

Universidade de Brasilia

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1 Contest 1	3
2 Mathematics 1	.bashrc 1 line
3 Data structures 3	alias comp='g++ -std=c++17 -g3 -ggdb3 -O3 -Wall -Wextra - fsanitize=address,undefined -Wshadow -Wconversion - D_GLIBCXX_ASSERTIONS -o test'
4 Dynamic Programming 4	hash.sh 3 line
5 Numerical 4	# Hashes a file, ignoring all whitespace and comments. Use for # verifying that code was correctly typed. CTRL+D to send EOF cpp -dD -P -fpreprocessed tr -d '[:space:]' md5sum cut -c
6 Number theory 4	_6
7 Combinatorial 4	troubleshoot.txt 52 line
8 Graph 4	Pre-submit: Write a few simple test cases if sample is not enough. Are time limits close? If so, generate max cases. Is the memory usage fine?
9 Geometry 8	Could anything overflow? Make sure to submit the right file.
10 Strings 8	Wrong answer: Print your solution! Print debug output, as well. Are you clearing all data structures between test cases?
11 Miscellaneous 10	
$\underline{\text{Contest}}$ (1)	Do you handle all corner cases correctly? Have you understood the problem correctly? Any uninitialized variables?
template.cpp 33 lines	Any overflows? Confusing N and M, i and j, etc.?
<pre>#include <bits stdc++.h=""> using namespace std; #define sws cin.tie(0)->sync_with_stdio(0)</bits></pre>	Are you sure your algorithm works? What special cases have you not thought of? Are you sure the STL functions you use work as you think? Add some assertions, maybe resubmit.
<pre>#define endl '\n' #define ll long long #define ld long double #define pb push_back #define ff first #define ss second #define pll pair<11, ll></pre>	Create some testcases to run your algorithm on. Go through the algorithm for a simple case. Go through this list again. Explain your algorithm to a teammate. Ask the teammate to look at your code. Go for a small walk, e.g. to the toilet. Is your output format correct? (including whitespace) Rewrite your solution from the start or let a teammate do it.
<pre>#define vll vector<ll></ll></pre>	Runtime error:
<pre>#define teto(a, b) ((a+b-1)/(b)) #define LSB(i) ((i) & -(i)) #define MSB(i) (32builtin_clz(i)) //64 - clzll #define BITS(i)builtin_popcountll(i) //count set bits</pre>	Have you tested all corner cases locally? Any uninitialized variables? Are you reading or writing outside the range of any vector? Any assertions that might fail?
<pre>mt19937 rng(chrono::steady_clock::now().time_since_epoch().</pre>	Any possible division by 0? (mod 0 for example) Any possible infinite recursion? Invalidated pointers or iterators?
<pre>#define debug(a) cerr<<#a<<": ";for(auto b:a)cerr<<b<<" ";<="" th=""><th>Are you using too much memory? Debug with resubmits (e.g. remapped signals, see Various).</th></b<<"></pre>	Are you using too much memory? Debug with resubmits (e.g. remapped signals, see Various).
<pre>template<typename a=""> void dbg(A const& a) {((cerr<<"{"<<a< th=""><th>Time limit exceeded: Do you have any possible infinite loops? What is the complexity of your algorithm?</th></a<></typename></pre>	Time limit exceeded: Do you have any possible infinite loops? What is the complexity of your algorithm?
<pre>const int MAX = 3e5+10; const int INF = INT32_MAX; const long long MOD = 1e9+7; const long long LLINF = INT64_MAX; const long double EPS = 1e-7;</pre>	Are you copying a lot of unnecessary data? (References) How big is the input and output? (consider scanf) Avoid vector, map. (use arrays/unordered_map) What do your teammates think about your algorithm?
<pre>const long double PI = acos(-1); int32_t main(){ sws;</pre>	Memory limit exceeded: What is the max amount of memory your algorithm should need? Are you clearing all data structures between test cases?

Mathematics (2)

2.1 Equations

$$ax^2 + bx + c = 0 \Rightarrow x = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}$$

The extremum is given by x = -b/2a.

$$ax + by = e$$

$$cx + dy = f$$

$$x = \frac{ed - bf}{ad - bc}$$

$$y = \frac{af - ec}{ad - bc}$$

In general, given an equation Ax = b, the solution to a variable x_i is given by

$$x_i = \frac{\det A_i'}{\det A}$$

where A'_i is A with the i'th column replaced by b.

2.2 Recurrences

If $a_n = c_1 a_{n-1} + \cdots + c_k a_{n-k}$, and r_1, \ldots, r_k are distinct roots of $x^k - c_1 x^{k-1} - \cdots - c_k$, there are d_1, \ldots, d_k s.t.

$$a_n = d_1 r_1^n + \dots + d_k r_k^n.$$

Non-distinct roots r become polynomial factors, e.g. $a_n = (d_1n + d_2)r^n$.

2.3 Trigonometry

$$\sin(v+w) = \sin v \cos w + \cos v \sin w$$
$$\cos(v+w) = \cos v \cos w - \sin v \sin w$$

$$\tan(v+w) = \frac{\tan v + \tan w}{1 - \tan v \tan w}$$
$$\sin v + \sin w = 2\sin\frac{v+w}{2}\cos\frac{v-w}{2}$$
$$\cos v + \cos w = 2\cos\frac{v+w}{2}\cos\frac{v-w}{2}$$

$$(V+W)\tan(v-w)/2 = (V-W)\tan(v+w)/2$$

where V, W are lengths of sides opposite angles v, w.

$$a\cos x + b\sin x = r\cos(x - \phi)$$

$$a\sin x + b\cos x = r\sin(x + \phi)$$

where
$$r = \sqrt{a^2 + b^2}$$
, $\phi = \operatorname{atan2}(b, a)$.

Geometry

2.4.1Triangles

Side lengths: a, b, c

Semiperimeter: $p = \frac{a+b+c}{2}$

Area: $A = \sqrt{p(p-a)(p-b)(p-c)}$

Circumradius: $R = \frac{abc}{4A}$

Inradius: $r = \frac{A}{}$

Length of median (divides triangle into two equal-area triangles): $m_a = \frac{1}{2}\sqrt{2b^2 + 2c^2 - a^2}$

Length of bisector (divides angles in two):

$$s_a = \sqrt{bc \left[1 - \left(\frac{a}{b+c}\right)^2\right]}$$

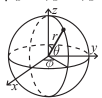
Law of sines: $\frac{\sin \alpha}{a} = \frac{\sin \beta}{b} = \frac{\sin \gamma}{c} = \frac{1}{2R}$ Law of cosines: $a^2 = b^2 + c^2 - 2bc \cos \alpha$

2.4.2 Quadrilaterals $\tan \frac{\alpha + \beta}{2}$ with of the nearths a, b, c, \overline{a} , diagonals e, f, diagonals angle θ , area A and magic flux $F = b^2 + a - 2 - c^2$:

$$4A = 2ef \cdot \sin \theta = F \tan \theta = \sqrt{4e^2f^2 - F^2}$$

2.4.3 Spherical coordinates

For cyclic quadrilaterals the sum of opposite angles is 180°, ef = ac + bd, and $A = \sqrt{(p-a)(p-b)(p-c)(p-d)}$.



$$\begin{aligned} x &= r \sin \theta \cos \phi & r &= \sqrt{x^2 + y^2 + z^2} \\ y &= r \sin \theta \sin \phi & \theta &= \arccos(z/\sqrt{x^2 + y^2 + z^2}) \\ z &= r \cos \theta & \phi &= \operatorname{atan2}(y, x) \end{aligned}$$

