n) time to find the evaluation of the polynomial then again it will take total $O(n^2)$ time. We are back to square one. So instead we will evaluate the polynomial in some special points and we will evaluate in all of them in $O(n \log n)$ time. So now the problem we will discuss now is to find some special n points where we can evaluate a n-1-degree polynomial in $O(n \log n)$ time.

Idea: Evaluate at roots of unity and use Fast Fourier Transform

Assume n is a power of 2. NWe have the polynomial $A(x) = \sum_{i=0}^{n-1} a_i x^i$. So now consider the following two polynomials

$$A^{0}(x) = a_{0} + a_{2}x + a_{4}x^{2} + \dots + a_{n-2}x^{\frac{n}{2}-1} \qquad A^{1}(x) = a_{1} + a_{3}x + a_{5}x^{2} + \dots + a_{n-1}x^{\frac{n}{2}-1}$$

Certainly we have

mials

$$A(x) = A^{0}(x^{2}) + xA^{1}(x^{2})$$

Hence we can get A(1) and A(-1) by

$$A(1) = A^{0}(1) + A^{1}(1)$$
 $A(-1) = A^{0}(1) - A^{1}(1)$

Hence like this by evaluating two $\frac{n}{2} - 1$ degree polynomials at one point we get evaluation of A at two points. More generally for any $y \ge 0$ we have

$$A(\sqrt{y}) = A^{0}(y) + \sqrt{y}A^{1}(y)$$
 $A(-\sqrt{y}) = A^{0}(y) - \sqrt{y}A^{1}(y)$

So by recursing like this evaluating at 1, -1 we can get evaluations of A at n^{th} roots of unity.

Let

$$\omega_n^k = n^{th}$$
 root of unity for $k \in [[n-1]] = e^{i\frac{k}{n}2\pi} = \cos\left(\frac{k}{n}2\pi\right) + i\sin s\left(\frac{k}{n}2\pi\right)$

Hence we have

$$\begin{split} A\left(\omega_{n}^{k}\right) &= A^{0}\left(\omega_{n}^{2k}\right) + \omega_{n}^{k}A^{1}\left(\omega_{n}^{2k}\right) = A^{0}\left(\omega_{\frac{n}{2}}^{k}\right) + \omega_{n}^{k}A^{1}\left(\omega_{\frac{n}{2}}^{k}\right) \\ A\left(-\omega_{n}^{k}\right) &= A\left(\omega_{n}^{\frac{n}{2}+k}\right) = A^{0}\left(\omega_{n}^{2k}\right) - \omega_{n}^{k}A^{1}\left(\omega_{n}^{2k}\right) = A^{0}\left(\omega_{\frac{n}{2}}^{k}\right) - \omega_{n}^{k}A^{1}\left(\omega_{\frac{n}{2}}^{k}\right) \end{split}$$

Hence now we will solve the following problem:

RECURSIVE-DFT

Input: (a_0, \ldots, a_{n-1}) representing (n-1)-degree polynomial $A(x) = \sum_{i=0}^{n-1} a_i x^i$

Question: Find the evaluations of the polynomial A(x) in all n^{th} roots of unity

Since A^0 and A^1 have degree $\frac{n}{2} - 1$ we can use recursion. Hence the algorithm is

```
Algorithm 9: Recursive-DFT(A)
```

```
Input: A = (a_0, ..., a_{n-1}) such that A(x) = a_0 + a_1 x + \cdots + a_{n-1} x^{n-1}
Output: A(x) evaluated at n^{th} roots of unity \omega_n^k for all k \in [n-1]
 1 begin
               if n == 1 then
 2
                 return A[0]
 3
               A^0 \leftarrow (A[0], A[2], \dots, A[n-2])

A^1 \leftarrow (A[1], A[3], \dots, A[n-1])
 4
               Y^0 \leftarrow \text{Recursive-DFT}(A^0)
               Y^1 \leftarrow \text{Recursive-DFT}(A^1)
 7
               for k = 0 to \frac{n}{2} - 1 do
                                                                                                                                                                                             // A\left(\omega_{n}^{k}\right) = A^{0}\left(\omega_{\frac{n}{2}}^{k}\right) + \omega_{n}^{k}A^{1}\left(\omega_{\frac{n}{2}}^{k}\right)
// A\left(-\omega_{n}^{k}\right) = A^{0}\left(\omega_{\frac{n}{2}}^{k}\right) - \omega_{n}^{k}A^{1}\left(\omega_{\frac{n}{2}}^{k}\right)
                       Y[k] \longleftarrow Y^0[k] + \omega_n^k Y^1[k]
 9
                       Y\left[k+\frac{n}{2}\right] \longleftarrow Y^{0}[k] - \omega_{n}^{\frac{n}{2}+k}Y^{1}[k]
10
               return Y
11
```

Time Complexity: $T(n) = 2T(\frac{n}{2}) + O(n) = O(n \log n)$.

Therefore we can evaluate a n-1 degree polynomial in all the n^{th} roots of unity in $O(n \log n)$ time. Hence with this algorithm we will get evaluations of the polynomial C(x) = A(x)B(x) in all the $2n^{th}$ roots of unity. Now we need to interpolate the polynomial C(x) from its evaluations. We will describe the process in the next subsection.

3.2.3 Interpolation from Evaluations at Roots of Unity

In this section we will show how to interpolate a n-1 degree polynomial from evaluations at all n^{th} roots of unity. Previously we had

$$\underbrace{\begin{bmatrix} C\left(\omega_{n}^{0}\right) \\ C\left(\omega_{n}^{1}\right) \\ C\left(\omega_{n}^{2}\right) \\ \vdots \\ C\left(\omega_{n}^{n-1}\right) \end{bmatrix}}_{Y} = \underbrace{\begin{bmatrix} 1 & \omega_{n}^{0} & \omega^{0\cdot2} & \cdots & \omega^{0\cdot(n-1)} \\ 1 & \omega_{n}^{1} & \omega^{1\cdot2} & \cdots & \omega^{1\cdot(n-1)} \\ 1 & \omega_{n}^{2} & \omega^{2\cdot2} & \cdots & \omega^{2\cdot(n-1)} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & \omega_{n}^{n-1} & \omega^{(n-1)\cdot2} & \cdots & \omega^{(n-1)\cdot(n-1)} \end{bmatrix}}_{V= \text{ Vandermonde Matrix}} \underbrace{\begin{bmatrix} c_{0} \\ c_{1} \\ c_{2} \\ \vdots \\ c_{n-1} \end{bmatrix}}_{C}$$

Now vandermonde matrix is invertible since all the n^{th} roots are distinct. Therefore $C = V^{-1}Y$. But we can not do a matrix inversion to interpolate the polynomial because that will take $O(n^2)$ time. Instead we have this beautiful result:

Lemma 3.2.2
$$\left(V^{-1}\right)_{j,k} = \frac{1}{n} \omega_n^{-jk} \text{ for all } 0 \leq j,k \leq n-1$$

Proof: Consider the matrix $n \times n$ matrix T such that $(T)_{j,k} = \frac{1}{n}\omega_n^{-jk}$. Now we will show VT = I This will confirm that $V^{-1} = T$. Now

$$\sum_{k=0}^{n-1} (V)_{i,j} (T)_{j,k} = \sum_{k=0}^{n-1} \omega_n^{ij} \times \frac{1}{n} \omega_n^{-jk} = \frac{1}{n} \sum_{k=0}^{n-1} \left(\omega_n^{i-k} \right)^j = \begin{cases} \frac{1}{n} \sum_{k=0}^{n-1} 1 = 1 & \text{when } i = k \\ \frac{1}{n} \frac{1 - \omega_n^n}{1 - \omega} = 0 & \text{when } i \neq k \end{cases}$$

Hence in VT there are 1's on the diagonal and rest of the locations are 0. Hence VT = I. So $V^{-1} = T$.

Hence we can see the inverse of the vandermonde matrix is also a vandermode matrix with a scaling factor. We will denote $y_i = C\left(\omega_n^i\right)$ for $i \in [n-1]$ since these values are given to us some how and we have to find the corresponding polynomial. Therefore we have

$$\underbrace{ \begin{bmatrix} c_0 \\ c_1 \\ c_2 \\ \vdots \\ c_{n-1} \end{bmatrix} }_{C} = \underbrace{ \frac{1}{n} \begin{bmatrix} 1 & 1 & 1 & \cdots & 1 \\ 1 & \omega_n^{-1} & \omega^{-1 \cdot 2} & \cdots & \omega^{-1 \cdot (n-1)} \\ 1 & \omega_n^{-2} & \omega^{-2 \cdot 2} & \cdots & \omega^{-2 \cdot (n-1)} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & \omega_n^{-(n-1)} & \omega^{-(n-1) \cdot 2} & \cdots & \omega^{-(n-1) \cdot (n-1)} \end{bmatrix}}_{V^{-1}} \underbrace{ \begin{bmatrix} y_0 \\ y_1 \\ y_2 \\ \vdots \\ y_{n-1} \end{bmatrix}}_{Y}$$

Observation.
$$nc_j = y_0 + y_1 \omega_n^{-j} + y_2 \omega_n^{-2j} + \dots + y_{n-1} \omega_n^{-(n-1)j}$$
 for all $j \in [[n-1]]$

We can also see this situation as we have the polynomial $Y(x) = y_0 + y_1x + y_2x^2 + \cdots + y_{n-1}x^{n-1}$ and c_j is just Y(x) evaluated as $\omega_n^{-j} = \omega_n^{n-j}$ multiplied by n. Hence we just reindex the n^{th} roots of unity and evaluate $Y(n^{th})$ roots of unity in $O(n \log n)$ time using the algorithm described in subsection 3.2.2

Dynamic Programming

Definition 4.1: Dynamic Programming

Dynamic Programming has 3 components:

- 1. Optimal Substructure: Reduce problem to smaller independent problems
- 2. Recursion: Use recursion to solve the problems by solving smaller independent problems
- 3. Table Filling: Use a table to store the result to solved smaller independent problems.

4.1 Longest Increasing Subsequence

Longest Increasing Subsequence

Input: Sequence of distinct integers $A = (a_1, ..., a_n)$

Question: Given an array of distinct integers find the longest increasing subsequence i.e. return maximum

size set $S \subseteq [n]$ such that $\forall i, j \in S, i < j \implies a_i < a_j$

4.1.1 $O(n^2)$ Time Algorithm

Given $A = (a_1, ..., a_n)$ first we will create a *n*-length array where i^{th} entry stores the length and longest increasing subsequence ending at a_i . Certainly we have the following recursion relation

$$LIS(k) = 1 + \max_{j < k, \ a_j < a_k} \{LIS(j)\}$$

since if a subsequence $S \subseteq [n]$ is the longest increasing subsequence ending at a_k then certainly $S - \{k\}$ is the longest increasing subsequence which ends at $a_j < a_k$ for some j < k. Hence in the table we start with 1st position and using the recursion relation we fill the table from left. And after the table is filled we look for which entry of the table has maximum length. So the algorithm will be following:

Algorithm 10: LIS (\overline{A})

```
Input: Sequence of distinct integers A = (a_1, ..., a_n)

Output: Maximum size set S \subseteq [n] such that \forall i, j \in S, i < j \implies a_i < a_j.

1 begin

2 | Create an array T of length n

3 | for i \in [n] do

4 | T[i][1] \leftarrow 1 + \max\{T[j][1] : j < k, a_j < a_k\} // Finds LIS[i]

5 | T[i][2] \leftarrow T[T[i][1] - 1][2]

6 | Index \leftarrow \max\{T[j][1] : j \in [n]\}

7 | return T[Index]
```

Time Complexity: For each iteration of the loop it takes O(n) time to find LIS[i]. Hence the time complexity of this algorithm is $O(n^2)$.

4.1.2 $O(n \log n)$ Time Algorithm

In the following algorithm we update the longest increasing sequence every time we see a new element of the given sequence. At any time we keep the best available sequence.

Idea. We can make an increasing subsequence longer by picking the smallest number for position k so that there is an increasing subsequence of length k. Doing this we can maximize the length of the subsequence.

Theorem 4.1.1

Is $S \subseteq A$ is the longest increasing subsequence of length t then for any $k \in [t]$ the number S(k) is the smallest number in subarray of A starting at first and ending at S(k) such that there is an increasing subsequence of length k ending at S(k).

Proof: Suppose $\exists k \in [t]$ such that k is the smallest number in [t] such that S(k) is not the smallest number to satisfy the condition. Now denote the subarray of A starting at first and ending at S(k) by A_k . Now let $x \in A_k$ be the smallest number in A_k such that there is an increasing subsequence of length k ending at k. Certainly k is the smallest index which does not satisfy the given condition, k is the smallest number in k such that there is an increasing subsequence of length k ending at k is the smallest number in k such that there is an increasing subsequence of length k ending at k is the smallest number in k such that there is an increasing subsequence of length k ending at k is the smallest number in k such that there is an increasing subsequence of length k ending at k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that there is an increasing subsequence of k and has length k is the smallest number in k such that k is not the smallest number in k in k such that k is not the smallest number in k such that k is not the smallest number in k in k such that k is not that k is not that k is not that k in k i

So we will construct an increasing subsequence by gradually where each step this property is followed, i.e. at each step we will ensure that the sequence built at some time have the above propert. So now we describe the algorithm.

```
Algorithm 11: QUICKLIS(A)
```

```
Input: Sequence of distinct integers A = (a_1, ..., a_n)
  Output: Maximum size set S \subseteq [n] such that \forall i, j \in S, i < j \implies a_i < a_j.
1 begin
      Create an array T of length n with all entries 0
2
       Create an array M of length n
3
      for i = 1, ..., n do
4
       M[i] \longleftarrow \infty
5
      for i = 1, ..., n do
          k \leftarrow Find smallest index i such that M[k] > a_i using BINARY-SEARCH
7
          M[k] \longleftarrow i
          T[i] \longleftarrow M[k-1]
                                       // Pointer to the previous element of the sequence
      l \leftarrow Largest l such that M[l] is finite
10
11
       Create an array S of length l
      for i = 1, ..., 1 do
12
          if i = l then
13
              S[l] \longleftarrow M[l]
14
              Continue
15
          S[i] \leftarrow T[S[i+1]] // T[S[i+1]] is pointer to previous value of sequence
16
      return (l, S)
17
```

Time Complexity: To create the arrays and the first for loop takes O(n) time. In each iteration of the for loop at line 6 it takes $O(\log n)$ time to find k and rest of the operations in the loop takes constant time. So the for loop takes $O(n \log n)$ time. Then To find l and creating S it takes O(n) time. Then in the for loop at line 12 in each iteration it takes constant time. So the for loop at line 12 takes in total O(n) time. Therefore the algorithm takes $O(n \log n)$ time.

