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Contest (1)		
tei	mplate.cpp	43 lines
<pre>#include <bits stdc++.h=""></bits></pre>		

using namespace std; #define sws cin.tie(0) -> sync with stdio(0) #define endl '\n' #define 11 long long #define ld long double #define pb push back #define ff first #define ss second #define pll pair<11, 11> #define vll vector<ll> #define teto(a, b) (((a)+(b)-1)/(b))#define LSB(i) ((i) & -(i)) #define MSB(i) (63 - __builtin_clzll(i)) // $for\ ll$ #define BITS(i) __builtin_popcountll(i) template<class A> void debug(A a) { cout << "container: ";</pre> for(auto b : a) cout << b << " "; cout << endl; template<class... A> void dbg(A const&... a) { ((cout << "{" << a << "} "), ...); cout << endl: const 11 MAX = 2e5+10; const 11 MOD = 998'244'353; const 11 INF = INT32_MAX; // INT64_MAX const ld EPS = 1e-7; const ld PI = acos(-1);

#include <chrono>

```
using namespace std::chrono;
int32 t main() { sws;
    auto start = high resolution clock::now();
    // function to be timed here
    auto stop = high_resolution_clock::now();
    auto duration = duration_cast<milliseconds>(stop - start);
    cout << duration.count() << endl;</pre>
.bashrc
alias comp='g++ -std=c++17 -02 -g3 -ggdb3 -fsanitize=address,
    undefined -Wall -Wextra -Wshadow -Wconversion -o test'
hash.sh
                                                           3 lines
# 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
```

troubleshoot.txt

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? Could anything overflow? Make sure to submit the right file.

Wrong answer:

Print your solution! Print debug output, as well. Are you clearing all data structures between test cases? Can your algorithm handle the whole range of input? Read the full problem statement again. Do you handle all corner cases correctly? Have you understood the problem correctly? Any uninitialized variables? Any overflows? Confusing N and M, i and j, etc.? 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. 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. Runtime error:

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?

Any possible division by 0? (mod 0 for example) Any possible infinite recursion?

Invalidated pointers or iterators? Are you using too much memory?

Debug with resubmits (e.g. remapped signals, see Various).

Time limit exceeded:

Do you have any possible infinite loops? What is the complexity of your algorithm? 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?
Memory limit exceeded:
What is the max amount of memory your algorithm should need?
Are you clearing all data structures between test cases?
```

Data structures (2)

2.1 Stack

An optimization for std::stack is to use a std::vector as the container, instead of std::deque!

```
stack<int, vector<int>> st;
```

A stack can be used to efficiently solve the maximum rectangle in a histogram problem:

max-rectangle-histogram.cpp

Description: solves the problem of finding the maximum rectangle area in a grid setting (different widths, different heights)

Time: $\mathcal{O}(nm)$

```
// Example Problem: You are given a map of a forest where some
```

```
squares are empty and some squares have trees.
// What is the maximum area of a rectangular building that can
    be placed in the forest so that no trees must be cut down?
```

```
ll maxRectangleHistogram (vector<ll> x) { // O(n)
    // add an end point with heighh 0 to compute the last
         rectangles
    ;(0)dq.x
    11 area = 0;
    11 n = x.size();
    stack<pl1, vector<pl1>> st; // fmaxLeft, height for this
         rectangle}
    for(ll i=0; i<n; i++) {</pre>
        11 h = x[i];
        11 maxLeft = i;
        while(!st.empty() and st.top().ss >= h) {
            auto [maxLeft2, h2] = st.top(); st.pop();
            // compute the area of the de-stacked rectangle
            area = max(area, (i-maxLeft2)*h2);
            // extend current rectangle width with previous
            maxLeft = maxLeft2;
        st.push({maxLeft, h});
    return area;
int32_t main() { sws;
    11 n, m; cin >> n >> m;
```

vector<vector<ll>> grid(n, vector<ll>(m));

for(ll i=0; i<n; i++) {</pre>

for(11 j=0; j<m; j++) {

// convert the problem into N histogram subproblems, O(n m)

```
char c; cin >> c;
    if (c == '*') grid[i][j] = 0;
    else if (i == 0) grid[i][j] = 1;
    else grid[i][j] = grid[i-1][j] + 1;
}

ll area = 0;
for(ll i=0; i<n; i++) {
    area = max(area, maxRectangleHistogram(grid[i]));
}

cout << area << endl;</pre>
```

Also can be used to solve the maximum rectangle in a grid, with some blocked spots:

max-rectangle-grid.cpp

rectangle}

11 n; cin >> n;
vector<11> x;

Description: solves the problem of finding the maximum rectangle area in a histogram setting (same bottom, different heights).

Time: $\mathcal{O}(n)$ 8610da, 38

ll maxRectangleHistogram(vector<ll> x) { // O(n)