2.5 Derivatives/Integrals

$$\frac{d}{dx}\arcsin x = \frac{1}{\sqrt{1-x^2}} \qquad \frac{d}{dx}\arccos x = -\frac{1}{\sqrt{1-x^2}}$$

$$\frac{d}{dx}\tan x = 1 + \tan^2 x \qquad \frac{d}{dx}\arctan x = \frac{1}{1+x^2}$$

$$\int \tan ax = -\frac{\ln|\cos ax|}{a} \qquad \int x\sin ax = \frac{\sin ax - ax\cos ax}{a^2}$$

$$\int e^{-x^2} = \frac{\sqrt{\pi}}{2}\operatorname{erf}(x) \qquad \int xe^{ax}dx = \frac{e^{ax}}{a^2}(ax-1)$$

Integration by parts:

$$\int_{a}^{b} f(x)g(x)dx = [F(x)g(x)]_{a}^{b} - \int_{a}^{b} F(x)g'(x)dx$$

2.6 Sums

$$c^{a} + c^{a+1} + \dots + c^{b} = \frac{c^{b+1} - c^{a}}{c-1}, c \neq 1$$

$$1 + 2 + 3 + \dots + n = \frac{n(n+1)}{2}$$

$$1^{2} + 2^{2} + 3^{2} + \dots + n^{2} = \frac{n(2n+1)(n+1)}{6}$$

$$1^{3} + 2^{3} + 3^{3} + \dots + n^{3} = \frac{n^{2}(n+1)^{2}}{4}$$

$$1^{4} + 2^{4} + 3^{4} + \dots + n^{4} = \frac{n(n+1)(2n+1)(3n^{2} + 3n - 1)}{30}$$

2.7Series

$$e^x = 1 + x + \frac{x^2}{2!} + \frac{x^3}{3!} + \dots, (-\infty < x < \infty)$$

$$\ln(1+x) = x - \frac{x^2}{2} + \frac{x^3}{3} - \frac{x^4}{4} + \dots, (-1 < x \le 1)$$

$$\sqrt{1+x} = 1 + \frac{x}{2} - \frac{x^2}{8} + \frac{2x^3}{32} - \frac{5x^4}{128} + \dots, (-1 \le x \le 1)$$

$$\sin x = x - \frac{x^3}{3!} + \frac{x^5}{5!} - \frac{x^7}{7!} + \dots, (-\infty < x < \infty)$$

$$\cos x = 1 - \frac{x^2}{2!} + \frac{x^4}{4!} - \frac{x^6}{6!} + \dots, (-\infty < x < \infty)$$

2.8 Probability theory

Let X be a discrete random variable with probability $p_X(x)$ of assuming the value x. It will then have an expected value (mean) $\mu = \mathbb{E}(X) = \sum_{x} x p_X(x)$ and variance $\sigma^2 = V(X) = \mathbb{E}(X^2) - (\mathbb{E}(X))^2 = \sum_x (x - \mathbb{E}(X))^2 p_X(x)$ where σ is the standard deviation. If X is instead continuous it will have a probability density function $f_X(x)$ and the sums above will instead be integrals with $p_X(x)$ replaced by $f_X(x)$.

Expectation is linear:

$$\mathbb{E}(aX + bY) = a\mathbb{E}(X) + b\mathbb{E}(Y)$$

For independent X and Y,

$$V(aX + bY) = a^2V(X) + b^2V(Y).$$

2.8.1 Discrete distributions Binomial distribution

The number of successes in n independent yes/no experiments, each which yields success with probability p is $Bin(n, p), n = 1, 2, ..., 0 \le p \le 1.$

$$p(k) = \binom{n}{k} p^k (1-p)^{n-k}$$

$$\mu = np, \, \sigma^2 = np(1-p)$$

Bin(n, p) is approximately Po(np) for small p.

First success distribution

The number of trials needed to get the first success in independent yes/no experiments, each which yields success with probability p is Fs(p), $0 \le p \le 1$.

$$p(k) = p(1-p)^{k-1}, k = 1, 2, \dots$$

$$\mu = \frac{1}{n}, \, \sigma^2 = \frac{1-p}{n^2}$$

Poisson distribution

The number of events occurring in a fixed period of time t if these events occur with a known average rate κ and independently of the time since the last event is $Po(\lambda)$, $\lambda = t\kappa$.

$$p(k) = e^{-\lambda} \frac{\lambda^k}{k!}, k = 0, 1, 2, \dots$$

$$\mu = \lambda, \, \sigma^2 = \lambda$$

2.8.2 Continuous distributions Uniform distribution

If the probability density function is constant between a and b and 0 elsewhere it is U(a, b), a < b.

$$f(x) = \begin{cases} \frac{1}{b-a} & a < x < b \\ 0 & \text{otherwise} \end{cases}$$

$$\mu = \frac{a+b}{2}, \, \sigma^2 = \frac{(b-a)^2}{12}$$

Exponential distribution

The time between events in a Poisson process is $\text{Exp}(\lambda)$, $\lambda > 0$.

$$f(x) = \begin{cases} \lambda e^{-\lambda x} & x \ge 0\\ 0 & x < 0 \end{cases}$$
$$\mu = \frac{1}{\lambda}, \, \sigma^2 = \frac{1}{\lambda^2}$$

Normal distribution

Most real random values with mean μ and variance σ^2 are well described by $\mathcal{N}(\mu, \sigma^2)$, $\sigma > 0$.

$$f(x) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(x-\mu)^2}{2\sigma^2}}$$

If $X_1 \sim \mathcal{N}(\mu_1, \sigma_1^2)$ and $X_2 \sim \mathcal{N}(\mu_2, \sigma_2^2)$ then

$$aX_1 + bX_2 + c \sim \mathcal{N}(\mu_1 + \mu_2 + c, a^2\sigma_1^2 + b^2\sigma_2^2)$$

2.9 Markov chains

A Markov chain is a discrete random process with the property that the next state depends only on the current state. Let X_1, X_2, \ldots be a sequence of random variables generated by the Markov process. Then there is a transition matrix $\mathbf{P} = (p_{ij})$, with $p_{ij} = \Pr(X_n = i | X_{n-1} = j)$, and $\mathbf{p}^{(n)} = \mathbf{P}^n \mathbf{p}^{(0)}$ is the probability distribution for X_n (i.e., $p_i^{(n)} = \Pr(X_n = i)$), where $\mathbf{p}^{(0)}$ is the initial distribution.

 π is a stationary distribution if $\pi = \pi \mathbf{P}$. If the Markov chain is irreducible (it is possible to get to any state from any state), then $\pi_i = \frac{1}{\mathbb{E}(T_i)}$ where $\mathbb{E}(T_i)$ is the expected time between two visits in state i. π_j/π_i is the expected number of visits in state j between two visits in state i.

For a connected, undirected and non-bipartite graph, where the transition probability is uniform among all neighbors, π_i is proportional to node i's degree.

A Markov chain is *ergodic* if the asymptotic distribution is independent of the initial distribution. A finite Markov chain is ergodic iff it is irreducible and *aperiodic* (i.e., the gcd of cycle lengths is 1). $\lim_{k\to\infty} \mathbf{P}^k = \mathbf{1}\pi$.

A Markov chain is an A-chain if the states can be partitioned into two sets \mathbf{A} and \mathbf{G} , such that all states in \mathbf{A} are absorbing $(p_{ii}=1)$, and all states in \mathbf{G} leads to an absorbing state in \mathbf{A} . The probability for absorption in state $i \in \mathbf{A}$, when the initial state is j, is $a_{ij} = p_{ij} + \sum_{k \in \mathbf{G}} a_{ik} p_{kj}$. The expected time until absorption, when the initial state is i, is $t_i = 1 + \sum_{k \in \mathbf{G}} p_{ki} t_k$.