We will do the proof of correctness of the algorithm now.

Lemma 4.1.2

For any index M[k] is non increasing

Proof: Every time we change a value of M[k] we replace by something smaller. So M[k] is non increasing.

We denote the state of array M at i^{th} iteration by M^i . Then we have the following lemma:

Lemma 4.1.3

At any time $i, M^i[1] \le M^i[2] \le \cdots \le M^i[n]$

Proof: We will prove this by induction on i. The base case follows naturally. Now for i^{th} iteration suppose $M^i[k]$ is replaced by x_i . Then we know $\forall j < k$ we have $M^i[j] \leq x_i$. By inductive hypothesis at time t-1 we have M as an increasing sequence. Now before replacing $M^i[k] \leq M^i[k+1] \leq \cdots M^i[n]$. Now by Lemma 4.1.2 $M^i[k]$ is nonincreasing. So So we still have $M^i[1] \leq \cdots M^i[k-1] \leq x_i \leq M^i[k+1] \leq \cdots \leq M^i[n]$. Hence bt mathematical induction it holds.

Now suppose at i^{th} iteration k_i is largest such that $M^i[k_i] < \infty$. Then S^i denote the set constructed like the way we constructed at line 12–16 in the algorithm i.e.

$$S^{i}[k_{i}] = M^{i}[k_{i}]$$
 and $S^{i}[j] = T[S^{i}[j+1]] \quad \forall j \in [k_{i}-1]$

Lemma 4 1 4

After any i^{th} iteration, for $k \in [n]$ if $M^i[k] < \infty$ then $S^i[k]$ stores the smallest value in x_1, \ldots, x_i such that there is an increasing subsequence of size k that ends in $S^i[k]$.

Proof: We will induction on *i*. Base case: This is true after first iteration since only $M^1[1] < \infty$. So this naturally follows. Suppose this is true after *i* iterations. Now at $(i+1)^{th}$ iteration suppose *t* be the smallest index such that $M^i[t] > x_{i+1}$. Then we have

$$M^{i}[1], \dots, M^{i}[t-1] < x_{i+1} < M^{i}[k], \dots, M^{i}[n] \implies S^{i}[1], \dots, S^{i}[t-1] < x_{i+1} < S^{i}[k], \dots, S^{i}[k_{i}]$$

Now for $k \le t-1$ it is true by the inductive hypothesis. For k > t and if $M^{i+1}[k] < \infty$ then $S^{i+1}[k]$ is the smallest value in x_1, \ldots, x_{i+1} such that there is an increasing subsequence of size k that ends in $S^{i+1}[k]$ since this was true for i^{th} iteration. Now only the case when k = t is remaining. If $S^{i+1}[k]$ is not the smallest value in x_1, \ldots, x_{i+1} to have an increasing subsequence of size k ending at $S^{i+1}[k]$ then let x_j was the smallest value to satisfy this condition where j < i+1. Then naturally $x_j < x_{i+1}$. Then $M^i[t] \le x_j < x_{i+1}$. But we t was the smallest number such that $M^i[t] > x_{i+1}$. Hence contradiction. Therefore $S^i[k]$ is the smallest value in x_1, \ldots, x_{i+1} to have an increasing subsequence of size k ending at $S^{i+1}[k]$.

Therefore by mathematical induction this is true for all iterations.

Theorem 4.1.5

S is the longest increasing subsequence of *A*.

Proof: After the n^{th} iteration $S^n = S$ and $k_n = l$. Hence by Lemma 4.1.4 we can say for all $k \in [l]$, S[k] is the smallest number such that there is an increasing sequence of length k ending at S[k]. Now we want to show that this increasing sequence is the longest increasing subsequence of A. Suppose S is not the longest increasing subsequence. Let T be the longest increasing subsequence of length t. Then suppose $j \leq l$ be the smallest index such that $S[j] \neq T[j]$. Now S[j] is the smallest number in x_1, \ldots, x_n such that there is an increasing subsequence of length j ending at S[j]. Hence we have S[j] < T[j]. Now for all i < j we have S[i] = T[i]. Then we form this new subsequence $\hat{T} = \{T[1], T[2], \ldots, T[j-1], S[j], T[j], \ldots, T[t]\}$. Certainly \hat{T} has length t+1 and it is also an increasing subsequence. But this contradicts the maximality condition of T. Hence S is indeed the longest increasing subsequence.

4.2 Optimal Binary Search Tree

Optimal BST

Input: A sorted array $A = (a_1, ..., a_n)$ of search keys and an array of their probability distributions P =

 $(p(a_1),\ldots,p(a_n))$

Question: Given array of keys A and their probabilities the probability of accessing a_i is $p(a_i)$ then return a

binary tree with the minimum cost where for any binary tree T, $Cost(T) = \sum_{i=1}^{n} p(a_i) \cdot height_T(a_i)$.

So let T be the optimal binary search tree with a_k as its root for some $k \in [n]$. Let T_l and T_r denote the tree rooted at the left child and right child of a_k in T respectively. Then:

$$\mathrm{Cost}(T) = p_k + \sum_{i < k} p_i \left(1 + height_{T_l}(a_i) \right) + \sum_{i > k} p_i \left(1 + height_{T_r}(a_i) \right) = \sum_{i = 1}^n p_i + \underbrace{\sum_{i < k} p_i \cdot height_{T_l}(a_i)}_{\mathrm{Cost}(T_l)} + \underbrace{\sum_{i > k} p_i \cdot height_{T_l}(a_i)}_{\mathrm{Cost}(T_r)}$$

We will use the use of notion in general OPTCost(i,k) = Cost(T_i^k) where T_i^k is the optimal binary tree of the subarray A[i...k] for any $i \le k \le n$. Therefore we arrive at the following recurrence relation

$$\mathsf{OPTCost}(i,k) = \begin{cases} 0 & \text{when } i > k \\ \sum\limits_{j=i}^k p(a_j) + \min_{i \leq r \leq k} \{\mathsf{OPTCost}(i,r-1) + \mathsf{OPTCost}(r+1,k)\} & \text{otherwise} \end{cases}$$

So the algorithm for constructing the optimal binary search tree is following:

```
Algorithm 12: OptimalBST(A, P)
```

```
Input: A sorted array A = (a_1, ..., a_n) of search keys and an array of their probability distributions P = (p(a_1), ..., p(a_n))
```

Output: Binary Tree T with the minimum search cost, $Cost(T) = \sum_{i=1}^{n} p(a_i) \cdot height_T(a_i)$

```
1 begin
       for i = 1, ..., n do
2
        for d = 2, ..., n do
           for i \in [n+1-d] do
               minval \leftarrow 0
               for k = i + 1, ..., i + d - 2 do
 7
                   newval \leftarrow OPTCost[i, k-1][1] + OPTCost[k+1, i+d-1][1]
 8
                   if minval > newval then
 9
                       minval \leftarrow newval
10
                     Index \longleftarrow k
11
              OPTCost[i, i+d-1] \leftarrow \left( minval + \sum_{k=1}^{i+d-1} p(a_k), k \right)
12
               a_k.left \leftarrow \text{OPTCost}[i, k-1][2]

a_k.right \leftarrow \text{OPTCost}[k+1, i+d-1][2]
13
14
       return OPTCost[1, n]
15
```

Time Complexity: To two for loops ar line 4 and line 5 takes $O(n^2)$ many iterations. Now the inner most for loop at line 7 runs O(n) iterations where in each iterations it takes constant runtime. So the total running time of the algorithm is $O(n^3)$.

Greedy Algorithm

5.1 Maximal Matching

MAXIMAL MATCHING

Input: Graph G = (V, E)

Question: Find a maximal matching $M \subseteq E$ of G

Before diving into the algorithm to find a matching or maximal matching we first define what is a matching.

Definition 5.1.1: Matching

For a graph G = (V, E) a matching $M \subseteq E$ is a set of edges such that no two edges in M are incident on same vertex.

Definition 5.1.2: Maximal Matching

For a graph G = (V, E) a matching $M \subseteq E$ is maximal if it cannot be extended and still by adding an edge.

There is also a maximum matching which can be easily understood from the name:

Definition 5.1.3: Maximum Matching

For a graph G = (V, E) a matching $M \subseteq E$ is maximum if it is maximal and has the maximum size among all the maximal matchings.

Idea. The idea is to create a maximal matching we will just go over each edge one by one and check if after adding them to the set M the matching property still holds.

Algorithm 13: MAXIMAL-MATCHING

```
Input: Graph G = (V, E)
Output: Maximal Matching M \subseteq E of G

1 begin
2 | M \longleftarrow \emptyset
3 | Order the edges E = \{e_1, \dots, e_k\} arbitrarily
4 | for e \in E do
5 | if M \cup \{u\} is matching then
6 | M \longleftarrow M \cup \{e\}
7 | return M
```

5.1 Maximal Matching Page 18

Question 5.1

Do we always get the largest possible matching?

Solution: Clearly algorithm output is not optimal always. We get a maximal matching sure. But we don't get a maximum matching always. For example the following graph



If we start from e_1 we get the matching $\{e_1.e_2\}$ which is maximum matching but if we start from e_3 then we get only the maximal matching $\{e_3\}$ which is not maximum.

Since the algorithm output may not be optimal always we can ask the following question

Question 5.2

How large is the matching obtained compared to the maximum matching?

This brings us to the following result:

Theorem 5.1.1

For any graph G let the greedy algorithm obtains the matching M and the maximum matching is M^* . Then

$$|M| \ge \frac{1}{2}|M^{\star}|$$

Proof: Consider an edge $e \in M^*$ but $e \notin M$. Since e wasn't picked in M, $\exists e' \in M \setminus M^*$ such that e and e' are incident on same vertex. Thus define the function $f: M^* \to M$ where

$$f(e) = \begin{cases} e & \text{when } e \in M \\ e' & \text{when } e \in M^* \setminus M \text{ where } e' \in M \setminus M^* \text{ such that } e' \cap e \neq \emptyset \end{cases}$$

Now note that there are at most two edges in M^* that are adjacent to an edge $e' \in M$ which will be mapped to e'. Hence

$$|M \setminus M^{\star}| \ge \frac{1}{2}|M^{\star} \setminus M|$$

Therefore $|f^{-1}(e')| \le 2 \forall e' \in M$. Hence

$$|M^{\star}| = |M \cap M^{\star}| + |M^{\star} \setminus M| \le |M \cap M^{\star}| + 2|M \setminus M^{\star}| \le 2|M|$$

Therefore we have the result $|M| \ge \frac{1}{2}|M^*|$.

Alternate Proof: Let M_1 and M_2 are two matchings. Consider the symmetric difference $M_1 \triangle M_2$. This consists of edges that are in exactly one of M_1 and M_2 . Now in $M \triangle M^*$ we have the following properties:

- (a) Every vertex in $M \triangle M^*$ has degree $\leq 2 \implies$ Each component is a path or an even cycle.
- (b) The edges of M and M^* alternate.

Now we will prove the following property about the connected components of $M \triangle M^*$.

Claim: No connected component is a single edge.

Proof: This is because let e be a connected component. So the two edges e_1, e_2 which are adjacent to e, they are either in both M and M^* or not in M and M^* . The former case is not possible because then e_1, e_2, e are all in either M or M^* which is not possible as they do not satisfy the condition of matching. For the later case since M^* is maximal matching, $e \in M^*$. Then $e \notin M$. That means $e, e_1, e_2 \notin M$ which is not possible since M is also a maximal matching. Therefore no connected component is a single edge.

Therefore every path has length ≥ 2 . Therefore ratio of # edges of M to # edges of M^* in a path is ≤ 2 . And for cycles we have # edges of M = # edges of M^* . So in every connected component C of $M \triangle M^*$ the ratio $\frac{|M^* \cap C|}{|M \cap C|} \leq 2$. Therefore we have

$$\frac{|M^{\star}|}{|M|} = \frac{|M \cap M^{\star}| + \sum\limits_{C} |M^{\star} \cap C|}{|M \cap M^{\star}| + \sum\limits_{C} |M \cap C|} \le 2$$

Hence we have $|M| \ge \frac{1}{2}|M^*|$.

5.2 Huffman Encoding

HUFFMAN CODING

Input: n symbols $A = (a - 1, ..., a_n)$ and their frequencies $P = (f_1, ..., f_n)$ of using symbols

Question: Create a binary encoding such that:

• Prefix Free: The code for one word can not be prefix for another code

• Minimality: Minimize $Cost(b) = \sum_{i=1}^{n} f_i \cdot Len(b(a_i))$ where $b: A \to \{0,1\}^*$ is the binary encoding

Assignment of binary strings can also be scene as placing the symbols in a binary tree where at any node 0 means left child and 1 means right child. Then the first condition implies that there can not be two codes which lies in the same path from the root to a leaf. I.e. it means that all the codes have to be in the leaves. Then the length of the binary coding for a symbol is the height of the symbol in the binary tree.

We can think the frequencies as the probability of appearing for a letter. We denote the probability of appearing of the letter a_i by $p(a_i) := \frac{f_i}{\sum_{i} f_i}$. So the we can see the updated cost function

$$Cost(b) = \sum_{i=1}^{n} p(a_i) \cdot Len(b(a_i))$$

And from now on we will see the frequencies as probabilities and cost function like this

5.2.1 Optimal Binary Encoding Tree Properties

Then our goal is to finding a binary tree with minimum cost where all the symbols are at the leaves. We have the following which establish the optimality of Huffman encoding over all prefix encodings where each symbol is assigned a unique string of bits.

Lemma 5.2.1

In the optimal encoding tree least frequent element has maximum height.

Proof: Suppose that is not the case. Let T be the optimal encoding tree and let the least frequent element x is at height h_1 and the element with the maximum height is y with height h_2 and we have $h_1 < h_2$. Then we construct a new encoding tree T' where we swap the positions of x and y. So in T' height of y is h_1 and height of x is h_2 . Then

$$Cost(T) - Cost(T') = (p(x)h_1 + p(y)h_2) - (p(x)h_2 + p(y)h_1) = (p(x) - p(y))(h_1 - h_2)$$

Since p(x) < p(y) and $h_1 < h_2$ we have Cost(T) - Cost(T') > 0. But that is not possible since T is the optimal encoding tree. So T should have the minimum cost. Hence contradiction. x has the maximum height.

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Lemma 5.2.2

The optimal encoding binary tree must be complete binary tree. (i.e. every non-leaf node has exactly 2 children)

Proof: Suppose T be the optimal binary tree and there is a non-leaf node r which has only one child at height h. By Lemma 5.2.1 the least frequent element x has the maximum height, h_m .