```
// Example Problem: A fence consists of n vertical boards. The width of each board is 1 and their heights may vary.
// You want to attach a rectangular advertisement to the fence.
What is the maximum area of such an advertisement?
```

```
// add an end point with heigth 0 to compute the last
    rectangles
x.pb(0);

11 area = 0;
11 n = x.size();
stack<pll, vector<pll>> st; // {maxLeft, height for this}
```

```
for(11 i=0; i<n; i++) {
    11 h = x(i];
    11 maxLeft = i;

while(!st.empty() and st.top().ss >= h) {
    auto [maxLeft2, h2] = st.top(); st.pop();

    // compute the area of the de-stacked rectangle
    area = max(area, (i-maxLeft2)*h2);
```

// extend current rectangle width with previous

```
st.push({maxLeft, h});
}
return area;
}
int32_t main() { sws;
```

for(ll i=0, a; i<n; i++) cin >> a, x.pb(a);

cout << maxRectangleHistogram(x) << endl;</pre>

maxLeft = maxLeft2;

2.2 List

std::list is a container that supports constant time insertion and removal of elements from anywhere in the container.

Adding, removing and moving the elements within the list or across several lists does not invalidate the iterators or references. An iterator is invalidated only when the corresponding element is deleted.

Element Access: O(1)

- *list.back()*
- list.front()

Modifiers: O(1)

- list.insert(itr, val) inserts val before itr and returns an itr to the inserted value
- list.erase(itr) erases the element referenced by itr and returns the itr for the next value (or .end())
- list.push_back(val)
- list.pop_back(val)
- list.push_front(val)
- list.pop_back(val)

2.3 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: $\mathcal{O}\left(log(N)\right)$ but with slow constant

2.4 Interval Set

interval-set.cpp

Description: A set that contains closed [l, r] interval which are disjoint (no intersection). This set is ordered and each interval [ll, r1] < [l2, r2] has r1 < l2. When a new interval is added, it checks which intersections will occur and rearranges the intervals.

```
Time: \mathcal{O}(log(N)) per insertion, slow constant
```

a3c7e0, 29 lines

```
// keeps track of disjoint closed intervals [l, r]
// a new interval added may replace parts of an older one
struct IntervalSet {
    using T = array<11, 3>;
    set<T> ranges;
    void add(T arr) {
        auto [1, r, k] = arr;
        while(ranges.upper_bound({r, INF, INF}) != ranges.begin
            auto itr = prev(ranges.upper_bound({r, INF, INF}));
            auto [12, r2, k2] = *itr;
            if (r2 < 1) break;
            // garantees that there is an intersection: l2 \le r
                  and r2 >= l
            ranges.erase(itr);
            if (12 <= 1-1) {
                ranges.insert({12, 1-1, k2});
            if (r+1 \le r2) {
                ranges.insert(\{r+1, r2, k2\});
        ranges.insert({1, r, k});
};
```

2.5 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.cp

Description: Disjoint Set Union with path compression and tree balancing **Time:** $\mathcal{O}\left(\alpha\right)$ 424a7d, 22 lines

```
struct DSU{
   vector<1l>   group, card;
   DSU (11 n){
        n += 1; // 0-idx -> 1-idx
        group = vector<1l>(n);
        iota(group.begin(), group.end(), 0);
        card = vll(n, 1);
   }
   ll find(ll i){
        return (i == group[i]) ? i : (group[i] = find(group[i])
        );
   }
}
```

trie seg-recursive-sum seg-recursive-minmax

```
// returns false if a and b are already in the same
         component
    bool join(ll a ,ll b) {
        a = find(a);
        b = find(b);
        if (a == b) return false;
        if (card[a] < card[b]) swap(a, b);</pre>
        card[a] += card[b];
        group[b] = a;
        return true;
};
```

Trie

Also called a **digital tree** or **prefix tree**.

Description: Creates a trie by pre-allocating the trie array, which contains the indices for the child nodes. The trie can be easily modified to support alphanumeric strings instead of binary strings.

Time: $\mathcal{O}(D)$, D = depth of trie