Data structures (3)

3.1 Ordered Set

Policy Based Data Structures (PBDS) from gcc compiler

Ordered Multiset can be created using ordered_set<pll>val, idx

order_of_key() can search for non-existent keys!

find_by_order() requires existent key and return the 0-idx position of the given value. Therefore, it returns the numbers of elements that are smaller than the given value:

ordered-set.cpp

Description: Set with index operators, implemented by gnu pbds. Remember to compile with gcc!!

Time: $\mathcal{O}(log(N))$ but with slow constant

3.2 Disjoint Set Union

There are two optional improvements:

-Tree Balancing

-Path Compression

If one improvement is used, the time complexity will become $O(\log N)$

If both are used, $O(\alpha) \approx O(5)$

dsu.cpp

Description: Disjoint Set Union with path compression and tree balancing **Time:** $\mathcal{O}(\alpha)$ 0479c4, 22 lines

```
struct DSU{
   v11 group, card;
   DSU (11 n) {
        n += 1; // 0-idx -> 1-idx
        group = v11(n);
}
```

3.3 Segment Tree

Each node of the segment tree represents the cumulative value of a range.

Observation: For some problems, such as range distinct values query, considerer offiline approach, ordering the queries by L for example.

segRecursive.cpp

Description: Basic Recursive Segment Tree for points increase and range sum query. When initializing the segmente tree, remeber to choose the range limits (L, R)

```
Time: \mathcal{O}(N \log N) to build, \mathcal{O}(\log N) to increase or query 156cd2, 70 line
```

```
// [0, n] segtree for range sum query, point increase
11 L=0, R;
struct Segtree {
    struct Node {
        // null element:
        11 ps = 0;
    };
    vector<Node> tree;
    vector<11> v;
    Segtree(ll n) {
        v.assign(n+1, 0);
        tree.assign(4*(n+1), Node{});
    Node merge (Node a, Node b) {
        return Node {
             // merge operaton:
            a.ps + b.ps
        };
    void build(l1 l=L, 11 r=R, 11 i=1 ) {
        if (1 == r) {
            tree[i] = Node {
                 // leaf element:
                v[1]
        else {
            11 \text{ mid} = (1+r)/2;
```

```
build(1, mid, 2*i);
        build(mid+1, r, 2*i+1);
        tree[i] = merge(tree[2*i], tree[2*i+1]);
void increase(11 idx=1, 11 val=0, 11 1=L, 11 r=R, 11 i=1)
    if (1 == r) {
        // increase operation:
        tree[i].ps += val;
    else {
        11 \text{ mid} = (1+r)/2;
        if (idx <= mid) increase(idx, val, 1, mid, 2*i);</pre>
        else increase(idx, val, mid+1, r, 2*i+1);
        tree[i] = merge(tree[2*i], tree[2*i+1]);
Node guery(ll left=L, ll right=R, ll l=L, ll r=R, ll i=1) {
    // left/right are the range limits for the query
    // l / r are the internal variables of the tree
    if (right < 1 or r < left) {</pre>
        // null element:
        return Node{};
    else if (left <= 1 and r <= right) return tree[i];</pre>
    else{
        11 \text{ mid} = (1+r)/2;
        return merge (
             query(left, right, 1, mid, 2*i),
             query(left, right, mid+1, r, 2*i+1)
```

3.4 Convex Hull Trick

If multiple transitions of the DP can be seen as first degree polynomials (lines). CHT can be used to optimized it

Some valid functions:

```
ax + b

cx^2 + ax + b (ignore cx^2 if c is independent)
```

cht-dynamic.cpp

Description: Dynamic version of CHT, thefore, one can insert lines in any order. There is no line removal operator

Time: $\mathcal{O}(\log N)$ per query and per insertion

707da4, 51 lines

```
// Convex Hull Trick Dinamico
//
// Para float, use LLINF = 1/.0, div(a, b) = a/b
//
// update(x) atualiza o ponto de intersecao da reta x
// overlap(x) verifica se a reta x sobrepoe a proxima
// add(a, b) adiciona reta da forma ax + b
// query(x) computa maximo de ax + b para entre as retas
// se quiser computar o minimo, eh soh fazer (-a)x + (-b)
//
// O(log(n)) amortizado por insercao
// O(log(n)) por query

struct Line {
 mutable 11 a, b, p;
```

```
bool operator<(const Line& o) const { return a < o.a; }</pre>
 bool operator<(ll x) const { return p < x; }</pre>
struct DynamicCHT : multiset<Line, less<>>> {
 11 div(ll a, ll b) {
   return a / b - ((a ^ b) < 0 and a % b);
 void update(iterator x) {
   if (next(x) == end()) x->p = LLINF;
   else if (x->a == next(x)->a) x->p = x->b >= next(x)->b?
        LLINF : -LLINF;
   else x->p = div(next(x)->b - x->b, x->a - next(x)->a);
 bool overlap(iterator x) {
   update(x);
   if (next(x) == end()) return 0;
   if (x->a == next(x)->a) return x->b >= next(x)->b;
    return x->p >= next(x)->p;
 void add(ll a, ll b) {
   auto x = insert({a, b, 0});
   while (overlap(x)) erase(next(x)), update(x);
   if (x != begin() and !overlap(prev(x))) x = prev(x), update
   while (x != begin() and overlap(prev(x)))
     x = prev(x), erase(next(x)), update(x);
 ll querv(ll x) {
   assert(!empty());
   auto 1 = *lower bound(x);
   return l.a * x + l.b;
};
```

3.5 Li-chao Tree

Works for any type of function that has the **transcending property**:

Given two functions f(x),g(x) of that type, if f(t) is greater than/smaller than g(t) for some x=t, then f(x) will be greater than/smaller than g(x) for x
eq t. In other words, once f(x) "win/lose" g(x), f(x) will continue to "win/lose" g(x).

The most common one is the line function: ax + b

Dynamic Programming (4)

Numerical (5)

Number theory (6)

Combinatorial (7)

7.1 Permutations

7.1.1 Factorial

Graph (8)

8.1 Fundamentals

8.2 Network flow

In optimization theory, maximum flow problems involve finding a feasible flow through a flow network that obtains the maximum possible flow rate.

dinic.cpp

Description: Run several bfs to compute the residual graph until a max flow configuration is discovered