Then consider the new tree \hat{T} where we place the least frequent element at height h and make it the second child of the node r. Then

$$Cost(T) - Cost(\hat{T}) = p(x)h_m - p(x)h = p(x)(h_m - h) > 0$$

But this is not possible as *T* is the optimal binary tree and it has the minimal cost. Hence contradiction. Therefore the optimal encoding binary tree must be a complete binary tree.

Lemma 5.2.3

There is an optimal binary encoding tree such that the least frequent element and the second least frequent element are siblings at the maximum height.

Proof: Let *T* be optimal binary encoding tree. Suppose x, y are the least frequent element and the second least frequent element. And suppose b, c be two siblings at the maximum height of the tree (There may be many such siblings, and if so pick any such pair.). If $\{x, y\} = \{b, c\}$ we are done. So suppose not. Let the frequencies of x, y, b, c are respectively p(x), p(y), p(b), p(c) and heights of x, y, b are h_x, h_y and h respectively. WLOG assume $p(x) \le p(y)$ and $p(b) \le p(c)$.

Now since we know x, y have the smallest frequencies we have $p(x) \le p(b)$ and $p(y) \le p(c)$. And since b, c have the maximum height we have $h_x, hy \ge h$. So we switch the position of x with b to form the new tree T'. And from T' we swap the positions fo y and c to form a new tree T''.

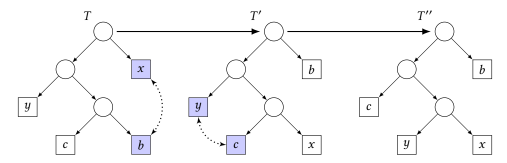


Figure 5.1: Showing that the lowest probability nodes are siblings at the tree's lowest level.

Now we will calculate how the cost changes as we go from T to T' and T' to T''. First check for $T \to T'$. Almost all the nodes contribute the same except x, b. So we have

$$Cost(T) - Cost(T') = (h_x \cdot p(x) + h \cdot p(b)) - (h_x \cdot p(b) + h \cdot p(x)) = (p(b) - p(x))(h - h_x) \ge 0$$

Therefore swapping x and b does not increase the cost and since T is the optimal binary encoding tree the cost doesn't decrease either. Therefore the costs are equal. Hence T' is also an optimal tree.

Similarly we calculate cost for going from T' to T'' we have

$$\mathsf{Cost}(T') - \mathsf{Cost}(T'') = (h_y \cdot p(y) + h \cdot p(c)) - (h_y \cdot p(c) + h \cdot p(y)) = (p(c) - p(y))(h - h_y) \ge 0$$

Therefore swapping y and c also does not increase the cost and since T' is the optimal binary encoding tree the cost doesn't decrease either. Therefore the costs are equal. Hence T'' is also an optimal tree. Hence T'' is the optimal tree where the least frequent element and second last frequent element are siblings.

By the Lemma 5.2.2 and Lemma 5.2.3 we have that the least frequent element and the second least frequent element are siblings and they have the maximum height.

Observation. The cost of the trees T_n and T_{n-1} differ only by the fixed term p(z) = p(x) + p(y) which does not depend on the tree's structure. Therefore minimizing the cost for T_n is equivalent to minimizing the cost of T_{n-1} .

Theorem 5.2.4

Given an instance with symbols \mathcal{I} :

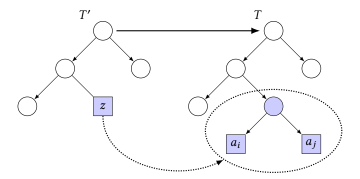
$$a_1,$$
 $a_2,$ $\cdots,$ $a_i,$ $\cdots,$ $a_j,$ $\cdots,$ a_n with probabilities $p(a_1),$ $p(a_2),$ $\cdots,$ $p(a_i),$ $\cdots,$ $p(a_j),$ $\cdots,$ $p(a_n)$

such that a_i , a_j are the least frequent and second least frequent elements respectively. Consider the instance with n-1 symbols \mathcal{I}' :

$$a_1,$$
 $a_2,$ $\cdots,$ $a_{i-1},$ $a_{i+1},$ $\cdots,$ $a_{j-1},$ $a_{j+1},$ $\cdots,$ $a_n,$ z $p(a_1),$ $p(a_2),$ $\cdots,$ $p(a_{i-1}),$ $p(a_{i+1})$ $\cdots,$ $p(a_{j-1}),$ $p(a_{j+1}),$ $\cdots,$ $p(a_n),$ $p(a_i)+p(a_j)$

Let T' be the optimal tree for this instance I'. Then there is an optimal tree for the original instance I obtained from T' by replacing the leaf of b by an internal node with children a_i and a_j .

Proof: We will prove this by contradiction. Suppose \hat{T} is optimal for I. Then $Cost(\hat{T}) < Cost(T)$. In \hat{T} we know a_i and a_j are siblings by Lemma 5.2.3. Now consider \hat{T}' for instance I' where we merge a_i, a_j leaves and their parent into a leaf for symbol z.



Then

$$\operatorname{Cost}(\hat{T}') = \operatorname{Cost}(\hat{T}) - p(a_i) - p(a_j) < \operatorname{Cost}(T) - p(a_i) - p(a_j) = \operatorname{Cost}(T')$$

This contradicts the fact that T' is optimal binary encoding tree for I'. Hence T is optimal.

5.2.2 Algorithm

Idea: We are going to build the tree up from the leaf level. We will take two characters x, y, and "merge" them into a single character, z, which then replaces x and y in the alphabet. The character z will have probability equal to the sum of x and y's probabilities. Then we continue recursively building the code on the new alphabet, which has one fewer character.

Since we always need the least frequent element and the second least frequent element we have to use the data structure called Min-Priority Queue. So the following algorithm uses a Min-Priority Queue Q keyed on the probabilities to identify the two least frequent objects.

Time Complexity: To create the priority queue it takes O(n) time in line 4-5. Then for each iteration of the for loop in line 6 the EXTRACT-MIN operation takes $O(\log n)$ time and then to insert an element it also takes $O(\log n)$ time. Hence each iteration takes $O(\log n)$ time. Since the for loop has n-1=O(n) many iterations the running time for the algorithm is $O(n \log n)$.

Remark: We can reduce the running time to $O(n \log \log n)$ by replacing the binary min-heap with a van Emde Boas tree.

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Algorithm 14: Huffman-Encoding(A, P)

```
Input: Set of n symbols A = \{a_1, \ldots, a_n\} and their probabilities P = \{p_1, \ldots, p_n\}
   Output: Optimal Binary Encoding b: A \to \{0,1\}^* for A with minimum Cost(b) = \sum_{i=1}^n p(a_i) \cdot Len(b(a_i)).
1 begin
2
       n \longleftarrow |A|
        Q \leftarrow Min-Priority Queue
3
       for x \in A do
4
         | Insert(Q, x)
        for i = 1, ... n - 1 do
6
            z \leftarrow New internal tree node
            x \leftarrow \text{Extract-Min}(Q), y \leftarrow \text{Extract-Min}(Q)
            left[z] \leftarrow x, right[z] \leftarrow y
            p(z) \longleftarrow p(x) + p(y)
10
            Insert(Q, z)
11
       return Last element left in Q as root
12
```

Theorem 5.2.5 Correctness of Huffman's Algorithm

The above Huffman's algorithm produces an optimal prefix code tree

Proof: We will prove this by induction on n, the number of symbols. For base case n = 1. There is only one tree possible. For n = k we know that by Lemma 5.2.3 and Lemma 5.2.1 that the two symbols x and y of lowest probabilities are siblings and they have the maximum height. Huffman's algorithm replaces these nodes by a character z whose probability is the sum of their probabilities. Now we have 1 less symbols. So by inductive hypothesis Huffman's algorithm computes the optimal binary encoding tree for the k-1 symbols. Call it T_{n-1} . Then the algorithm replaces z with a parent node with children x and y which results in a tree T_n whose cost is higher by a fixed amount p(z) = p(x) + p(y). Now since T_{n-1} is optimal by Theorem 5.2.4 we have T_n is also optimal.

5.3 Matroids

Definition 5.3.1: Matroid

A matroid M = (E, I) has a ground set E and a collection I of subsets of E called the *Independent Sets* st

- 1. Downward Closure: If $Y \in \mathcal{I}$ then $\forall X \subseteq Y, X \in \mathcal{I}$.
- 2. Exchange Property: If $X, Y \in I$, |X| < |Y| then $\exists e \in Y X$ such that $X \cup \{e\}$ also written as $X + e \in I$

An element $x \in E$ extends $A \in I$ if $A \cup \{x\} \in I$. And A is maximal if no element can extend A.

Lemma 5.3.1

If A, B are maximal independent set, then |A| = |B| i.e. all maximal independent sets are also maximum

Proof: Suppose $|A| \neq |B|$. WLOG assume |A| > |B|. Then by the exchange property $\exists \ e \in A - B$ such that $B \cup \{e\} \in I$. But we assumed that B is maximal independent set. Hence contradiction. We have |A| = |B|.

```
Base: Maximal Independent sets are called bases. 
Rank of S \in I: max{|X|: X \subseteq S, X \in I}
```

Rank of a Matroid: Size of the base.

Span of $S \in I$: $\{e \in E : rank(S) = rank(S+e)\}$

5.3.1 Examples of Matroid

• Uniform Matroid: Given $E = \{e_1, \dots, e_n\}$, and $k \in \mathbb{Z}_0$ take $I = \{S \subseteq E : |S| \le k\}$

Lemma 5.3.2

M = (E, I) defined as above is a matroid

Proof:

- 1 Downward Closure: $A \in I$, $B \subseteq A \implies |B| \le k \implies B \in I$
- (2) Exchange Property: $A, B \in I$, $|B| < |A| \le k \implies |B| < k \implies \forall e \in A B$, $|B \cup \{e\}| \le k \implies B \cup \{e\} \in I$

Therefore M is a matroid

• **Partition Matroid:** Given E, $\{P_1, \ldots, P_l\}$ such that $E = \bigcap_{i=1}^l P_i$ and $k_1, \ldots, k_l \in \mathbb{Z}_0$ then take

$$I = \{ S \subseteq E \colon \forall \ k \in [l], \ |S \cap P_j| \le k_j \}$$

Lemma 5.3.3

M = (E, I) defined as above is a matroid

Proof:

- (1) Downward Closure: $A \in I$, $B \subseteq A \implies \forall j \in [l] |B \cap P_j| \le |A \cap P_j| \le k_j \implies B \in I$
- ② Exchange Property: $A, B \in I$, $|B| < |A| \implies \exists j \in [l]$, $|B \cap P_j| < |A \cap P_j| \le k_j \implies e \in (A \cap P_j) (B \cap P_j)$, $|(B \cup \{e\}) \cap P_j| = |B \cap P_j| + 1 \le k \implies B \cup \{e\} \in I$

Therefore M is a matroid

• Laminar Matroid: Given E, $\mathcal{L} = \{L_1, ..., L_l\}$ such that $\forall i, j \in [l]$, either $L_i \subseteq L_j$ or $L_i \supseteq L_j$ or $L_i \cap L_j = \emptyset$ and also given $k_1, ..., k_l \in \mathbb{Z}_0$. Then take

$$I = \{S \subseteq E : \forall j \in [l], |S \cap L_i| \le k_i\}$$

For any $L \in \mathcal{L}$ we denote k(L) be the given number corresponding to L.

Lemma 5.3.4

M = (E, I) defined as above is a matroid

Proof:

- 1 Downward Closure: $A \in I$, $B \subseteq A \implies \forall j \in [l] |B \cap L_j| \le |A \cap L_j| \le k_j \implies B \in I$
- ② Exchange Property: Let $A, B \in \mathcal{I}$ with |B| < |A|. If there exists $e \in A \setminus B$ such that $e \notin L$ for any $L \in \mathcal{L}$, then $|(B+e) \cap L| = |B \cap L| \le k(L)$ for any $L \in \mathcal{L}$.

Hence assume that for each $e \in A \setminus B$ there exists $L \in \mathcal{L}$ with $e \in L$. For each $e \in A \setminus B$, let \mathcal{L}_e be the collection of $L \in \mathcal{L}$ with $e \in L$. For each $e \in A \setminus B$ and any $L \in \mathcal{L} \setminus \mathcal{L}_e$, we have $|(B + e) \cap L| = |B \cap L| \le k(L)$.

Hence it remains to show that there exists $e \in A \setminus B$ such that $|(B+e) \cap L| \leq k(L)$ for any $L \in \mathcal{L}_e$. Note that \mathcal{L}_e is a chain, as \mathcal{L} is a laminar. Let $\mathcal{L}' = \{L_{e_1}, \dots, L_{e_l}\}$ be the collection of inclusion-wise maximal sets in \mathcal{L} such that $|B \cap L_{e_i}| \leq k(L_{e_i})$ with $e_i \in A \setminus B$. Then $L_{e_i} \cap L_{e_j} = \emptyset$. Moreover, |A| > |B| and $|A \cap L_{e_i}| \leq k(L_{e_i})$ imply that $|A \setminus (\cup L_{e_i})| > |B \setminus (\cup L_{e_i})|$. Hence there $\exists e_i$ such that $|A \cap L_{e_i}| > |B \cap L_{e_i}|$.

Now we take a look at the chain \mathcal{L}_{e_i} . For brevity we will use e instead of e_i . So in the chain $\mathcal{L}_e = \{L_1, \ldots, L_n\}$ such that we have

$$L_n \supseteq L_{n-1} \supseteq \cdots \supseteq L_2 \supseteq L_1$$

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Then take $i \in [n]$ to be the largest index such that $|A \cap L_i| \leq |B \cap L_i|$. There will be such index because otherwise we will have $|A| \leq |B|$ which is not possible. Then take $e^* \in (A \cap L_{i+1}) - (L_i \cup B)$. Such an e^* will exist because $|A \cap L_{i+1}| > |A \cap L_{i+1}| \implies A \cap (L_{i+1} - L_i \neq \emptyset)$ and also $A \cap (L_{i+1} - L_i \not\subseteq B \cap (L_{i+1} - L_i))$ because otherwise we will have

$$|A \cap L_{i+1}| = |A \cap (L_{i+1} - L_i)| + |A \cap L_{i+1}| \le |B \cap (L_{i+1} - L_i)| + |B \cap L_i| = |B \cap L_{i+1}|$$

which is not possible. Hence there exists e^* such that $e^* \in (A \cap L_{i+1}) - (L_i \cup B)$. Therefore take $B^* = B \cup \{e^*\}$. Then for all j < i we have $B^* \cap L_j = B \cap L_j$ so we don't have a problem there. Now for all $j \ge i$ we have $|A \cap L_j| > |B \cap L_j|$. Hence now $|B^* \cap L_j| \le |B \cap L_j| + 1 \le |A \cap L_j| \le k(L_j)$. Therefore we have $B^* \in I$. Hence the exchange property follows.