```
efeb7e, 40 lines
// MAX = maximum number of nodes that can be created
struct Trie{
    11 trie[MAX][26];
    bool isWordEnd[MAX];
   11 \text{ nxt} = 1, wordsCnt = 0;
    void add(string s) { // O(Depth)
        11 node = 0;
        for(auto c: s) {
            if (trie[node][c-'a'] == 0) { // create new node
                trie[node][c-'a'] = nxt++;
            node = trie[node][c-'a'];
        if(!isWordEnd[node]){
            isWordEnd[node] = true;
            wordsCnt++;
   bool find(string s, bool remove=false) { // O(Depth)
        11 \text{ node} = 0;
        for(auto c: s) {
            if(trie[node][c-'a'] == 0) {
                 return false;
            else {
                node = trie[node][c-'a'];
        if(remove and isWordEnd[node]){
            isWordEnd[node] = false;
             wordsCnt--;
        return isWordEnd[node];
};
```

Segment Trees

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.

2.7.1 Recursive SegTree

seg-recursive-sum.cpp

}

Description: Basic Recursive Segment Tree for points increase and range sum query. When initializing, choose an appropriate value for n.

```
Time: \mathcal{O}(N \log N) to build, \mathcal{O}(\log N) to increase or query
// [0, n] segtree for range sum query, point increase
// 0 or 1-idx
11 L=0, R;
struct Segtree {
    struct Node {
         // null element:
         11 ps = 0;
    vector<Node> tree;
    vector<ll> v;
    Segtree(11 n) {
        v.assign(n+1, 0);
        tree.assign(4*(n+1), Node{});
    Node merge (Node a, Node b) {
         return Node {
             // merge operation:
             a.ps + b.ps
        };
    void build(ll l=L, ll r=R, ll 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 \times i], tree[2 \times 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 query(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>
             11 \text{ mid} = (1+r)/2;
             return merge (
                 query(left, right, 1, mid, 2*i),
                  query(left, right, mid+1, r, 2*i+1)
};
```

seg-recursive-minmax.cpp

Description: Basic Recursive Segment Tree for point update, range min/max query When initializing, choose an appropriate value for n.

```
Time: \mathcal{O}(N \log N) to build, \mathcal{O}(\log N) to update or query
// [0, n] segtree for point update, range min/max guery
// 0 or 1-idx
11 L=0, R;
struct Segtree {
    struct Node {
        // null element:
        11 mn = INF, mx = -INF;
    vector<Node> tree;
    vector<ll> v;
    Segtree(ll n) {
    R = n:
        v.assign(n+1, 0);
        tree.assign(4*(n+1), Node{});
    Node merge (Node a, Node b) {
        return Node {
             // merge operation:
            min(a.mn, b.mn),
            max(a.mx, b.mx)
        };
    void build(ll l=L, ll r=R, ll i=1 ) {
        if (1 == r) {
            tree[i] = Node {
                 // leaf element:
                 v[1],
                 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 update(11 idx=1, 11 val=0, 11 l=L, 11 r=R, 11 i=1 ) {
        if (1 == r) {
             // increase operation:
             tree[i].mn = tree[i].mx = val;
        else {
```

2.7.2 Inverted Segtree

Instead of keeping the prefix sum for all the children in each node, store only the delta encoding value.

```
11 mid = (1+r)/2;
return merge(
    query(left, right, 1, mid, 2*i),
    query(left, right, mid+1, r, 2*i+1)
);
```

Therefore, to check a value in a certain position, iterate and sum all delta values from root to leaf.

seg-inverted.cpp

Description: Basic Inverted Segment Tree for point query stored value, range increase When initializing, choose an appropriate value for n. Time: $\mathcal{O}(N \log N)$ to build $\mathcal{O}(\log N)$ to proper increase or point query.

```
Time: \mathcal{O}(N \log N) to build, \mathcal{O}(\log N) to range increase or point query
// [0, n] segtree for point query stored value, range increase
// 0 or 1-idx
11 L=0, R;
struct Segtree {
    struct Node {
        // null element:
        11 ps = 0;
    };
    vector<Node> tree:
    vector<ll> v;
    Segtree(ll n) {
    R = n:
        v.assign(n+1, 0);
        tree.assign(4*(n+1), Node{});
    Node merge (Node a, Node b) {
        return Node {
             // merge operation:
            a.ps + b.ps
        };
    void build(11 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] = Node{};
void increase(ll left, ll right, ll val=0, ll l=L, ll r=R,
    ll i=1 ) {
    if (right < 1 or r < left) {</pre>
        return;
    else if (left <= l and r <= right) {
        // increase operation
        tree[i].ps += val;
    else {
        11 \text{ mid} = (1+r)/2;
        increase(left, right, val, 1, mid, 2*i);
        increase(left, right, val, mid+1, r, 2*i+1);
Node query(ll idx, ll l=L, ll r=R, ll i=1) {
    if (1 == r) {
        return tree[i];
    else {
        11 \text{ mid} = (1+r)/2;
        if (idx <= mid)</pre>
             return merge(tree[i], query(idx, 1, mid, 2*i));
             return merge(tree[i], query(idx, mid+1, r, 2*i
```

2.7.3 PA Segtree

seg-pa.cpp

Description: Seg with PA (Progressao Aritmetica / Arithmetic Progression) When initializing the segmente tree, remeber to choose the range limits (L, R) and call build()