Time: General Case, $\mathcal{O}\left(V^2E\right)$; Unit Capacity, $\mathcal{O}\left((V+E)\sqrt{E}\right)$; Bipartite and unit capacity, $\mathcal{O}\left((V+E)\sqrt{V}\right)$

```
// remember to duplicate vertices for the bipartite graph
//N = number of nodes, including sink and source
const 11 N = 700;
struct Dinic {
    struct Edge {
        11 from, to, flow, cap;
    vector<Edge> edges;
    vector<ll> q[N];
    11 \text{ ne} = 0, 1v1[N], vis[N], pass;
    ll qu[N], px[N], qt;
    ll run(ll s, ll sink, ll minE) {
        if (s == sink) return minE;
        11 \text{ ans} = 0;
        for(; px[s] < (int)g[s].size(); px[s]++){</pre>
            11 e = g[s][px[s]];
            auto &v = edges[e], &rev = edges[e^1];
            if( lvl[v.to] != lvl[s]+1 || v.flow >= v.cap)
            11 tmp = run(v.to, sink, min(minE, v.cap - v.flow))
```

v.flow += tmp, rev.flow -= tmp;

ans += tmp, minE -= tmp;

if (minE == 0) break;

```
return ans:
bool bfs(ll source, ll sink) {
    qt = 0;
    qu[qt++] = source;
    lvl[source] = 1;
    vis[source] = ++pass;
    for(11 i=0; i<qt; i++) {
        11 u = qu[i];
        px[u] = 0;
        if (u == sink) return 1;
        for(auto& ed :g[u]) {
            auto v = edges[ed];
            if (v.flow >= v.cap || vis[v.to] == pass)
                 continue;
            vis[v.to] = pass;
            lvl[v.to] = lvl[u]+1;
            qu[qt++] = v.to;
    return false;
11 flow(11 source, 11 sink) { // max_flow
    reset_flow();
    11 \text{ ans} = 0;
    while(bfs(source, sink))
        ans += run(source, sink, LLINF);
    return ans;
void addEdge(ll u, ll v, ll c, ll rc = 0) { // c = capacity
     , rc = retro-capacity;
    Edge e = \{u, v, 0, c\};
    edges.pb(e);
    q[u].pb(ne++);
    e = \{v, u, 0, rc\};
    edges.pb(e);
    g[v].pb(ne++);
void reset flow() {
    for (ll i=0; i<ne; i++) edges[i].flow = 0;</pre>
    memset(lvl, 0, sizeof(lvl));
    memset(vis, 0, sizeof(vis));
    memset(qu, 0, sizeof(qu));
    memset(px, 0, sizeof(px));
    qt = 0; pass = 0;
// cut set cost = minimum cost = max flow
// cut set is the set of edges that, if removed,
// will disrupt flow from source to sink and make it 0.
vector<pll> cut() {
    vector<pll> cuts;
    for (auto [from, to, flow, cap]: edges)
        if (flow == cap and vis[from] == pass and vis[to]
              pass and cap > 0)
            cuts.pb({from, to});
    return cuts:
```

};

8.2.1 Minimum Cut

In computer science and optimization theory, the max-flow min-cut theorem states that, in a flow network, the maximum amount of flow passing from the source to the sink is equal to the total weight of the edges in a minimum cut, i.e., the smallest total weight of the edges which if removed would disconnect the source from the sink.

Let's define an s-t cut $C=(S\text{-component},\,T\text{-component})$ as a partition of $V\in G$ such that source $s\in S\text{-component}$ and sink $t\in T\text{-component}$. Let's also define a cut-set of C to be the set $(u,v)\in E-u\in S\text{-component},\,v\in T\text{-component}$ such that if all edges in the cut-set of C are removed, the Max Flow from s to t is 0 (i.e., s and t are disconnected). The cost of an s-t cut C is defined by the sum of the capacities of the edges in the cut-set of C

The by-product of computing Max Flow is Min Cut! After Max Flow algorithm stops, we run graph traversal (DFS/BFS) from source s again. All reachable vertices from source s using positive weighted edges in the residual graph belong to the S-component. All other unreachable vertices belong to the T-component. All edges connecting the S-component to the T-component belong to the cut-set of C. The Min Cut value is equal to the Max Flow value. This is the minimum over all possible s-t cuts values.

8.2.2 Matching with Flow

By modeling a bipartite graph, with some Vertices (that will choose a match) to be on the L graph and some Vertices (that will be chosen) on the R. Set the correct capacities for these edges and also for the edges that connects the sink and source. After this modeling and running the dinic max flow algorithm, one will generate a possible matching (if it is possible).

A special case of matching is the perfect matching, which includes all vertices from the bipartite graph L and R.

A maximum matching has the maximum cadinality. A perfect matching is a maximum matching. But the opposite is not necessarity true.

It's possible to access dinic.edges, which is a vector that contains all edges and also its respective attributes, like the *flow* passing through each edge. Remember to consider that negative flow exist for reverse edges.

8.3 Matching

8.4 Coloring

8.5 Undirected Graph

Bridges and Articulation Points are concepts for undirected graphs!

8.5.1 Bridges (Cut Edges)

Also called **isthmus** or **cut arc**.

A back-edge is never a bridge!

A **lowlink** for a vertice U is the closest vertice to the root reachable using only span edges and a single back-edge, starting in the subtree of U.

After constructing a DFS Tree, an edge (u, v) is a bridge \iff there is no back-edge from v (or a descendent of v) to u (or an ancestor of u)

To do this efficiently, it's used tin[i] (entry time of node i) and low[i] (minimum entry time considering all nodes that can be reached from node i).

In another words, a edge (u, v) is a bridge \iff the low[v] ξ tin[u].

bridges.cpp

Description: Using the concepts of entry time (tin) and lowlink (low), an edge is a bridge if, and only if, low[v] > tin[u]

Time: $\mathcal{O}\left(V+E\right)$

87e0d3, 25 lines

```
vector<vll> q(MAX);
11 timer = 1;
11 tin[MAX], low[MAX];
vector<pll> bridges;
void dfs(ll u, ll p = -1){
    tin[u] = low[u] = timer++;
    for (auto v : g[u]) if (v != p) {
        if (tin[v]) // v was visited (\{u,v\}) is a back-edge)
            // considering a single back-edge:
            low[u] = min(low[u], tin[v]);
        else { // v wasn't visited ({u, v} is a span-edge)
            dfs(v, u);
            // after low[v] was computed by dfs(v, u):
            low[u] = min(low[u], low[v]);
            if (low[v] > tin[u])
                bridges.pb({u, v});
void findBridges(ll n) {
    for(ll i=1; i<=n; i++) if (!tin[i])</pre>
        dfs(i);
```

8.5.2 Bridge Tree

After merging *vertices* of a **2-edge connected component** into single vertices, and leaving only bridges, one can generate a Bridge Tree.

Every **2-edge connected component** has following properties:

• For each pair of vertices A, B inside the same component, there are at least 2 distinct paths from A to B (which may repeat vertices).

bridgeTree articulation blockCutTree

```
bridgeTree.cpp
```

Description: After finding bridges, set an component id for each vertice, then merge vertices that are in the same 2-edge connected component

Time: $\mathcal{O}(V+E)$

```
// g: u \Rightarrow \{v, edge id\}
vector<vector<pll>> q(MAX);
vector<vll> gc(MAX);
11 timer = 1;
11 tin[MAX], low[MAX], comp[MAX];
bool isBridge[MAX];
void dfs(ll u, ll p = -1) {
    tin[u] = low[u] = timer++;
    for(auto [v, id] : g[u]) if (v != p) {
        if (tin[v])
            low[u] = min(low[u], tin[v]);
            dfs(v, u);
            low[u] = min(low[u], low[v]);
            if (low[v] > tin[u])
                isBridge[id] = 1;
void dfs2(11 u, 11 c, 11 p = -1) {
    comp[u] = c;
    for (auto [v, id] : g[u]) if (v != p) {
       if (isBridge[id]) continue;
       if (!comp[v]) dfs2(v, c, u);
void bridgeTree(ll n) {
    // find bridges
    for(ll i=1; i<=n; i++) if (!tin[i])
        dfs(i);
    // find components
    for(ll i=1; i<=n; i++) if (!comp[i])
        dfs2(i, i);
    // condensate into a TREE (or TREES if disconnected)
    for(11 u=1; u<=n; u++) {
        for(auto [v, id] : q[u]) {
            if (comp[u] != comp[v]) {
                gc[comp[u]].pb(comp[v]);
```

8.5.3 Articulation Points

One Vertice in a graph is considered a Articulation Points or Cut Vertice if its removal in the graph will generate more

disconnected components

articulation.cpp

Description: if low[v] >= tin[u], u is an articulation points The root is a corner case