Therefore M is a matroid.

• **Graphic Matroid:** Given a graph G = (V, E) E is the ground set and take

$$I = \{E' \subseteq E \colon E' \text{ is acyclic}\}\$$

Lemma 5.3.5

M = (E, I) defined as above is a matroid

Proof:

- ① Downward Closure: If a set of edges *S* is acyclic then naturally any subset of edges of *S* is also acyclic. Hence downward closure property follows.
- ② Exchange Property: $A, B \in I$, and |B| < |A|. Let G_1, \ldots, G_k are the connected components due to B. For each component G_i , we have $|G_i \cap A| \le |G_i \cap B|$ since each component is a tree and B has maximum number of edges for that component. Then A contains an edge e connecting 2 components G_i and G_j . Then $B \cup \{e\} \in I$.

Therefore M is a matroid

• Linear Matroid: Given a $m \times n$ matrix $M \in \mathbb{Z}^{m \times n}$, E = [n] and take

 $I = \{S \subseteq E : \text{Columns of } M \text{ corresponding to } S \text{ are linearly independent} \}$

Lemma 5.3.6

M = (E, I) defined as above is a matroid

Proof:

- ① Downward Closure: $A \in I$, $B \subseteq A$. Subset of linearly independent set is also linearly independent. Hence $B \in I$.
- ② Exchange Property: $A, B \in \mathcal{I}$, |B| < |A|. Then take span $\langle A \rangle$ over \mathbb{Q} . Now we know a set of integral vectors are linearly independent over integers if and only if they are linearly independent over rationals. Hence $|A| = \dim_{\mathbb{Q}} \langle A \rangle > \dim_{\mathbb{Q}} \langle B \rangle = |B|$. Hence we can extend B by an element $e \in A B$ such that $\langle B \cup \{e\} \rangle = |B| + 1$. Hence $B \cup \{e\} \in \mathcal{I}$.

Therefore M is a matroid

This matroid is also called Metric Matroid.

5.3.2 Finding Max Weight Base

MAX WEIGHT BASE

Input: A matroid M = (E, I) is given as an input as an oracle and a weight function $W : E \to \mathbb{R}$.

Question: Find the maximum weight base of the matroid.

We will solve this using greedy algorithm.

Algorithm 15: Max-Weight-Base(E, W)

Input: A matroid M = (E, I) is given as an input as an oracle and a weight function $W : E \to \mathbb{R}$.

Output: Find the maximum weight base of the matroid

```
1 begin
```

Theorem 5.3.7

The above algorithm outputs a maximum weight base

Proof: Let M be a matroid. We will prove that this greedy algorithm works by inducting on i. At any iteration i we need to prove the following claim:

Claim 5.3.1

At any iteration *i* there is a max weight base B_i such that $S_i \subseteq B_i$ and $B_i \setminus S_i \subseteq \{i+1,\ldots,n\}$.

Proof: Base case: $S = \emptyset$. So for base case the statement is true trivially. Assume that the statement is true up to (i-1) iterations.

Now $S_{i-1} \subseteq B_{i-1}$ where B_{i-1} is a maximum weight base and $B_{i-1} - S_{i-1} \subseteq \{i, ..., n\}$. Now three cases arise:

- **Case 1:** If $i \in B_{i-1}$ then $S_{i-1} + i \subseteq B_{i-1}$. Therefore $S_{i-1} + i$ is independent. So now $B_i = B_{i-1}$ and $S_i = S_{i-1} + i$ and $B_i S_i \subseteq \{i+1, \ldots, n\}$.
- **Case 2:** If $i \notin B_{i-1}$ and $S_{i-1} + i \notin I$. Then $S_i = S_{i-1}$ and $B_i = B_{i-1}$. And $B_i S_i \subseteq \{i+1, ..., n\}$.
- **Case 3:** If $i \notin B_{i-1}$ but $S_{i-1} + i \in I$. Then $S_i = S_{i-1} + i$. Now S_i can be extended to a B' by adding all but one element of B_{i-1} . So $|B'| = |B_{i-1}|$. Let the element which is not added is $j \in B_{i-1}$. So $B' = B_{i-1} + i j$.

$$wt(B') = Wt(B_{i-1}) - wt() + wt(i)$$

But we have $wt(i) \ge wt(j)$. So $wt(B') \ge wt(B_{i-1})$. Now since B_{i-1} has maximum weight we have $wt(B') = wt(B_{i-1})$. Then our $B_i = B'$. So $B_i - S_i \subseteq \{i+1, \ldots, n\}$.

Hence the claim is true for the *i*th stage as well. Therefore the claim is true.

Claim 5.3.2

At any iteration, $T_i = \{t_1, \dots, t_k\}$, then T_i is a maximum weight independent set with at most *i* elements

Proof: We will prove by induction. Base Case: i = 0. Then $T_i = \emptyset$. So the statement follows naturally. Assume T_{i-1} is maximum weight independent set with at most i - 1 elements. Now for a contradiction, say $\hat{T}_i \in I$ of size at most i with strictly larger weight than T_i . Then $\exists x \in \hat{T}_i - T_{i-1}$ such that $T_{i-1} \cup \{x\} \in I$. Then we have

$$wt(\hat{T}_i - x) \le wt(T_{i-1})$$

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by inductive hypothesis. The only element that extend T_{i-1} are those t_{i-1} . Therefore $wt(x) \le wt(t_i)$. Hence we have

$$wt(\hat{T}_i - x) + wt(x) \le wt(T_{i-1}wt(t_i) \implies wt(\hat{T}_I) \le wt(T_i)$$

But we assumes $wt(\hat{T}_i) > wt(T_i)$. Hence contradiction.

Therefore using the claims, after the algorithm finished we have no elements left to check, so the current set has the maximum weight which is also an independent set. So the algorithm successfully returns a maximum weight base.

5.3.3 Job Selection with Penalties

FIND FEASIBLE SCHEDULE

Input: Set *J* of *n* jobs with deadlines d_1, \ldots, d_n and rewards w_1, \ldots, w_n

Question: Each jobs unit time and we have a single machine to process their jobs. Give a feasible schedule of jobs

with maximum reward

First lets define what is a schedule and what is a feasible schedule:

Definition 5.3.2: Feasible Schedule

For a subset *S* of jobs:

- \bigcirc A schedule is an ordering of *S*
- ② A feasible schedule is one where one job in *S* gets finished by deadline.
- (3) A set $S \subseteq J$ is feasible if S has a feasible schedule.

Now for any $S \subseteq J$, and $t \in \mathbb{Z}_+$, define $N_t(S) = \{j \in S : d_i \le t\}$. Then we have the following lemma:

Lemma 5.3.8

The following are equivalent:

- \bigcirc *S* is feasible
- ② $\forall t \in \mathbb{Z}_t, |N_t(S)| \leq t$
- (3) The schedule that orders jobs by deadline is feasible

Proof:

 $3 \implies 1$: This follows naturally

 $1 \Longrightarrow 2$: Suppose not. Then $\exists t$ such that $|N_t(S)| > t$. Then by time t, greater than t many jobs have to be completed. But S is feasible so every job is finished by deadlines and each job takes unit take. Hence by time t, more than t jobs can not finished. Hence contradiction.

 $2\Longrightarrow 3$: The schedule orders the jobs by deadline. We induction on t. For t=1 we have $|N_1(S)|\le 1$. Hence by t=1 at most one job is completed. At t=1 the jobs are completed within deadline. Suppose till time t-1 the jobs are completed within deadlines. At time t we have $|N_t(S)|\le t$. Therefore all the jobs with deadlines $\le t$ in S. So they all can be completed within time t in any order. Therefore if we complete the jobs with deadline $\le t$ first then also we can complete all the jobs with deadline t within time t. Hence at time t all the jobs are completed within their deadlines. Hence by mathematical induction at time t=n all the jobs are completed within deadline. Therefore the schedule orders jobs by deadline then it is a feasible schedule.

Lemma 5.3.9

Consider M = (J, I) where S is feasible $\implies S \in I$. Then M is a matroid. (Assume that no two jobs have same deadline)

Proof: Suppose D := the maximum of all deadlines. Consider the set

$$\mathcal{L} = \{N_t(J) : t \in [D]\}$$

Then take $I' = \{S \subseteq J : |N_t(S)| \le t \ \forall t \in [D]\}$. By Lemma 5.3.4 M = (J, I') is a laminar matroid. And by Lemma 5.3.8 I' is the set of feasible schedules. Therefore I' = I. Hence M is a matroid.

Alternate Proof:

- ① Downward Closure: If $S \in I$ then S is feasible. Then for any subset T of S all the jobs are completed within deadlines since S is feasible. So $T \in I$.
- ② Exchanges Property: Given $S, T \in I$ and |T| < |S|. Now order S and T by deadlines. Let j be the job with largest deadline that is not in S i.e. $j = \max_{i \in S \setminus T} d_i$. Then we claim that $T \cup \{j\} \in I$.

Now define

$$T^{<} = \{i \in T : d_i < d_j\}$$
 $T^{>} = \{i \in T : d_i > d_j\}$

And also similarly define

$$S^{<} = \{i \in S : d_i < d_i\}$$
 $S^{>} = \{i \in S : d_i > d_i\}$

As we defined j we have $T^> = S^>$. Since we have |S| > |T| we have $|S^<| \ge |T^<|$.

Now if $T \cup \{j\}$ is not feasible then $\exists t$ such that $|N_t(T \cup \{j\})| > t$. Since T is feasible we have $|N_t(T)| \le t$. Hence $t \ge d_j$ otherwise $N_t(T \cup \{j\}) = N_t(T)$. But then

$$|N_t(T \cup \{j\})| = |T^{<}| + 1 + |\{i \in T \cup \{j\}: d_i < d_i \le t\}| \le |S^{<}| + 1 + |\{i \in S \cup \{j\}: d_i < d_i \le t\}| = |N_t(S)| \le t$$

Therefore we obtain $|N_t(T \cup \{j\})| \le t$. Hence contradiction. Therefore $T \cup \{j\}$ is feasible.

Dijkstra Algorithm with Data Structures

MINIMUM WEIGHT PATH

Input: Directed Graph G = (V, E), $s \in V$ is source and $W = \{w_e \in \mathbb{Z}_+ : e \in E\}$

Question: $\forall v \in V - \{s\}$ find minimum weight path $s \rightsquigarrow v$.

This is the problem we will discuss in this chapter. In this chapter we will often use the term 'shortest distance' to denote the minimum weight path distance. One of the most famous algorithm for finding out minimum weight paths to all vertices from a given source vertex is Dijkstra's Algorithm

6.1 Dijkstra Algorithm

We will assume that the graph is given as adjacency list. Dijkstra Algorithm is basically dynamic programming. Suppose $\delta(v)$ is the shortest path distance from $s \rightsquigarrow v$. Then we have the following relation:

$$\delta(v) = \min_{u:(u,v)\in E} \{\delta(u) + e(u,v)\}$$

And suppose for any vertex $v \in V - \{s\}$, dist(v) be the distance from s estimated by the algorithm at any point. This is why Dijkstra's algorithm maintains a set S of vertices whose final shortest-path weights from the source s have already been determined. The algorithm repeatedly selects the vertex $u \in V - S$ with minimum shortest-path estimate and estimates the distances of neighbors of u. So here is the algorithm:

Algorithm 16: DIJKSTRA(G, s, W)

```
Input: Adjacency Matrix of digraph G = (V, E), source vertex s \in V and weight function W = \{w_e \in \mathbb{Z}_+ : e \in E\}

Output: \forall v \in V - \{s\} minimum weight path from s \rightsquigarrow v

1 begin

2 |S \longleftarrow \emptyset, U \longleftarrow V|

3 |dist(s) \longleftarrow 0, \forall v \in V - \{s\}, dist(v) \longleftarrow \infty

4 while U \neq \emptyset do

5 |u \longleftarrow \min_{u \in U} dist(u)| and remove u from U

6 |S \longleftarrow S \cup \{u\}|

7 |for \ e = (u, v) \in E \ do

|dist(v) \longleftarrow \min \{dist(v), dist(u) + w(u, v)\}|
```

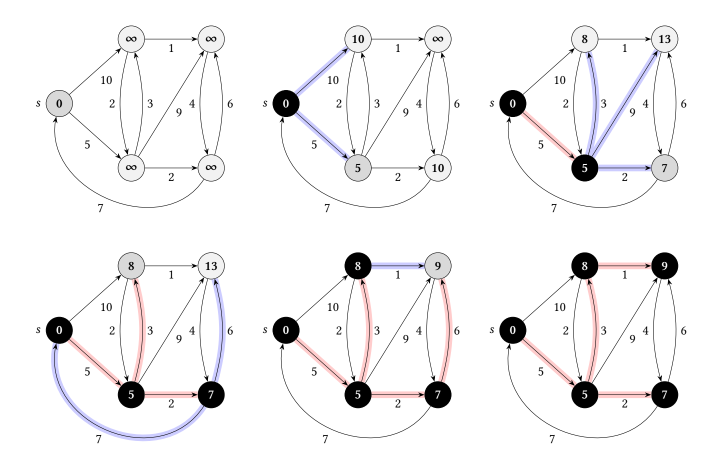


Figure 6.1: The execution of Dijkstra's algorithm. The source s is the leftmost vertex. The shortest-path estimates appear within the vertices, and shaded edges indicate predecessor values. Black vertices are in the set *S* and at any iteration of while loop the shaded vertex has the minimum value. At any iteration the red edges are the edges considered in minimum weight path from *s* using only vertices in *S*.

Suppose at any iteration t, let $dist_t(v)$ denotes the distance v from s calculated by algorithm for any $v \in V$ and $S^{(t)}$ denote the content of S at t^{th} iteration. In order to show that the algorithm correctly computes the distances we prove the following lemma:

Theorem 6.1.1

For each $v \in S^{(t)}$, $\delta(v) = dist_t(v)$ for any iteration t.

Proof: We will prove this induction. Base case is $|S^{(1)}| = 1$. S grows in size. Then only time $|S^{(1)}| = 1$ is when $S^{(1)} = \{s\}$ and $d(s) = 0 = \delta(s)$. Hence for base case this is correct.