Time: $\mathcal{O}(V+E)$

```
8707a0, 29 lines
vector<vll> g(MAX);
11 timer = 1;
11 low[MAX], tin[MAX], isAP[MAX];
// when vertex i is removed from graph
// isAP[i] is the quantity of new disjoint components created
```

```
// isAP[i] >= 1 \{i \ is \ a \ Articulation \ Point\}
void dfs(ll u, ll p = -1) {
    low[u] = tin[u] = timer++;
    for (auto v : q[u]) if (v != p) {
        if (tin[v]) // visited
            low[u] = min(low[u], tin[v]);
        else { // not visited
            dfs(v, u);
            low[u] = min(low[u], low[v]);
            if (low[v] >= tin[u])
                 isAP[u]++;
    // corner case: root
    if (p == -1 \text{ and } isAP[u]) isAP[u]--;
void findAP(ll n) {
    for(ll i=1; i<=n; i++) if (!tin[i])</pre>
        dfs(i);
```

8.5.4 Block Cut Tree

After merging *edges* of a **2-vertex connected component** into single vertices, one can obtain a block cut tree.

2-vertex connected components are also called as biconnected component

Every bridge by itself is a biconnected component

Each edge in the block-cut tree connects exactly an Articulation Point and a biconnected component (bipartite graph)

Each biconnected component has the following properties:

- For each pair of edges, there is a cycle that contains both edges
- For each pair of vertices A, B inside the same connected component, there are at least 2 distinct paths from A to B (which do not repeat vertices).

blockCutTree.cpp

Description: After Merging 2-Vertex Connected Components, one can generate a block cut tree

```
Time: \mathcal{O}(V+E)
                                                     f752d5, 100 lines
// Block-Cut Tree (bruno monteiro)
// Cria a block-cut tree, uma arvore com os blocos
// e os pontos de articulação
// Blocos sao as componentes 2-vertice-conexos maximais
// Uma 2-coloração da arvore eh tal que uma cor são
  os componentes, e a outra cor sao os pontos de articulação
// Funciona para grafo nao conexo
// isAP[i] responde o numero de novas componentes conexas
// criadas apos a remocao de i do grafo g
// Se isAP[i] >= 1, i eh ponto de articulação
''// Para todo i < blocks.size()
```

```
// blocks[i] eh uma componente 2-vertce-conexa maximal
// blockEdges[i] sao as arestas do bloco i
// tree eh a arvore block-cut-tree
// tree[i] eh um vertice da arvore que corresponde ao bloco i
// comp[i] responde a qual vertice da arvore vertice i pertence
// Arvore tem no maximo 2n vertices
// O(n+m)
// 0-idx graph!!!
vector<vll> g(MAX), tree, blocks;
vector<vector<pll>>> blockEdges;
stack<11> st; // st for vertices,
stack<pl1> st2; // st2 for edges
vector<ll> low, tin, comp, isAP;
11 timer = 1;
void dfs(ll u, ll p = -1) {
    low[u] = tin[u] = timer++;
    st.push(u);
    // add only back-edges to stack
    if (p != -1) st2.push({u, p});
    for (auto v : g[u]) if (v != p) {
        if (tin[v] != -1) // visited
            st2.push({u, v});
    for (auto v : g[u]) if (v != p) {
        if (tin[v] != -1) // visited
            low[u] = min(low[u], tin[v]);
        else { // not visited
            dfs(v, u);
            low[u] = min(low[u], low[v]);
            if (low[v] >= tin[u]) {
                isAP[u] += 1;
                blocks.pb(vll(1, u));
                while(blocks.back().back() != v)
                    blocks.back().pb(st.top()), st.pop();
                blockEdges.pb(vector<pll>(1, st2.top())), st2.
                while(blockEdges.back().back() != pair<11, 11>(
                    blockEdges.back().pb(st2.top()), st2.pop();
    // corner case: root
    if (p == -1 \text{ and } isAP[u]) isAP[u]--;
void blockCutTree(ll n) {
    // initialize vectors and reset
    tree.clear(), blocks.clear(), blockEdges.clear();
    st = stack<11>(), st2 = stack<pl1>();
    tin.assign(n, -1);
    low.assign(n, 0), comp.assign(n, 0), isAP.assign(n, 0);
    timer = 1;
    // find Articulation Points
```

kruskal toposort kosaraju 2sat

```
for(ll i=0; i<n; i++) if (tin[i] == -1)
   dfs(i);
// set component id for APs
tree.assign(blocks.size(), vll());
for(ll i=0; i<n; i++) if (isAP[i])</pre>
   comp[i] = tree.size(), tree.pb(vll());
// set component id for non-APs and construct tree
for(11 u=0; u<(11)blocks.size(); u++) {
    for(auto v : blocks[u]) {
        if (!isAP[v])
            comp[v] = u;
       else
           tree[u].pb(comp[v]), tree[comp[v]].pb(u);
```

8.5.5 Minimum Spanning Tree

A minimum spanning tree (MST) or minimum weight spanning tree is a subset of the edges of a connected, edge-weighted undirected graph that connects all the vertices together, without any cycles and with the minimum possible total edge weight. That is, it is a spanning tree whose sum of edge weights is as small as possible.

kruskal.cpp

Description: Sort all edges in crescent order by weight, include all edges which joins two disconnected trees. In case of tie, choose whichever. Dont include edges that will join a already connected part of the tree.

Time: $\mathcal{O}\left(E\log E\alpha\right)$

206ba3, 21 lines

```
// use DSU struct
struct DSU{};
set<array<11, 3>> edges;
int32_t main() { sws;
   11 n, m; cin >> n >> m;
   DSU dsu(n+1);
   for(11 i=0; i<m; i++) {
       11 u, v, w; cin >> u >> v >> w;
        edges.insert({w, u, v});
   11 \min Cost = 0;
    for(auto [w, u, v] : edges) {
       if (dsu.find(u) != dsu.find(v)) {
           dsu.join(u, v);
            minCost += w;
    cout << minCost << endl;</pre>
```

Directed Graph

8.6.1 Topological Sort

Sort a directed graph with no cycles (DAG) in an order which each source of an edge is visited before the sink of this edge.

Cannot have cycles, because it would create a contradition of which vertices whould come before.

It can be done with a DFS, appending in the reverse order of transversal. Also a stack can be used to reverse order

toposort.cpp

Description: Using DFS pos order transversal and inverting the order, one can obtain the topological order

```
Time: \mathcal{O}(V+E)
                                                        75f781, 17 lines
vector<vll> q(MAX, vll());
vector<bool> vis;
vll topological;
void dfs(ll u) {
    vis[u] = 1;
    for(auto v : q[u]) if (!vis[v]) dfs(v);
    topological.pb(u);
// 1 - indexed
void topological_sort(ll n) {
    vis.assign(n+1, 0);
    topological.clear();
    for(ll i=1; i<=n; i++) if (!vis[i]) dfs(i);</pre>
```

8.6.2 Kosaraju

A Strongly Connected Component is a maximal subgraph in which every vertex is reachable from any vertex inside this same subgraph.

reverse(topological.begin(), topological.end());

A important property is that the inverted graph or transposed graph has the same SCCs as the original graph.