Suppose this is also true for t-1. Let at t^{th} iteration the vertex $u \in V-S$ is picked. By induction hypothesis for all $v \in S^{(t)} - \{u\}$, $dist_t(v) = dist_{t-1}(v) = \delta(v)$. So we have to show that $dist_t(u) = \delta(u)$.

Suppose for contradiction the shortest path from $s \rightsquigarrow u$ is P and has total weight $= \delta(u) = w(P) < dist_t(u)$. Now P starts with vertices from $S^{(t)}$ by eventually leaves S. Let (x, y) be the first edge in P which leaves S i.e. $x \in S$ but $y \notin S$. By inductive hypothesis $dist_t(x) = \delta(x)$. Let P_y denote the path $s \rightsquigarrow y$ following P. Since y appears before u we have

$$w(P_u) = \delta(y) \le \delta(u) = w(P)$$

Now

$$dist_t(y) \leq dist_t(x) + w(x, y)$$

since y is adjacent to x. Therefore

$$dist_t(y) \le dist_t(x) + w(x, y) = \delta(y) \le dist_t(y) \implies dist_t(y) = \delta(y)$$

Now since both $u, y \notin S^{(t)}$ and the algorithm picked up u we have $\delta(u) < dist_t(u) \le dist_t(y) = \delta(y)$. But we can not have both $\delta(y) \le \delta(u)$ and $\delta(u) < \delta(y)$. Hence contradiction. Therefore $\delta(u) = dist_t(u)$. Hence by mathematical induction for any iteration t, for all $v \in S^{(t)}$, $\delta(v) = dist_t(v)$.

Therefore by the lemma after all iterations S has all the vertices with their shortest distances from s and henceforth the algorithm runs correctly.

- 6.2 Data Structure 1: Linear Array
- 6.3 Data Structure 2: Min Heap
- 6.4 Amortized Analysis

6.5 Data Structure 3: Fibonacci Heap

Instead of keeping just one Heap we will no keep an array of Heaps. We will also discard the idea of binary trees. We will now use a data structure which will take the benefit of the faster time of both the data structure. I.e. The * is because in Fibonacci Heap the amortized time taken by Extract-Min is $O(\log n)$.

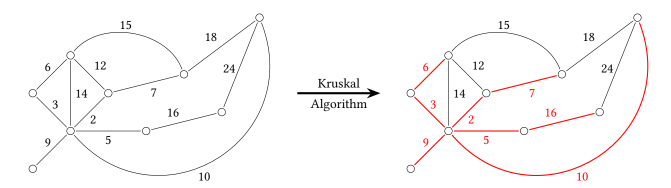
Since Fibonacci heap is an array of heaps there is a *rootlist* which is the list of all the roots of all the heaps in the Fibonacci heap. There is a *min-pointer* which points to the root with the minimum key. For each node in the Fibonacci heap we have a pointer to its parent and we keep 3 variables. The 3 variables are *degree*, *size* and *lost* where *lost* is a Boolean Variable. For any node x in the Fibonacci heap the x-degree is the number of children x has, x-size is the number of nodes in the tree rooted at x and x-lost is 1 if and only if x has lost a child before. Why any node will lost a child that explanation we will give later. With this set up let's dive into the data structure.

6.5.1 Inserting Node

To insert a node we call the Fib-Insert function and in the function the algorithm initiates the node with setting up all the pointers and variables then add the node to the *rootlist*.

Kruskal Algorithm with Data Structure

7.1 Kruskal Algorithm



- 7.2 Data Structure 1: Array
- 7.3 Data Structure 2: Left Child Right Siblings Tree
- 7.4 Data Structure 3: Union Find

7.4.1 Analyzing the Union-Find Data-Structure

We call a node in the union-find data-structure a leader if it is the root of the (reversed) tree.

Lemma 7.4.1

Once a node stop being a leader (i.e. the node in top of a tree). it can never become a leader again.

Proof: A node x stops being a leader only because of the Union operation which made x child of a node y which is a leader of a tree. From this point on, the only operation that might change the parent pointer of x is the Find operation which traverses through x. Since path-compression only change the parent pointer of x to point to some other node y. Therefore the parent pointer of x will never become equal to itself i.e. x can never be a leader again. Hence once x stops being a leader it can never be a leader again.

Lemma 7.4.2

Once a node stop being a leader then its rank is fixed.

Proof: The rank of a node changes only by an Union operation. But the Union operation only changes the rank of nodes that are leader after the operation is done. Therefore once a node stops being a leader it's rank will not being changed by an Union operation. Hence once a node stop being a leader then its rank is fixed.

Lemma 7.4.3

Ranks are monotonically increasing in the reversed trees, as we travel from a node to the root of the tree.

Proof: To show that the ranks are monotonically increasing it suffices to prove that for all edge $u \to v$ in the data structure we have $\operatorname{rank}(u) < \operatorname{rank}(v)$.

Lemma 7.4.4

When a node gets rank k than there are at least $\geq 2^k$ elements in its subtree.

Corollary 7.4.5

For all vertices v, v. $rank \leq \lfloor \log n \rfloor$

Corollary 7.4.6

Height of any tree $\leq |\log_2 n|$

Lemma 7.4.7

The number of nodes that get assigned rank k throughout the execution of the Union-Find data-structure is at most $\frac{n}{2^k}$.

Define N(r) = #vertices with rank at least k. Then by the above lemma we have $N(r) \leq \frac{n}{2k}$.

Lemma 7.4.8

The time to perform a single find operation when we perform union by rank and path compression is $O(\log n)$ time.

We will show that we can do much better. In fact we will show that for m operations over n elements the overall running time is $O((n+m)\log^* n)$

Lemma 7.4.9

During a single FIND(x) operation, the number of jumps between blocks along the search path is $O(\log^* n)$.

Lemma 7.4.10

At most $|Block(i)| \le Tower(i)$ many FIND operations can pass through an element x which is in the i^{th} block (i.e. $INDEX_B(x) = i$) before x.parent is no longer in the i^{th} block. That is $INDEX_B(x.parent) > i$.

Lemma 7.4.11

There are at most $\frac{n}{Tower(i)}$ nodes that have ranks in the i^{th} block throughout the algorithm execution.

Lemma 7.4.12

The number of internal jumps performed, inside the i^{th} block, during the lifetime of Union-Find data structure is O(n).

Theorem 7.4.13

The number of internal jumps performed by the Union-Find data structure overall $O(n \log^* n)$.

Theorem 7.4.14

The overall time spent on m FIND operations, throughout the lifetime of a Union-Find data structure defined over n elements is $O((n+m)\log^* n)$.

Red Black Tree

Definition 8.1: Perfect Binary Tree

df

Definition 8.2: Red Black Tree

df

Lemma 8.1

Every perfect binary tree with k leaves has 2k - 1 nodes (i.e. k - 1 internal nodes).

Maximum Flow

9.1 **Flow**

Suppose we are given a directed graph G = (V, E) with a source vertex s and a target vertex t. And additionally for every edge $e \in E$ we are given a number $c_e \in \mathbb{Z}_0$ which is called the capacity of the edge.

Definition 9.1.1: Flow

1
$$\forall e \in E, f(e) \leq c_e$$

An
$$s-t$$
 flow is a function $f:E\to\mathbb{R}_0$ which satisfies the following:

① $\forall \ e\in E, f(e)\leq c_e$
② $\forall \ v\in V\setminus\{s,t\}, \sum\limits_{e\in in(v)}f(e)=\sum\limits_{e\in out(v)}f(e)$

Also the value of a flow f is denoted by $|f| := \sum_{e \in out(s)} f(e)$.

Before proceeding into the setup and the problem first we will assume some things

• $in(s) = \emptyset$ i.e. there is no edge into s. Assumption.

- $out(t) = \emptyset$ i.e. there is no edge out of t.
- There are no parallel edges

Lemma 9.1.1

For any flow f, $|f| = \sum_{e \in in(t)} f(e)$

Proof: We have for every edge $e \in E$, $\exists v \in V$ such that $e \in in(v)$ and $\exists u \in V$ such that $e \in out(u)$. Hence we get

$$\sum_{e \in E} f(e) = \sum_{v \in V} \sum_{e \in in(v)} f(e) = \sum_{v \in V} \sum_{e \in out(v)} f(e) \implies \sum_{v \in V} \left[\sum_{e \in in(v)} f(e) - \sum_{e \in out(v)} f(e) \right] = 0$$

Now we know $\forall v \in V \setminus \{s, t\}$. $\sum_{e \in in(v)} f(e) = \sum_{e \in out(v)} f(e)$. Therefore we get

$$\sum_{v \in V} \left[\sum_{e \in in(v)} f(e) - \sum_{e \in out(v)} f(e) \right] = 0 \implies \sum_{v \in \{s,t\}} \left[\sum_{e \in in(v)} f(e) - \sum_{e \in out(v)} f(e) \right] = 0 \implies \sum_{e \in out(s)} f(e) - \sum_{e \in in(t)} f(e)$$

Hence we have $|f| = \sum_{e \in in(t)} f(e)$.

Max Flow

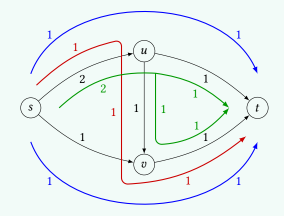
Input: A directed graph G = (V, E) with source vertex s and target vertex t and for all edge $e \in E$ capacity

of the edge $c_e \in \mathbb{Z}_+$

Question: Given such a graph and its capacities find an s - t flow which has the maximum value

Example 9.1.1

Consider the following directed graph with capacities: $V = \{s, t, u, v\}$, $c_{s,u} = 2$, $c_{s,v} = c_{u,t} = c_{v,t} = c_{u,v} = 1$. Firstly the following function: f': f'(s, u) = 2 = f(u, t). It is not a flow since $f(u, t) = 2 > 1 = c_{u,t}$. Now we define three different flow functions:



- f: f(s, u) = f(u, v) = f(v, t) = 1 and otherwise 0. Therefore |f| = 1
- g: g(s, u) = g(u, t) = 1, g(s, v) = g(v, t) = 1 and otherwise 0. Therefore |g| = 2
- h: h(s, u) = 2, h(u, t) = h(u, v) = h(v, t) = 1 and otherwise 0. Therefore |h| = 2

Notice here q and h has the maximum flow value.

9.2 Ford-Fulkerson Algorithm

Definition 9.2.1: Residual Graph

Given a directed graph G = (V, E) and capacities C_e for all $e \in E$ and an s - t flow f the residual graph $G_f = (V, E_f)$ has the edges with the following properties:

- ① If $(u,v) \in E$ and f(u,v) > 0 then $(v,u) \in E_f$ and $c_{v,u}^f = f(u,v)$. Such an edge is called a *backward* edge.
- ② If $(u, v) \in E$ and $f(u, v) < c_{u,v}$ then $(u, v) \in E_f$ and $c_{u,v}^f = c_{u,v} f(u,v)$. It is called *forward* edge.

Algorithm 17: Ford-Fulkerson

```
Input: Directed graph G=(V,E), source s, target t and edge capacities C_e for all e\in E
```

Output: Flow f with maximum value

```
1 begin
2 | for e \in E do
3 | \int f(e) = 0
4 | while \exists s \leadsto t \ path \ P \ in \ G_f do
5 | \delta \longleftarrow \min_{e \in P} \{c_e^f\}  for e = (u, v) \in P do
6 | if e is Forward Edge then
7 | \int f(u, v) \longleftarrow f(u, v) + \delta
8 | else
9 | \int f(u, v) \longleftarrow f(v, u) - \delta
```

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Lemma 9.2.1

At any iteration the f' obtained after the flow augmentation of the flow f is a valid flow

Proof: At any iteration let P be the path from $s \rightsquigarrow t$ and $\delta = \min_{e \in P} c_f(e)$. Let f' be the new function such that for each $(u,v) \in P$ if (u,v) is forward edge in G_f then $f'(u,v) = f(u,v) + \delta$ and if (u,v) is backward edge in G_f then $f'(v,u) = f(v,u) - \delta$ and for other edges $e \in E \setminus P$, f'(e) = f(e).

Now since $\delta = \min_{e \in P} c_f(e)$, $c_f(e) \ge \delta$ for all $e \in P$. Hence if (u, v) is backward edge then $(v, u) \in E$ and $c_f(u, v) = f(u, v)$. Hence $f'(v, u) = f(v, u) - \delta \ge 0$. Therefore for all $e \in E$, $f'(e) \ge 0$.

Now first we will show $f'(e) \le c_e$ for all $e \in E$. If $(u,v) \in P$ is a forward edge then $(u,v) \in E$ and $c_f(u,v) = c_{u,v}f(u,v)$. Therefore $f'(u,v) = f(u,v) + \delta \le f(u,v) + c_{u,v} - f(u,v) = c_{u,v}$. Now if $(u,v) \in P$ is a backward edge then $(v,u) \in E$ and $c_f(u,v) = f(u,v)$. Therefore $f'(u,v) = f(u,v) - \delta \le f(u,v) \le c_{u,v}$. For other edges $e \in E \setminus P$, $f'(e) = f(u) \le c_e$. Therefore $f'(e) \le c_e$ for all $e \in E$

Now we will prove for all $v \in V \setminus \{s, t\}$, $\sum_{e \in in(v)} f'(e) = \sum_{e \in out(v)} f'(e)$. If v is not in the path P in G_f then, f'(e) = f(e)

for all edges $e \in in(v) \cup out(v)$. Hence the condition is satisfied for such vertices. Suppose v is in the path P. Then there are two edges e_1 and e_2 in P which are incident on e. If both are forward edges or both are backward edges then one of them is in in(v) and other one is in out(v). WLOG suppose $e_1 \in in(v)$ and $e_2 \in out(v)$ we have

$$\sum_{e \in in(v)} f'(e) = \sum_{e \in in(v) \setminus \{e_1\}} f(e) + f(e_1) \pm \delta = \sum_{e \in out(v) \setminus \{e_2\}} f(e) + f(e_2) \pm \delta = \sum_{e \in out(v)} f'(e)$$

If one of e_1 , e_2 forward edge and other one is backward edge then either e_1 , $e_2 \in in(v)$ (when e_1 is forward and e_2 is backward) or e_1 , $e_2 \in out(v)$ (when e_1 is backward and e_2 is forward). Now if e_1 , $e_2 \in in(v)$, $f'(e_1) + f'(e_2) = f(e_1) + \delta + f(e_2) - \delta = f(e_1) + f(e_2)$ and if e_1 , $e_2 \in out(v)$ then $f'(e_1) + f'(e_2) = f(e_1) - \delta + f'(e_2) + \delta = f(e_1) + f(e_2)$. Hence

$$\sum_{e \in in(v)} f'(e) = \sum_{e \in in(v)} f(e) = \sum_{e \in out(v)} f(e) = \sum_{e \in out(v)} f'(e)$$

Hence f' is a valid flow.

Lemma 9.2.2

At any iteration Given G_f if the flow, f' obtained after flow augmentation of f by δ then

$$|f'| = |f| + \delta$$

Proof: Since we augment flow along an $s \rightsquigarrow t$ path, the first edge of the path is always in out(s). Let the first edge is e = (s, u). Now e has to be a forward edge because otherwise $(u, s) \in E$ and then there is an incoming edge in G which is not possible. Hence

$$|f'| = \sum_{e \in out(s)} f'(e) = \sum_{e \in out(s) \setminus \{e\}} f(e) + f'(e) = \sum_{e \in out(s) \setminus \{e\}} f(e) + f(e) + \delta = \sum_{e \in out(s)} f(e) + \delta = |f| + \delta$$

Hence we have the lemma.

Lemma 9.2.3

At every iteration of the Ford-Fulkerson Algorithm the flow values and the residual capacities of the residual graph are non-negative integers.

Proof: Initial flow and the residual capacities are non-negative integers. Let till i^{th} iteration the flow values and the residual capacities were non-negative integers. Let the flow after i^{th} iteration was f. Hence $\forall e \in E, f(e) \in \mathbb{Z}_0$. Therefore in the G_f for all $e \in E_f$, $c_f(e) \in \mathbb{Z}_0$. Hence $\delta \in \mathbb{Z}_0$. Therefore $\forall e \in E, f'(e) \in \mathbb{Z}_0$. And therefore for all $e \in E_{f'}$ where $G_{f'}$ is the residual graph of the flow f', $c_{f'}(e) \in \mathbb{Z}_0$. Hence by mathematical induction the lemma follows.

At any iteration let P be the path from $s \rightsquigarrow t$. Then for all $e \in P$, $c_f(e) > 0$. Therefore $\delta = \min_{e \in P} c_f(e) \ge 1$. Therefore the algorithm must stop in at most $\sum_{e \in out(s)} c_e$ since we can have the value of a flow to be at max the value of the sum of capacities of edges in out(s) and therefore we can increase the flow at max that many times.

Lemma 9.2.4

If f is a max flow then there is no $s \rightsquigarrow t$ path in G_f .

Proof: Suppose there is an $s \rightsquigarrow t$ path P in G_f . We will show that then f is not a max flow following the algorithm. Then $\forall e \in P$, $c_f(e) > 0$. Hence $\delta = \min_{e \in P} c_f(e) \ge 1$. Now after the flow augmentation process of f by δ we get a new valid flow f' by Lemma 9.2.1 and by Lemma 9.2.2 we have $|f'| = |f| + \delta > ||f|$. Hence f is not a maximum flow. Hence contradiction. Therefore there is no $s \rightsquigarrow t$ path in G_f .

9.2.1 Max Flow Min Cut

Definition 9.2.2: Cut Set

For a graph G = (V, E) and a subset $A \subseteq V$, the cut $(A, V \setminus A)$ is a bipartition of V where the edges E_A of the graph $G_A = (A, V \setminus A, E_A)$ is the set $E_A = E \cap (A \times (V \setminus A))$.

Now if s, t are two vertices of G then an s-t Cut $(A, V \setminus A)$ is a cut such that $s \in A$ and $t \in V \setminus A$.

Now we define for a cut $(A, V \setminus A)$ the Capacity of the Cut $(A, V \setminus A) = \sum_{e \in E_A} c_e$. For an s - t cut $(A, V \setminus A)$ we denote the capacity of the cut by cap(A) A Min s - t Cut is a s - t cut of minimum capacity. Then we have the following relation between cut and flow.

Lemma 9.2.5

Given a graph G = (V, E), $s, t, c_e \in \mathbb{Z}_0$ for all $e \in E$ for any flow f and a s - t cut $(A, V \setminus A)$

$$|f| \le cap(A)$$

Proof: Given f and the s - t cut $(A, V \setminus A)$ we have

$$|f| = \sum_{e \in out(s)} f(e)$$

$$= \sum_{v \in A} \left[\sum_{e \in out(v)} f(e) - \sum_{e \in in(v)} f(e) \right]$$

$$= \sum_{e = (u,v), \ u \notin A, v \notin A} f(e) - \sum_{e = (u,v), \ u \notin A, v \in A} f(e)$$

$$= \sum_{e \in out(A)} f(e) - \sum_{e \in in(A)} f(e)$$

$$\leq \sum_{e \in out(A)} f(e) \leq \sum_{e \in out(A)} c_e = cap(A)$$
[Edges for both endpoints in A are canceled out]

Hence we have the lemma.

Having this lemma we have for any flow f and s - t cut $(A, V \setminus A)$ we have

$$|f| \leq cap(A) \implies \max_{f} |f| \leq \min_{s-t \ cut \ (A,V \setminus A)} cap(A)$$

So we have the following theorem that the value of maximum flow is equal to the capacity of minimum cut.

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Theorem 9.2.6 Max Flow Min Cut

Given a graph G = (V, E), $s, t, c_e \in \mathbb{Z}_0$ for all $e \in E$. Then the following are equivalent:

- (1) f is a maximum flow.
- (2) There is no $s \rightsquigarrow t$ path in G_f
- (3) There exists an s t cut of capacity |f|

Proof:

- $(1) \Longrightarrow (2)$: This is by Lemma 9.2.4.
- (2) \Longrightarrow (3): We are given a flow f such that there is no $s \leadsto t$ path in G_f . We will construct a s-t cut which has the capacity |f|. Now take A to be all the vertices reachable from s in G_f . This is a valid s-t cut since $s \in A$ and as there is no $s \leadsto t$ path in G_f , $t \notin A$. Now

$$|f| = \sum_{e \in out(A)} f(e) - \sum_{e \in in(A)} f(e)$$

Now $\forall e = (u, v) \in E$ where $u \in A$ and $v \notin A$ we have $c_{u,v} = f(u,v) \implies c_{u,v} - f(u,v) = 0$ since otherwise $c_{u,v} - f(u,v) \neq 0 \implies c_{u,v} > f(u,v) \implies (u,v) \in E_f$ and therefore v is reachable from $v \notin A$ contradiction. Therefore $v \in A$ and $v \in A$ we have $v \in A$ and $v \in A$ we have $v \in A$ and $v \in A$ we have $v \in A$ and $v \in A$ and $v \in A$ and $v \in A$ contradiction. Hence we have

$$|f| = \sum_{e \in out(A)} f(e) - \sum_{e \in in(A)} f(e) = \sum_{e \in out(A)} c_e = cap(A)$$

(3) \Longrightarrow (1): Now by Lemma 9.2.5 we have for any flow f and s - t cut

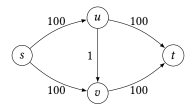
$$|f| \le cap(A) \implies \max_{f} |f| \le \min_{s-t \ cut \ (A, V \setminus A)} cap(A)$$

Now given f there exists an s - t cut of capacity |f|. Hence f is a max flow.

Hence at the end of the Ford-Fulkerson Algorithm let the flow returned by the algorithm is f. The algorithm terminates when there is no $s \rightsquigarrow t$ path in G_f . Hence by Max Flow Min Cut Theorem we have f is a maximum flow. This completes the analysis of the Ford-Fulkerson Algorithm.

Since the capacities of the edges can be very large we want an algorithm return the maximum flow with running time $poly(n, m, \log c_e)$ where n is the number of vertices and m is number of edges and $\log c_e$ basically means number of bits at most needed to represent the capacities.

But Ford-Fulkerson algorithm takes does not run in $poly(n, m, \log c_e)$ instead $poly(n, m, c_e)$ as the while loop in the algorithm takes $poly(c_e)$ many iterations. For example in the following graph: it takes around 100 steps



and in general Ford-Fulkerson takes $O(|f_{\text{max}}|)$ time. For this reason we will now discuss a modification of the Ford-Fulkerson Algorithm which takes $poly(n, m, \log c_e)$ time, Edmonds-Karp Algorithm.

9.2.2 Edmonds-Karp Algorithm

To get a $poly(n, m, \log c_e)$ time algorithm we will always pick the shortest $s \rightsquigarrow t$ path in the residual graph. This algorithm is known as the Edmonds-Karp Algorithm

Suppose f_i be the total flow after i^{th} iteration. And G_{f_i} be the residual graph with respect f_i . Then $f_0(e) = 0$ for all $e \in E$ and $G_{f_0} = G$. Also suppose $dist_i(v) = \text{Shortest } s \rightsquigarrow v$ path distance in the residual graph G_{f_i} . Hence $dist_i(s) = 0$ for all i and $dist_i(t) = \infty$ at the end of the algorithm.

Note:-

In i^{th} iteration of the Ford-Fulkerson Algorithm or Edmonds-Karp Algorithm if P is the $s \leadsto t$ in the residual graph G_{f_i} where $e = (u, v) \in P$ and $c_{f_i}(u, v) = \delta = \min_{e \in P} c_{f_i}(e)$ then the edge (u, v) is not present in the next residual graph $G_{f_{i+1}}$. Thus at least one edge disappears in each iteration of Ford-Fulkerson or Edmonds-Karp Algorithm.

Now we will prove following two lemmas which will help us to prove that the Edmond-Karp algorithm takes O(mn) iterations.

Lemma 9.2.7

At any iteration $i, \forall v \in V$, $dist_i(v) \leq dist_{i+1}(v)$

Proof: Suppose this is not true. Then let i be the first iteration in which there exists a vertex $v \in V$ such that $dist_i(v) > dist_{i+1}(v)$. We pick such v which minimizes $dist_{i+1}(v)$. Consider the shortest path P from $s \leadsto v$ in $G_{f_{i+1}}$. Hence length of P, $|P| = dist_{i+1}(v)$. Let (u, v) be the last edge of P.

$$P: (s) \longrightarrow (u) \longrightarrow (v)$$

Then

$$dist_{i+1}(v) = dist_{i+1}(u) + 1 \ge dist_i(u) + 1$$

Here the last inequality follows because v is the vertex which has the minimum $dist_{i+1}(v)$ among all the vertices $w \in V$ which follows $dist_i(w) > dist_{i+1}(w)$. Now we will analyze case wise.

• Case 1: $(u, v) \in E_{f_i}$. Then

$$dist_i(v) \leq dist_i(u) + 1 \leq dist_{i+1}(v)$$

But this is not possible since $dist_i(v) > dist_{i+1}(v)$.

• Case 2: $(u,v) \notin E_{f_i}$. Then $(v,u) \in E_{f_i}$. Since $(u,v) \in E_{f_{i+1}}$ then we must have sent flow along (v,u). Since we take the shortest $s \leadsto t$ path in G_{f_i} in the algorithm we have $dist_i(u) = dist_i(v) + 1$. But then

$$dist_i(u) \le dist_{i+1}(v) - 1 \implies dist_{i+1}(v) \ge dist_i(v) + 2$$

But this is not possible.

Hence contradiction f Therefore for all iterations i, for all vertices $v \in V$, $dist_i(v) \leq dist_{i+1}(v)$.

Lemma 9.2.8

For any edge $e = (u, v) \in E$ the number of iterations where either (u, v) appears or (v, u) appears is at most O(n) i.e.

$$\left|\left\{i\colon (u,v)\notin G_{f_{i}},(u,v)\in G_{f_{i+1}}\right\}\right|+\left|\left\{i\colon (v,u)\notin G_{f_{i}},(v,u)\in G_{f_{i+1}}\right\}\right|=O(n)$$

Proof: Following the proof of Lemma 9.2.7 in the second case we showed if $(u,v) \notin G_{f_i}$ but $(u,v) \in G_{f_{i+1}}$ then $dist_{i+1}(v) \ge dist_i(v) + 2$. Hence the distance increases by at least 2. Now this can happen at most O(n) many times since $\forall i, dist_i(v) \le n-1$. Hence the number of iterations where either (u,v) appears or (v,u) appears is at most O(n).

With this this lemma we will prove that the Edmonds-Karp Algorithm takes O(mn) iterations.

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Theorem 9.2.9

Edmonds-Karp Algorithm terminates in O(mn) many iterations.

Proof: For k iterations at least k edges must disappear. Since each edge can reappear O(n) times by Lemma 9.2.8, it can disappear at most O(n) many times. In each iteration at least one edge disappears. Now after O(mn) iterations number of disappearances is at most O(mn). But after O(mn) many disappearances there are no edge remaining and therefore there is no $s \rightsquigarrow t$ path. Hence the algorithm terminates. Therefore the Algorithm terminates in O(mn) iterations.

Hence Edmond-Karp Algorithm takes $O(m^2n)poly(\log c_e) = O\left(m^2n\log^{O(1)}(c_e)\right)$ time since it takes O(mn) iteratives tions and in each iteration it finds the shortest $s \rightsquigarrow t$ path in G_f in O(m) time and in each iteration it does addition and subtraction and finds minimum of the capacities which takes polynomial of the bits needed to represent them time.

9.3 Preflow-Push/Push-Relabel Algorithm

In this algorithm we will maintain something called "Preflow" which is not a valid flow. Unlike Ford-Fulkerson, Edmonds-Karp it does not maintain a $s \rightsquigarrow t$ path in the residual graph and the algorithm stops when the preflow is actually a valid flow.