kosaraju.cpp

Description: By using the fact that the inverted graph has the same SCCs, just do a DFS twice to find all SCCs. A condensated graph can be created if wished. The condensated graph is a DAG!!

```
Time: \mathcal{O}(V+E)
                                                      381904, 45 lines
struct Kosaraju {
   11 n;
    vector<vll> g, gi, gc;
    vector<bool> vis;
    vector<11> comp;
   stack<ll, vll> st;
    void dfs(ll u) { // g
        vis[u] = 1;
        for(auto v : g[u]) if (!vis[v]) dfs(v);
        st.push(u);
    void dfs2(11 u, 11 c) { // gi
        comp[u] = c;
        for (auto v : gi[u]) if (comp[v] == -1) dfs2(v, c);
    Kosaraju (vector<vll> &g_)
      : g(g_), n(g_size()-1) { // 1-idx}
        gi.assign(n+1, vll());
        for(11 i=1; i<=n; i++) {
            for(auto j : g[i])
                gi[j].pb(i);
```

```
gc.assign(n+1, vll());
        vis.assign(n+1, 0);
        comp.assign(n+1, -1);
        st = stack<11, v11>();
        for(ll i=1; i<=n; i++) if (!vis[i]) dfs(i);
        while(!st.empty()) {
            auto u = st.top(); st.pop();
            if (comp[u] == -1) dfs2(u, u);
        for(11 u=1; u<=n; u++)
            for(auto v : g[u])
                if (comp[u] != comp[v])
                    gc[comp[u]].pb(comp[v]);
};
```

8.6.3 2-SAT

SAT (Boolean satisfiability problem) is NP-Complete.

2-SAT is a restriction of the SAT problem, in 2-SAT every clause has exactly two variables: $(X_1 \vee X_2) \wedge (X_2 \vee X_3)$

Every restriction or implication are represented in the graph as directed edges.

The algorithm uses kosaraju to check if any $(X \text{ and } \neg X)$ are in the same Strongly Connected Component (which implies that the problem is impossible).

If it doesn't, there is at least one solution, which can be generated using the topological sort of the same kosaraju (opting for the variables that appers latter in the sorted order)

2sat.cpp

void dfs(ll u) {

vis[u] = 1;

Description: Kosaraju to find if there are SCCs. If there are not cycles, use toposort to choose states

```
Time: \mathcal{O}(V+E)
                                                         87417c, 83 lines
// 0-idx graph !!!!
struct TwoSat {
    11 N; // needs to be the twice of the number of variables
    // node with idx \ 2x \Rightarrow variable \ x
    // node with idx 2x+1 \Rightarrow variable !x
    vector<vll> a, ai;
    // q = qraph; qi = transposed qraph (all edges are inverted
    TwoSat(ll n) { // number of variables (add +1 faor 1-idx)
        N = 2 * n;
        g.assign(N, vll());
        gi.assign(N, vll());
    11 idx; // component idx
    vector<11> comp, order; // topological order (reversed)
    vector<bool> vis, chosen;
    // chosen[x] = 0 \Rightarrow x was assigned
    // chosen [x] = 1 \Rightarrow !x was assigned
    // dfs and dfs2 are part of kosaraju algorithm
```

```
for (ll v : g[u]) if (!vis[v]) dfs(v);
        order.pb(u);
    void dfs2(11 u, 11 c) {
        comp[u] = c;
        for (ll v : gi[u]) if (comp[v] == -1) dfs2(v, c);
    bool solve() {
        vis.assign(N, 0);
        order = vector<11>();
        for (ll i = 0; i < N; i++) if (!vis[i]) dfs(i);</pre>
        comp.assign(N, -1); // comp = 0 \ can \ exist
        idx = 1:
        for(ll i=(ll)order.size()-1; i>=0; i--) {
            11 u = order[i];
            if (comp[u] == -1) dfs2(u, idx++);
        }
        chosen.assign(N/2, 0);
        for (11 i = 0; i < N; i += 2) {
             // x and !x in the same component \Rightarrow contradiction
            if (comp[i] == comp[i+1]) return false;
            chosen[i/2] = comp[i] < comp[i+1]; // choose latter</pre>
                   node
        }
        return true;
    // a (with flagA) implies \Rightarrow b (with flagB)
    void add(ll a, bool fa, ll b, bool fb) {
        // \{fa == 0\} \Rightarrow a
        // \{fa == 1\} \Rightarrow !a
        a = 2*a + fa;
        b = 2*b + fb;
        q[a].pb(b);
        gi[b].pb(a);
    // force a state for a certain variable (must be true)
    void force(ll a, bool fa) {
        add(a, fa^1, a, fa);
    // xor operation: one must exist, and only one can exist
    void exclusive(ll a, bool fa, ll b, bool fb) {
        add(a, fa^0, b, fb^1);
        add(a, fa^1, b, fb^0);
        add(b, fb^0, a, fa^1);
        add(b, fb^1, a, fa^0);
    // nand operation: no more than one can exist
    void nand(ll a, bool fa, ll b, bool fb) {
        add(a, fa^0, b, fb^1);
        add(b, fb^0, a, fa^1);
};
       Trees
Description: Solves LCA for trees
Time: \mathcal{O}(N \log(N)) to build, \mathcal{O}(\log(N)) per query
```

8.7

7afc1a, 54 lines

```
struct BinaryLifting {
    11 n, logN = 20; // \sim 1e6
    vector<vll> q;
```

```
vector<11> depth;
    vector<vll> up;
    BinaryLifting(vector<vll> &g_)
    : g(g_), n(g_size() + 1) { // 1-idx}
        depth.assign(n, 0);
        while((1 << logN) < n) logN++;</pre>
        up.assign(n, vll(logN, 0));
        build();
    void build(ll u = 1, ll p = -1) {
        for(ll i=1; i<logN; i++) {</pre>
            up[u][i] = up[up[u][i-1]][i-1];
        for (auto v : q[u]) if (v != p) {
            up[v][0] = u;
            depth[v] = depth[u] + 1;
            build(v, u);
    ll go(ll u, ll dist) { // O(log(n))
        for(ll i=logN-1; i>=0; i--) { // bigger jumps first
            if (dist & (1LL << i)) {
                u = up[u][i];
        return u;
    ll lca(ll a, ll b) { // O(log(n))
        if (depth[a] < depth[b]) swap(a, b);</pre>
        a = go(a, depth[a] - depth[b]);
        if (a == b) return a;
        for(ll i=logN-1; i>=0; i--) {
            if (up[a][i] != up[b][i]) {
                a = up[a][i];
                b = up[b][i];
        return up[a][0];
    11 lca(11 a, 11 b, 11 root) { // lca(a, b) when tree is
         rooted at 'root'
        return lca(a, b) ^lca(b, root) ^lca(a, root); //magic
};
queryTree.cpp
Description: Binary Lifting for min, max weight present in a simple path
Time: \mathcal{O}(N \log(N)) to build; \mathcal{O}(\log(N)) per query
                                                      75ba37, 67 lines
struct BinaryLifting {
    11 n, logN = 20; // \sim 1e6
   vector<vpll> q;
    vector<ll> depth;
   vector<vll> up, mx, mn;
    BinaryLifting(vector<vpll> &g )
```