Definition 9.3.1: Preflow

Given a graph G = (V, E) and the edge capacities c_e , a function $f : E \to \mathbb{R}_0$ is a preflow if is satisfies:

1
$$\forall e \in E, f(e) \leq c_e$$

(2)
$$\forall v \in V \setminus \{s\}, \sum_{ein(v)} f(e) \ge \sum_{e \in out(v)} f(e)$$

Notice here unlike the definition of Flow here in the second criteria we need $\sum_{ein(v)} f(e) \ge \sum_{e \in out(v)} f(e)$ instead of

$$\sum_{e \in n(v)} f(e) = \sum_{e \in out(v)} f(e).$$

 $\sum_{ein(v)} f(e) = \sum_{e \in out(v)} f(e).$ Now define for all $v \in V$ and for all preflow f, $excess_f(v) = \sum_{e \in in(v)} f(e) - \sum_{e \in out(v)} f(e)$. If f is a preflow then $excess_f(s) \le 0$ and $\forall v \in V \setminus \{s\}, excess_f(v) \ge 0$

Lemma 9.3.1

For all preflow *f*

$$\sum_{v \in V} excess_f(v) = 0$$

Proof:

$$\sum_{v \in V} excess_f(v) = \sum_{v \in V} \left[\sum_{e \in in(v)} f(e) - \sum_{e \in out(v)} f(e) \right]$$
$$= \sum_{v \in V} \sum_{e \in in(v)} f(e) - \sum_{v \in V} \sum_{e \in out(v)} f(e)$$
$$= \sum_{e \in E} f(e) - \sum_{e \in E} f(e) = 0$$

Now for each $v \in V$ we assign a label $l(v) \in \mathbb{Z}_0$. The algorithm then sends flow from $u \to v$ if l(v) = l(u) - 1.

$$(u)$$

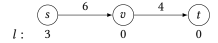
$$(v) = l(u)$$

Algorithm 18: Preflow-Push

```
Input: Directed graph G = (V, E), source s, target t and edge capacities C_e for all e \in E
   Output: Flow f with maximum value
1 begin
        Initially \forall e = (s, u) \in E, f(e) = c_e and f(e) for all other edges.
2
        l(s) \leftarrow n
3
        for v \in V \setminus \{s\} do
4
            l(v) \longleftarrow 0
5
        while \exists v \neq t, excess_f(v) > 0 do
6
            if \exists u, such that (v, u) \in E_f and l(u) = l(v) - 1 then
7
                 \delta \leftarrow \min \left\{ excess_f(v), c_f(v, u) \right\}
 8
                 if (v, u) is Forward Edge then
                   f(v,u) \longleftarrow f(v,u) + \delta
10
11
                      f(u,v) \longleftarrow f(u,v) - \delta
12
13
                 l(v) \longleftarrow l(v) + 1
                                                                                                                            //Relabeling
14
```

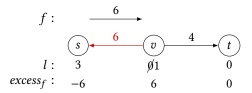
In the algorithm in line 8 if $\delta = c_f(v, u)$ then we call it *saturating push* and if $\delta = excess_f(v)$ then we call it *non-saturating push*.

Now we will show an example of how the algorithm on a graph. We will start the algorithm with the following graph:



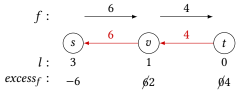
Below we will show change of the residual graph and preflow in each iteration of the While loop:

• Step 1:



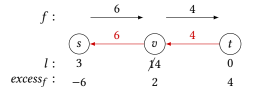
Since $excess_f(v) = 6 > 0$. So in first iteration v is taken. Since there is no edge (v, u) with l(u) = l(v) - 1, label of v got increased

• Step 2:



Since $excess_f f(v) = 2 > 0$, in second iteration again v is selected. There is an edge (v,t) with l(t) = 0 = l(v) - 1 = 1 - 1. Now $\delta = c_f(v,t) = 4$. Hence saturating push. The preflow gets updated, f(s,v) = 6, f(v,t) = 4.

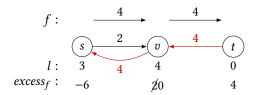
• Step 5:



Since $excess_f f(v) = 2 > 0$, in next 3 iterations again v is selected. Since there is no edge (v, u) with l(u) = l(v) - 1, label of v gets increased every time. Which becomes 4 after 3 iterations.

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• Step 6:



Since $excess_f f(v) = 2 > 0$, in this iteration again v is selected. There is an edge (v, s) with l(t) = 3 = l(v) - 1 = 4 - 1. Now $\delta =$ $excess_f(v, s) = 2$. Hence it's non-saturating push. So the preflow gets updated f(s, v) = 6 - 2 = 4, f(v, t) = 4. Now it's a valid flow. Now there is no vertex with postive excess. Hence the algorithm stops.

Observation 1. Labels are monotone non-decreasing.

Observation 2. For every iteration f is always a preflow. The proof is similar to Lemma 9.2.1 but use inequalities.

Observation 3. $\sum_{v} excess_f(v) = 0$ and $\forall v \in V \setminus \{s\}$, $excess_f(v) \ge 0$. Hence $excess_f(s) \le 0 \implies l(s)$ is unchanged.

Now suppose f^i denote the preflow after the i^{th} iteration of the algorithm. Then

$$f^{0}(e) = \begin{cases} c_{e} & \text{when } e = (s, u) \\ 0 & \text{otherwise} \end{cases}$$

Now we will show the correctness of the algorithm.

Lemma 9.3.2 $\forall v \in V, \forall i, excess_{f^i}(v) > 0 \implies \exists v \leadsto s \text{ in } G_{f^i}$

Proof: First we fix v and i such that $excess_{f^i} > 0$. Let X be the set of vertices reachable from v in G_{f^i} . Now

$$\sum_{u \in X} excess_{f^i}(u) = \sum_{u \in X} \left[\sum_{e \in in(v)} f^i(e) - \sum_{e \in out(v)} f^i(e) \right] = \sum_{e \in in(X)} f^i(e) - \sum_{e \in out(X)} f^i(e)$$

Now if $\sum_{e \in in(X)} f^i(e) > 0$ then $\exists e = (u', u) \in E$ such that $u' \notin X$ and $u \in X$ and $f^i(e) > 0$. Then the backward edge $(u, u') \in E_{f^i}$. Then u' is reachable from v in G_{f^i} . But $u' \notin X$. Contradiction f Therefore $\sum_{e \in V(Y)} f^i(e) = 0$. Hence

$$\sum_{u \in X} excess_{f^i}(u) = \sum_{e \in in(X)} f^i(e) - \sum_{e \in out(X)} f^i(e) \leq 0$$

But from Observation 3 we have $\forall w \in V \setminus \{s\}$, $excess_{f^i}(w) \geq 0$. But at the same time $\sum_{u \in X} excess_{f^i}(u) \leq 0$ and $excess_{fi}(v) > 0$. Hence \exists a vertex $u \in X$ such that $excess_{fi}(u) < 0$. But we know only vertex with negative excess is s. Therefore $s \in X$. Hence s is reachable from v.

Lemma 9.3.3

 $\forall i$, if $(u, v) \in G_{f^i}$ then $l(v) \ge l(u) - 1$.

Proof: We will prove this using induction on *i*. Initially l(s) = n and l(v) = 0 for all $v \in V \setminus \{s\}$. Hence for all edges (u, v)where $u, v \neq s$ this is satisfied. All the other edges incident on s are in in(s) in the residual graph. And $l(s) = n \geq l(u) = 0$. Therefore the base case is followed.

Now suppose the condition is true for f^{i-1} . Now in the i^{th} iteration suppose the selected vertex is $v \in V \setminus \{t\}$ with $excess_{f^{i-1}} > 0$. Now there are two possible cases.

• Case 1: If the step is relabeling then $f^{i-1} = f^i$, $G_{f^{i-1}} = G_{f^i}$ but v is relabeled by l(v) + 1. Now for any edge $e = (u,v) \in in(v)$ by Inductive Hypothesis $l(v) \ge l(u) - 1 \implies l(v) + 1 \ge l(u) - 1$. Now consider any edge $e = (v, w) \in out(v)$. By Inductive Hypothesis we have $l(w) \ge l(v) - 1$. Now if l(w) = l(v) - 1 then we would have pushed flow along the edge (v, w). Since that is not the case we have l(w) > l(v) - 1. Therefore $l(w) \ge (l(v) + 1) - 1$. Hence the condition is satisfied.

• Case 2: If the step is pushing flow then suppose we push flow along the edge $(v, w) \in E_{f^{i-1}}$ and l(w) = l(v) - 1. Now if we push flow along the edge (v, w) we might introduce the reverse edge (w, v) in G_{f^i} . In that case $l(v) = l(w) + 1 \ge l(w) - 1$. Hence the condition is satisfied.

Therefore by mathematical induction $\forall i, \forall (u, v) \in E_{f^i}, l(v) \ge l(u) - 1$.

Corollary 9.3.4

There is no $s \leadsto t$ path in G_{f^i} in any iteration i. Thus when the algorithm terminates f is a max flow.

Proof: Now l(s) = n and l(t) = 0. We fix v and i. If there is a $s \leadsto v$ path in G_{f^i} then length of the path is at most n-1. For each edge in the path the label decreases by at most 1 by Lemma 9.3.3. Hence $l(v) \ge 1$. Therefore for every vertex $v \in V$, reachable from s we have $l(v) \ge 1$. But l(t) = 0. Hence t is not reachable from s. Hence if the algorithm terminates, band if f is a valid flow then y Max Flow Min Cut Theorem it is a max flow.

Corollary 9.3.5

 $\forall v \in V, \forall i, l(v) \leq 2n.$

Proof: Suppose $\exists v, i$ such that l(v) = 2n and $excess_{f^i}(v) > 0$. By Lemma 9.3.2 there exists an $v \leadsto s$ path in G_{f^i} . Now by Lemma 9.3.3 for each edge in the path the label decreases by at most 1 and the length of the path is at most n-1. Since $l(v) = 2n, l(s) \ge n+1$. But we know l(s) for all i by Observation 3. Hence contradiction f Therefore for all $v \in V$ and f in f

Corollary 9.3.6

Total number relabeling operations is $\leq 2n^2$

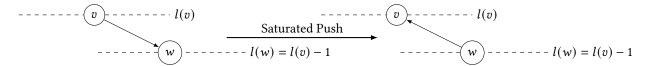
Proof: By Corollary 9.3.5 each vertex label can be at most 2n. So total number of relabeling operations done in the algorithm is at most $2n^2$

Now we need a bound on the number of push operations. We will count separately the number of Saturating Pushes and number of Non-Saturating Pushes.

Lemma 9.3.7

Total number of saturating pushes is $\leq 2mn$

Proof: We first fix an edge (v, w). Now we will count the number of saturating pushes along (v, w). Then $\delta = c_f(v, w)$. Now consider the scenario of two consecutive saturating pushes along (v, w). When the first saturating push along (v, w) occurred we have l(w) = l(v) - 1. Now if (v, w) is forward edge then $\delta = c_f(v, w) = c_{v,w} - f(v, w)$. Then new flow along (v, w) is $f(v, w) + c_f(v, w) = c_{v,w}$. Hence the edge (v, w) vanishes and the flow along (w, v) is $c_{v,w}$. If (v, w) is a backward edge then $\delta = c_f(w, v) = f(w, v)$. Hence then new flow along (w, v) is $f(w, v) - \delta = 0$. Hence again the (w, v) edge vanishes and the flow along (w, v) is f(w, v).



Therefore after a saturated push along (v, w) the edge vanishes and the (w, v) edge is there. Hence in order for another push along (v, w) the algorithm must push flow along (w, v). And this happens when we have the new labels of

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v, w follow the condition l'(w) = l'(v) + 1. Since by Observation 1 the labels never decreases in order for l(w) = l(v) + 1 the label of v must increase by at least 2.

Now starting from l(v) = 0 we have by Lemma 9.3.5 $l(v) \le 2n$ and for each saturating push along (v, w) the l(v) increase by 2. Hence at most n many saturating pushes occurred along (v, w). Now in the original graph since there are m edges the total number of saturating pushes is $\le 2mn$.

Now we will count the number of non-saturating pushes. For such pushes along any edge (v, u) the $excess_f(v)$ goes to 0. We define the potential function for a preflow f,

$$\phi(f) = \sum_{v: excess_f(v) > 0} l(v)$$

Now $\phi(f) \ge 0$ for all preflow f and initially at the start of the algorithm $\phi(f^0) = 0$.

Lemma 9.3.8

For each non-saturating push $\phi(f)$ decreases by at least 1.

Proof: Suppose at any iteration i a non-saturating push occur along an edge (v, w). Therefore l(w) = l(v) - 1. We will show that $\phi(f^i) \le \phi(f^{i-1}) - 1$. We have $\delta = excess_{f^{i-1}}(v)$. Now if (v, w) is a forward edge then new flow along (v, w) is $f^i(v, w) = f^{i-1}(v, w) + excess_{f^{i-1}}(v)$. Since $(v, w) \in out(v)$

$$excess_{f^{i}}(v) = \sum_{e \in in(v)} f^{i}(e) - \sum_{e \in out(v)} f^{i}(e) = \sum_{e \in in(v)} f^{i-1}(e) - \sum_{e \in out(v) \setminus \{(v,w)\}} f^{i-1}(e) - f^{i}(v,w) = excess_{f^{i-1}}(v) - \delta = 0$$

Otherwise if (v, w) is a backward edge. Then ew flow along (w, v) is $f^i(w, v) = f^{i-1}(w, v) - excess_{f^{i-1}}(v)$. Since $(w, v) \in in(v)$

$$excess_{f^{i}}(v) = \sum_{e \in in(v)} f^{i}(e) - \sum_{e \in out(v)} f^{i}(e) = f^{i}(w, v) + \sum_{e \in in(v) \setminus \{(w, v)\}} f^{i-1}(e) - \sum_{e \in out(v)} f^{i-1}(e) = -\delta + excess_{f^{i-1}}(v) = 0$$

In both cases $excess_{f^i}(v) = 0$. Therefore v goes out of the summation. Now there are two cases depending on the value of $excess_{f^{i-1}}(w)$

- Case 1: If $excess_{f^{i-1}}(w) > 0$ i.e. w had excess flow before push operation then $\phi(f^{i-1})$ decreases by l(v) i.e. $\phi(f^i) = \phi(f^{i-1}) l(v)$. Since l(w) = l(v) 1 and by Observation 1 $l(v) \ge 1$. Therefore $\phi(f^i) = \phi(f^{i-1}) l(v) \le \phi(f^{i-1}) 1$.
- Case 2: If $excess_{f^{i-1}}(w) = 0$, then $excess_{f^{i}}(w) = excess_{f^{i-1}}(w) + \delta > 0$ since $\delta = excess_{f^{i-1}}(v) > 0$ and therefore $\phi(f^{i}) = \phi(f^{i-1}) l(v) + l(w) = \phi(f^{i-1}) 1$

Hence for both the cases $\phi(f^i) \leq \phi(f^{i-1}) - 1$. Therefore $\phi(f^{i-1})$ decreases by at least 1.

Observation 4. For relabeling operation $\phi(f)$ increases by 1.

Since there are at most $2n^2$ relabeling operations by Corollary 9.3.6, $\phi(f)$ increases by at most $2n^2$ with relabeling operations.

Observation 5. For each saturating push excess f(v, w) might not go to 0 and therefore ϕ might increase.

Now by Lemma 9.3.7 total number of saturated pushes is at most 2mn. And by Corollary 9.3.5 each vertex has label at most 2n. Hence in total $\phi(f)$ can increase at most $2mn \times 2n = 4mn^2$ by saturated pushes. Hence $\phi(f)$ increases at most $2n^2 + 2mn \times 2m = O(mn^2)$.