: $g(g_)$, $n(g_size() + 1) { // 1-idx}$

while ((1 << logN) < n) logN++;

up.assign(n, vll(logN, 0));

depth.assign(n, 0);

```
mx.assign(n, vll(logN, -INF));
        mn.assign(n, vll(logN, INF));
        build();
    void build(ll u = 1, ll p = -1) {
        for(ll i=1; i<logN; i++) {</pre>
            mx[u][i] = max(mx[u][i-1], mx[up[u][i-1]][i-1]);
            mn[u][i] = min(mn[u][i-1], mn[up[u][i-1]);
            up[u][i] = up[up[u][i-1]][i-1];
        for(auto [v, w] : g[u]) if (v != p) {
            mx[v][0] = mn[v][0] = w;
            up[v][0] = u;
            depth[v] = depth[u] + 1;
            build(v, u);
    }
    array<11, 3> go(11 u, 11 dist) { // O(log(n))
        11 mxval = -INF, mnval = INF;
        for(11 i=logN-1; i>=0; i--) { // bigger jumps first
            if (dist & (1LL << i)) {</pre>
                mxval = max(mxval, mx[u][i]);
                mnval = min(mnval, mn[u][i]);
                u = up[u][i];
        }
        return {u, mxval, mnval};
    array<11, 3> query(11 u, 11 v) { // O(log(n))
        if (depth[u] < depth[v]) swap(u, v);</pre>
        auto [a, mxval, mnval] = go(u, depth[u] - depth[v]);
        11 b = v;
        if (a == b) return {a, mxval, mnval};
        for(ll i=logN-1; i>=0; i--) {
            if (up[a][i] != up[b][i]) {
                mxval = max(\{mxval, mx[a][i], mx[b][i]\});
                mnval = min({mnval, mn[a][i], mn[b][i]});
                a = up[a][i];
                b = up[b][i];
        mxval = max(\{mxval, mx[a][0], mx[b][0]\});
        mnval = min(\{mnval, mn[a][0], mn[b][0]\});
        return {up[a][0], mxval, mnval};
};
```

8.8Math

Geometry (9)

Strings (10)

10.1 Hashing

Hashing consists in generating a Polynomial for the string, therefore, assigning each distint string to a specific numeric value In practice, there will always be some collisions:

hashing zfunction kmp suffix-array

```
Probability of colision: =\frac{n^2}{2m}
n = Comparissons, m = mod size
```

when using multiple mods, they multiply: m = m1 * m2

hashing.cpp

Description: Create a numerical value for a string by using polynomial hashing

```
Time: \mathcal{O}(n) to build, \mathcal{O}(1) per query
// s[0]*P^n + s[1]*P^(n-1) + ... + s[n]*P^0
// 0-idx
struct Hashing {
    11 n, mod;
    string s;
    vector<11> p, h; // p = P^i, h = accumulated hash sum
    const 11 P = 31; // can be 53
    Hashing(string &s , ll m)
      : n(s_.size()), s(s_), mod(m), p(n), h(n) {
        for(11 i=0; i<n; i++)
            p[i] = (i ? P*p[i-1] : 1) % mod;
        for(11 i=0; i<n; i++)</pre>
            h[i] = (s[i] + P*(i ? h[i-1] : 0)) % mod;
    ll query(ll l, ll r) { // [l, r] inclusive (0-idx)
        ll hash = h[r] - (1 ? (p[r-1+1]*h[1-1]) % mod : 0);
        return hash < 0 ? hash + mod : hash;
};
// for codeforces:
mt19937 rng(chrono::steady_clock::now().time_since_epoch().
     count());
int32_t main() { sws;
    vector<11> mods = {
        1000000009,1000000021,1000000033,
        1000000087,1000000093,1000000097,
        1000000103,1000000123,1000000181,
        1000000207,1000000223,1000000241,
        1000000271,1000000289,1000000297
    };
    shuffle(mods.begin(), mods.end(), rng);
    string s; cin >> s;
```

10.2 Z-Function

Hashing hash(s, mods[0]);

Suppose we are given a string s of length n. The Z-function for this string is an array of length n where the i-th element is equal to the greatest number of characters starting from the position i that coincide with the first characters of s (the prefix of s)

The first element of the Z-function, z[0], is generally not well defined. This implementation assumes it as z[0] = 0. But it can also be interpreted as z[0] = n (all characters coincide).

Can be used to solve the following simples problems:

- Find all ocurrences of a pattern p in another string s. (p + '\$' + s) (z[i] == p.size())
- Find all borders. A border of a string is a prefix that is also a suffix of the string but not the whole string. For example, the borders of abcababcab are ab and abcab. (z[8] = 2, z[5] = 5) (z[i] = n-i)
- Find all period lengths of a string. A period of a string is a prefix that can be used to generate the whole string by repeating the prefix. The last repetition may be partial. For example, the periods of *abcabca* are **abc**, **abcabc** and **abcabca**.

It works because $(z[i] + i \ \xi = n)$ is the condition when the common characters of z[i] in addition to the elements already passed, exceeds or is equal to the end of the string. For example:

abaababaab z[8] = 2

abaababa is the period; the remaining (z[i] characters) are a prefix of the period; and when all these characters are combined, it can form the string (which has n characters).

zfunction.cpp

Description: For each substring starting at position i, compute the maximum match with the original prefix. z[0] = 0**Time:** $\mathcal{O}(n)$

```
vector<ll> z_function(string &s) { // O(n)
    ll n = (ll) s.length();
    vector<ll> z(n);
    for (ll i=1, l=0, r=0; i<n; i++) {
        if (i <= r) z[i] = min(r - i + 1, z[i - l]);
        while (i + z[i] < n and s[z[i]] == s[i + z[i]]) z[i]++;
        if (r < i + z[i] - 1) l = i, r = i + z[i] - 1;
    }
    return z;
}</pre>
```

10.3 KMP

KMP stands for Knuth-Morris-Pratt and computes the prefix function.

You are given a string s of length n. The prefix function for this string is defined as an array π of length n, where $\pi[i]$ is the length of the longest proper prefix of the substring $s[0\ldots i]$ which is also a suffix of this substring. A proper prefix of a string is a prefix that is not equal to the string itself. By definition, $\pi[0]=0$.

For example, prefix function of string "abcabcd" is [0,0,0,1,2,3,0], and prefix function of string "aabaaab" is [0,1,0,1,2,2,3].

```
kmp.cpp Description: Computes the prefix function Time: \mathcal{O}(n) 48408b, 13 lines
```

10.4 Suffix Array

The suffix array is the array with size n, whose values are the indexes from the longest substring (0) to the smallest substring (n) after ordering it lexicographically. Example:

Note that the length of the string i is: (s.size()-sa[i])

```
suffix-array.cpp Description: Creates the Suffix Array Time: O(N \log N)
```

49608b, 20 lines

```
vector<ll> suffixArray(string s) {
   s += "!";
   11 n = s.size(), N = max(n, 260LL);
   vector<ll> sa(n), ra(n);
   for (ll i = 0; i < n; i++) sa[i] = i, ra[i] = s[i];
   for (11 k = 0; k < n; k ? k *= 2 : k++) {
       vector<11> nsa(sa), nra(n), cnt(N);
       for (ll i = 0; i < n; i++) nsa[i] = (nsa[i]-k+n)%n, cnt
            [ra[i]]++;
       for (ll i = 1; i < N; i++) cnt[i] += cnt[i-1];
       for (ll i = n-1; i+1; i--) sa[--cnt[ra[nsa[i]]]] = nsa[
            i];
       for (ll i = 1, r = 0; i < n; i++) nra[sa[i]] = r += ra[
            sa[i]] !=
            ra[sa[i-1]] or ra[(sa[i]+k)%n] != ra[(sa[i-1]+k)%n]
                ];
       ra = nra;
       if (ra[sa[n-1]] == n-1) break;
   return vector<ll>(sa.begin()+1, sa.end());
```

UnB kasai manacher booth 10

Kasai generates an array of size n (like the suffix array), whose values indicates the length of the longest common prefix beetwen (sa[i] and sa[i+1])

kasai.cpp

Description: Creates the Longest Common Prefix array (LCP) **Time:** $O(N \log N)$

vector<1l> kasai(string s, vector<1l> sa) {
 ll n = s.size(), k = 0;
 vector<1l> ra(n), lcp(n);
 for (ll i = 0; i < n; i++) ra[sa[i]] = i;

 for (ll i = 0; i < n; i++, k -= !!k) {
 if (ra[i] == n-1) { k = 0; continue; }
 ll j = sa[ra[i]+1];
 while (i+k < n and j+k < n and s[i+k] == s[j+k]) k++;
 lcp[ra[i]] = k;
 }
 return lcp;</pre>

Problems that can be solved:

Numbers of Distinct Substrings:

• $\frac{n(n+1)}{2} - lcp[i]$ (for all i)

Longest Repeated Substring:

• biggest lcp[i]. The position can be found in sa[i]

Find how many distinct substrings there are for each len in [1:n]:

 Use delta encoding and the fact that lcp[i] counts the repeated substring between s.substr(sa[i]) and s.substr(sa[i+1]), which are the substrings corresponding to the common prefix.

Find the k-th distinct substring:

```
string s; cin >> s;
ll n = s.size();
auto sa = suffix_array(s);
auto lcp = kasai(s, sa);

ll k; cin >> k;

for(ll i=0; i<n; i++) {
    ll len = n-sa[i];
    if (k <= len) {
        cout << s.substr(sa[i], k) << endl;
        break;
    }
    k += lcp[i] - len;
}</pre>
```

10.5 Manacher

Manacher's Algorithm is used to find all palindromes in a string.

For each substring, centered at i, find the longest palindrome that can be formed.

Works best for odd size string, so we convert all string to odd ones by adding and extra characters between the original ones

Therefore, the value stored in the vector cnt is actually palindrome-len + 1.

manacher.cpp

Description: Covert String to odd length to use manacher, which computes all the maximum lengths of all palindromes in the given string

Time: $\mathcal{O}(2n)$ 0c2a2b, 46 lines struct Manacher { string s, t; vector<11> cnt; // t is the transformed string of s, with odd size Manacher(string &s_) : s(s_) { t = "#"; for(auto c : s) { t += c, t += "#"; count(); // perform manacher on the odd string cnt will give all the palindromes centered in i // for the odd string t void count() { ll n = t.size();string aux = $"$" + t + "^";$ vector<11> p(n + 2);11 1 = 1, r = 1;for(11 i = 1; i <= n; i++) { p[i] = max(0LL, min(r - i, p[1 + (r - i)]));while (aux[i - p[i]] == aux[i + p[i]]) { p[i]++; if(i + p[i] > r) { 1 = i - p[i], r = i + p[i];cnt = vector < 11 > (p.begin() + 1, p.end() - 1);

// compute a longest palindrome present in s

for(ll i=0; i<(ll)t.size(); i++) {</pre>

return s.substr(pos/2 - len/2, len);

11 sz = cnt[i]-1;

len = sz;

pos = i:

if (sz > len) {

10.6 Booth

};

string getLongest() {
 ll len = 0, pos = 0;

An efficient algorithm which uses a modified version of KMP to compute the least amount of rotation needed to reach the lexicographically minimal string rotation.

A rotation of a string can be generated by moving characters one after another from beginning to end. For example, the rotations of *acab* are *acab*, *caba*, *abac*, and *baca*.

booth.cpp

Description: Use a modified version of KMP to find the lexicographically minimal string rotation

Time: $\mathcal{O}\left(n\right)$

64184b, 30 lines

```
// Booth Algorithm
ll least_rotation(string &s) { // O(n)
    11 n = s.length();
    vector<11> f(2*n, -1);
    11 k = 0;
    for(ll j=1; j<2*n; j++) {
        11 i = f[j-k-1];
        while (i != -1 and s[j % n] != s[(k+i+1) % n] ) {
            if (s[j % n] < s[(k+i+1) % n])
                k = j - i - 1;
            i = f[i];
        if (i == -1 \text{ and } s[j % n] != s[(k+i+1) % n]) {
            if (s[j % n] < s[(k+i+1) % n])
                k = j;
            f[j - k] = -1;
        else
            f[j - k] = i + 1;
    return k:
int32_t main() { sws;
    string s; cin >> s;
    ll n = s.length();
    11 ans_idx = least_rotation(s);
    string tmp = s + s;
    cout << tmp.substr(ans_idx, n) << endl;</pre>
```

Miscellaneous (11)

Techniques (A)

techniques.txt

Combinatorics

159 lines

Recursion Divide and conquer Finding interesting points in N log N Algorithm analysis Master theorem Amortized time complexity Greedy algorithm Scheduling Max contiquous subvector sum Invariants Huffman encoding Graph theory Dynamic graphs (extra book-keeping) Breadth first search Depth first search * Normal trees / DFS trees Dijkstra's algorithm MST: Prim's algorithm Bellman-Ford Konig's theorem and vertex cover Min-cost max flow Lovasz toggle Matrix tree theorem Maximal matching, general graphs Hopcroft-Karp Hall's marriage theorem Graphical sequences Floyd-Warshall Euler cycles Flow networks * Augmenting paths * Edmonds-Karp Bipartite matching Min. path cover Topological sorting Strongly connected components Cut vertices, cut-edges and biconnected components Edge coloring * Trees Vertex coloring * Bipartite graphs (=> trees) * 3^n (special case of set cover) Diameter and centroid K'th shortest path Shortest cycle Dynamic programming Knapsack Coin change Longest common subsequence Longest increasing subsequence Number of paths in a dag Shortest path in a dag Dynprog over intervals Dynprog over subsets Dynprog over probabilities Dynprog over trees 3^n set cover Divide and conquer Knuth optimization Convex hull optimizations RMQ (sparse table a.k.a 2^k-jumps) Bitonic cycle Log partitioning (loop over most restricted)

Computation of binomial coefficients Pigeon-hole principle Inclusion/exclusion Catalan number Pick's theorem Number theory Integer parts Divisibility Euclidean algorithm Modular arithmetic * Modular multiplication * Modular inverses * Modular exponentiation by squaring Chinese remainder theorem Fermat's little theorem Euler's theorem Phi function Frobenius number Ouadratic reciprocity Pollard-Rho Miller-Rabin Hensel lifting Vieta root jumping Game theory Combinatorial games Game trees Mini-max Nim Games on graphs Games on graphs with loops Grundy numbers Bipartite games without repetition General games without repetition Alpha-beta pruning Probability theory Optimization Binary search Ternary search Unimodality and convex functions Binary search on derivative Numerical methods Numeric integration Newton's method Root-finding with binary/ternary search Golden section search Matrices Gaussian elimination Exponentiation by squaring Sorting Radix sort Geometry Coordinates and vectors * Cross product * Scalar product Convex hull Polygon cut Closest pair Coordinate-compression Ouadtrees KD-trees All segment-segment intersection Sweeping Discretization (convert to events and sweep) Angle sweeping Line sweeping Discrete second derivatives Strings Longest common substring Palindrome subsequences

Knuth-Morris-Pratt Tries Rolling polynomial hashes Suffix array Suffix tree Aho-Corasick Manacher's algorithm Letter position lists Combinatorial search Meet in the middle Brute-force with pruning Best-first (A*) Bidirectional search Iterative deepening DFS / A* Data structures LCA (2^k-jumps in trees in general) Pull/push-technique on trees Heavy-light decomposition Centroid decomposition Lazy propagation Self-balancing trees Convex hull trick (wcipeg.com/wiki/Convex_hull_trick) Monotone queues / monotone stacks / sliding queues Sliding queue using 2 stacks Persistent segment tree

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