Now

#Non-saturating Pushes \leq Total decrease in $\phi \leq$ Total increase in $\phi \leq 2n^2 + 4mn^2 = O(mn^2)$

Therefore total number of iterations of the While loop is #Relabeling+#Saturated Push+#Non-saturated Push= $2n^2 + 2mn + O(mn^2) = O(mn^2)$. There fore the algorithm takes $O(mn^2)$ iteration. In each iteration it takes O(m+n) time. Therefore the runtime of the algorithm is $O(mn^2)O(n+m) = O(m^2n^2)$.

CHAPTER 10

Randomized Algorithm

- 10.1 Estimated Binary Search Tree Height
- 10.2 Solving 2-SAT

CHAPTER 11 CHAPTER 11 CHAPTER 11

Matching Algorithms

12.1 Bipartite Matching

- 12.1.1 Using Max Flow
- 12.1.2 Using Augmenting Paths

12.1.3 Using Matrix Scaling

Here we will show a new algorithm for bipartite matching using matrix scaling. The paper which we will follow is [Linial-Samorodnitsky-Wigderson, STOC'1998]

Suppose $G = (L \cup R, E)$ a bipartite graph. If bipartite adjacency matrix of the graph G is A then the permanent of the matrix A,

$$per(A) = \sum_{\sigma \in S_n} \prod_{i=1}^n x_{i,\sigma(i)}$$

counts the number of perfect matchings in G. So we want to check if for a given bipartite graph $(L \cup R, E)$, per(A) > 0 or not where A is the bipartite adjacency matrix. Now there is a necessary and sufficient condition for existence of perfect matching in a bipartite graph which is called Hall's condition.

Theorem 12.1.1 Hall's Condition

A bipartite graph $G = (L \cup R, E)$ has an L-perfect matching if and only if $\forall S \subseteq L, |S| \le |N(S)|$ where $N(S) = \{v \in R: \exists u \in L, (u, v) \in E\}$

Proof: Now if *G* has a *L*-perfect matching then for every $S \subseteq L$, *S* is matched with some $T \subseteq R$ such that |S| = |T|. Therefore $T \subseteq N(S) \implies |S| = |T| \le |N(S)|$.

Now we will prove the opposite direction. Suppose for all $S \subseteq L$ we have $|S| \le |N(S)|$. Assume there is no L-perfect matching in G. Let M be a maximum L-matching in G. Let $u \in L$ is unmatched. Now consider the following sets:

$$X = \{x \in L : \exists M \text{-alternating path from } u \text{ to } x\},$$
 $Y = \{y \in R : \exists M \text{-alternating path from } u \text{ to } y\}$

Now notice that $N(X) \subseteq Y$. Since in a M-alternating path from u whenever the odd edges are not matching edges and the even edges are matching edges. So in the odd edges we can pick any neighbor except the one it is matched with and the immediate even edge before that connects that vertex with the vertex in R it is matched with. Hence we have $N(X) \subseteq Y$.

Now it suffices to prove that |X| > |Y|. Now let $y \in Y$. Suppose $u \leadsto x' \to y$ be the M-alternating path. If y is not matched then we could increase the matching by taking the odd edges of the path and thus obtain a matching with larger size than M. But M is maximum matching. Hence y is matched. Therefore we can extend the path by taking the matching edge incident on y and go the the vertex $x'' \in L$ i.e. the new M-alternating path becomes $u \leadsto x' \to y \to x''$ to have a M-alternating path $u \leadsto x''$. So |X| > |Y|.

Therefore we obtained a set of vertices $X \subseteq Y$ such that $|X| > |Y| \ge N(X)|$. This contradicts the assumption. Hence contradiction. Therefore G has a L-perfect matching.

We will use hall's condition on the adjacency matrix to check if per(A) is positive or not. Now multiplying a row or a column of a matrix by some constant c also multiplies the permanent of the matrix by c as well. In fact if $d_1, d_2 \in \mathbb{R}^n_+$ and $D_1 = diag(d_1)$ and $D_2 = diag(D_2)$ then $per(D_1AD_2) = \begin{pmatrix} n \\ \prod_{i=1}^n d_{1_i} \end{pmatrix} \begin{pmatrix} n \\ \prod_{i=1}^n d_{2_i} \end{pmatrix} per(A)$. So we can scale our original matrix A to obtain a different matrix B and from B we can approximate per(A) by approximating per(B). A natural strategy is to seek an efficient algorithm for scaling A to a doubly stochastic B.

Definition 12.1.1: Doubly Stochastic

A matrix $M \in \mathbb{R}^{m \times m}$ is doubly stochastic if entries are non-negative and each row and column sum to 1.

First we will show that Hall's Condition holds for doubly stochastic matrix. First let's see what it means for a matrix to satisfy hall's condition. A matrix with all entries non-negative holds Hall's Condition if for all $S \subseteq [n]$ if $T = \{i \in [n]: \exists j \in S, A(i, j) \neq 0\}$ then $|T| \geq |S|$. This also corresponds to the bipartite adjacency matrix satisfying the hall's condition since for any set of rows S the number of columns for which in the S rows at least one entry is non zero should be greater than or equal to |S|.

Lemma 12.1.2

Hall's Condition holds for doubly stochastic matrix.

Proof: Let M be the doubly stochastic matrix. Let $S \subseteq [n]$. So consider the $|S| \times n$ matrix which only consists of the rows in S. Call this matrix M_S^r . Now suppose T be the set of columns in M_S^r which has nonzero entries. Now consider the $n \times |T|$ matrix which only consists of the columns in T. Call this matrix M_T^c . Now since M is doubly stochastic we know sum of entries of M_S^r is |S| and sum of entries of M_T^c is |T|. Our goal is to show $|S| \le |T|$. Now since T is the only set of columns which have nonzero columns in M_S^r the elements which contributes to the sum of entries in M_S^r are in the T columns in M_S^r . Since these elements are also present in M_T^c we have $|T| \ge |S|$.

Hence for doubly stochastic matrices the permanent is positive. Now not all matrices are doubly stochastic. And in fact matrices with permanent zero will not be doubly stochastic so no amount of scaling will make it doubly stochastic. So we will settle for approximately doubly stochastic matrix. In order to make a matrix doubly stochastic first for each row we will divide the row with their row some. Now it becomes row stochastic. Then if its not approximately doubly stochastic for each column we will divide the column entries with their column sum. But first what ϵ -approximate doubly stochastic matrix means.

Definition 12.1.2: ϵ -Approximate Doubly Stochastic Matrix

A matrix is ϵ -approximate doubly stochastic if for each column, the column sum is in $(1 - \epsilon, 1 + \epsilon)$ and for each row, the row sum is in $(1 - \epsilon, 1 + \epsilon)$

Now we will show that even for ϵ -approximate doubly stochastic matrix the hall's condition holds.

Lemma 12.1.3

Halls's Condition holds for ϵ -approximate doubly stochastic matrix for $\epsilon < \frac{1}{10n}$

Proof: Let M is ϵ -approximate doubly stochastic matrix. Let $S \subseteq [n]$. So consider the $|S| \times n$ matrix which only consists of the rows in S. Call this matrix M_S^r . Now suppose T be the set of columns in M_S^r which has nonzero entries. Now consider the $n \times |T|$ matrix which only consists of the columns in T. Call this matrix M_T^c . Now the sum of entries in M_S^r is $\geq |S|(1-\epsilon)$ and sum of entries in M_T^c is $\leq |T|(1-\epsilon)$. Now since T is the only set of columns which have nonzero columns in M_S^r the elements which contributes to the sum of entries in M_S^r are in the T columns in M_S^r . Since these elements are also present in M_T^c we have $||T|(1+\epsilon) \geq |S|(1-\epsilon)$. Therefore we have

$$|T| \geq |S| \frac{1-\epsilon}{1+\epsilon} = |S| \left(1 - \frac{2\epsilon}{1+\epsilon}\right) \geq |S| \left(1 - 2\epsilon\right) > |S| \left(1 - \frac{1}{5n}\right) \geq |S| \left(1 - \frac{1}{|S|}\right) > |S| - 1$$

Since *T* is an integer we have $|T| \ge |S|$. Hence the Hall's condition holds.

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Therefore permanent of ϵ -approximate doubly stochastic matrix is also positive. Hence our algorithm for bipartite perfect matching is:

```
Algorithm 19: BP-MATRIX-SCALING
```

```
Input: Bipartite adjacency matrix A of G = (L \cup R, E)

Output: Decide if G has a perfect matching.

1 begin

2 | while True do

3 | A \leftarrow Scale every rows of A to make it row stochastic.

4 | if All column-sums are in (1 - \epsilon, 1 + \epsilon) then

5 | C return Yes

6 | A \leftarrow Scale every column of A to make it column stochastic.

7 | if All row-sums are in (1 - \epsilon, 1 + \epsilon) then

8 | C return Yes
```

In both if conditions we are checking if the matrix is ϵ -approximate doubly stochastic matrix. The moment it becomes a ϵ -approximate doubly stochastic matrix we are done.

Now if G doesn't have a perfect matching then we will never reach a ϵ -approximate doubly stochastic matrix since otherwise Hall's condition will hold and then we will have that the permanent is positive. So if G doesn't have a perfect matching the algorithm will run in an infinite loop. We only need to check if G has a perfect matching the algorithm returns Yes.

We will now define a potential function $\Phi \colon \mathbb{Z}_0 \to \mathbb{R}_+$. Let $\sigma \in S_n$ such that $a_{i,\sigma(i)} \neq 0$ for all $i \in [n]$. Now if an entry of the matrix is nonzero then it is always nonzero since all the entries are non-negative. Now since the scalings are symmetric we will define the potential function for i^{th} scaling (row/column) is $\Phi(i) = \prod_{i=1}^n a_{i,\sigma(i)}$. So we have $\Phi(0) = 1$ since at first all the entries of the matrix are from $\{0,1\}$. Also we know $\Phi(t) \leq 1$ for all t since every time we are scaling the matrix. Now $\Phi(1) \geq \frac{1}{n^n}$ since every row-sum can be at most n so it will be divided by n and therefore $a_{i,\sigma(i)} \geq \frac{1}{n}$ for all $i \in [n]$. Now to show the while loop stops if G has a perfect matching it suffices to show that $\Phi(t)$ increases by a multiplicative factor. So we have the following lemma.

```
Lemma 12.1.4 For all t, \Phi(t+1) \ge \Phi(t)(1+\alpha) for some \alpha \in (0,1).
```

Proof: Let A' denote the matrix at the t^{th} scaling where the $(t-1)^{th}$ scaling was column-scaling. Let A'' denote the matrix after row-scaling. Now since we went to the next iteration not all column-sums are in $(1-\epsilon, 1+\epsilon)$ after scaling the rows. Now the row sums of A'' are 1. Therefore we have

$$\frac{\Phi(t)}{\Phi(t+1)} = \prod_{i=1}^{n} Col\text{-}sum_{i}(A'') \leq \left(\frac{\sum\limits_{i=1}^{n} Col\text{-}sum_{i}(A'')}{n}\right)^{n} = \left(\frac{\sum\limits_{i=1}^{n} Row\text{-}sum_{i}(A'')}{n}\right)^{n} = 1 \implies \Phi(t) \leq \Phi(t+1)$$

Similarly we can say the same if $(t-1)^{th}$ scaling was row-scaling. Since not all column-sums are in $(1-\epsilon, 1+\epsilon)$ we have $\sum_{i=1}^{n} (Col\text{-}sum_i(A'') - 1)^2 \ge \epsilon^2$. Therefore using Lemma 12.1.5 we have

$$\frac{\Phi(t)}{\Phi(t+1)} \le 1 - \frac{\epsilon^2}{2} \implies \Phi(t+1) \ge \left(1 + \frac{\epsilon^2}{2}\right) \Phi(t)$$

Therefore we have the lemma.

We have $\epsilon < \frac{1}{10n}$. Therefore if $t \ge 200n^4$ then we have

$$1 \ge \Phi(t) \ge \frac{1}{n^n} \left(1 + \frac{1}{200n^2} \right)^t \ge \frac{1}{n^n} e^{n^2} > 1$$

Hence the while loop will iterate for at most $200n^4$ iterations. Hence this algorithm takes $O(n^4)$ time. Hence if G has a perfect matching the algorithm runs for at most $O(n^4)$ iterations. And if G doesn't have a perfect matching then the loop never stops. So we have the new modified algorithm to prevent infinite looping:

Algorithm 20: BP-MATRIX-SCALING

```
Input: Bipartite adjacency matrix A of G = (L \cup R, E)

Output: Decide if G has a perfect matching.

1 begin

2 | \epsilon \leftarrow \frac{1}{20n}

3 | for i \in [200n^4] do

4 | A \leftarrow Scale every rows of A to make it row stochastic.

5 | if All \ column-sums are in (1 - \epsilon, 1 + \epsilon) then

6 | return Yes

7 | A \leftarrow Scale every column of A to make it column stochastic.

8 | if All \ row-sums are in (1 - \epsilon, 1 + \epsilon) then

9 | return Yes
```

We will prove the helping lemma needed to prove Lemma 12.1.4.

Lemma 12.1.5

Suppose
$$x_1, \ldots, x_n \ge 0$$
 and $\sum_{i=1}^n x_i = n$ and $\sum_{i=1}^n (1 - x_i)^2 \ge \delta$. Then

$$\prod_{i=1}^{n} x_i \le 1 - \frac{\delta}{2} + o(\delta)$$

Proof: Denote
$$\rho_i = x_i - 1$$
. So $\sum_{i=1}^n \rho_i = 0$ and $\sum_{i=1}^n \rho^2 \ge \delta$. Now

$$\log(1+\rho_i) = \sum_{i=1}^{\infty} (-1)^{j-1} \frac{\rho_i^j}{j} \implies \log(1+\rho_i) \le \rho_i - \frac{\rho_i^2}{3} + \frac{\rho_i^3}{3} \implies 1+\rho_i \le e^{\rho_i - \frac{\rho_i^2}{3} + \frac{\rho_i^3}{3}}$$

Therefore we have

$$\prod_{i=1}^{n} x_i \le \exp\left[\sum_{i=1}^{n} \rho_i - \sum_{i=1}^{n} \frac{\rho_i^2}{3} + \sum_{i=1}^{n} \frac{\rho_i^3}{3}\right] \le \exp\left[0 - \frac{\delta}{2} + \frac{\left(\sum_{i=1}^{n} \rho_i^2\right)^{\frac{3}{2}}}{3}\right] = \exp\left[-\frac{\delta}{2} + \frac{\delta^{\frac{3}{2}}}{3}\right] \le 1 - \frac{\delta}{2} + o(\delta)$$

Therefore we have the lemma.