```
Time: \mathcal{O}(N \log N) to build, \mathcal{O}(\log N) to increase or query
                                                        22f4a0, 10<u>0 lines</u>
// [0, n] segtree for range sum query, point increase
11 L=0, R;
struct SegtreePA {
    struct Node {
         // null element:
         11 ps = 0;
    };
    vector<Node> tree;
    vector<11> v;
    vector<pll> lazy; // {x, y} of {x*i + y}
    // x = razao da PA, y = constante
    SegtreePA(11 n) {
    R = n;
        v.assign(n+1, 0);
        tree.assign(4*(n+1), Node{});
        lazy.assign(4*(n+1), pll());
    Node merge (Node a, Node b) {
        return Node {
```

```
// merge operaton:
        a.ps + b.ps
   };
}
inline pll sum(pll a, pll b) {
    return {a.ff+b.ff, a.ss+b.ss};
void build(ll l=L, ll r=R, ll 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]);
    lazv[i] = \{0, 0\};
}
void prop(ll l=L, ll r=R, ll i=1) {
    auto [x, y] = lazy[i];
    if (x == 0 \text{ and } y == 0) \text{ return;}
    11 len = r-1+1;
    // (l_{val} + r_{val}) * len / 2
    Node val{ ((y + y + x*(len-1))*len) / 2 };
    tree[i] = merge(tree[i], val);
    if (1 != r) {
        11 \text{ mid} = (1+r)/2;
        lazy[2*i] = sum(lazy[2*i], lazy[i]);
        lazy[2*i+1] = sum(lazy[2*i+1], {x, y + x*(mid-1+1)}
             );
    lazv[i] = \{0, 0\};
// left/right are the range limits for the query
// l / r are the internal variables of the tree
void increase(ll left, ll right, ll x, ll y, ll l=L, ll r=R
    , 11 i=1 ) {
    prop(l, r, i);
    if (right < 1 or r < left) return;
    else if (left <= 1 and r <= right) {
        lazy[i] = \{x, y\};
        prop(1, r, i);
    else{
        11 \text{ mid} = (1+r)/2;
        increase(left, right, x, y, 1, mid, 2*i);
        ll ny = y + max( x*(mid-max(left, 1) + 1), OLL);
        increase(left, right, x, ny, mid+1, r, 2*i+1);
        tree[i] = merge(tree[2*i], tree[2*i+1]);
Node query(11 left=L, 11 right=R, 11 l=L, 11 r=R, 11 i=1) {
    prop(l, r, i);
    if (right < 1 or r < left) {</pre>
        // null element:
        return Node{};
```

Dynamic Programming (3)

3.1 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 ll 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<>>> {
 ll 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) {
```

3.2 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

3.3 SOS DP

Sum over Subsets DP (SOS DP) computes how many elements there are for each mask which are a subset of this mask.

This can be modified for other operations in which the subset contributes for the mask . *Example:*

```
1001 if a subset of 1101;
0001 if a subset of 1101;
1100 if a subset of 1101;
1101 if a subset of 1101;
```

```
10110,4

10110,3

10110,2

10110,1

10110,1

10110,0

10110,0

10110,0

10110,0

10110,0

10110,0

10110,0

10110,0

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```

sos-dp.cpp

Description: Efficiently compute a bitmask dp, in which a subset of this bitmask contributes for the value of this bitmask.

```
Time: \mathcal{O}\left(2^N N\right), N = number of bits
```

19e50a, 35 lines

```
// problem: Given a list of n integers, your task is to
     calculate for each element x:
// the number of elements y such that x \mid y = x
// the number of elements y such that x \& y = x
// the number of elements y such that x \& y != 0
const 11 LOGMAX = 20;
11 dp[1 << LOGMAX];
11 dp2[1 << LOGMAX];
int32_t main() { sws;
    11 n; cin >> n;
    vector<ll> a(n);
    for(auto &val : a) cin >> val;
    ll full = (1LL << LOGMAX) - 1;
    for(auto val : a) dp2[full^val] += 1;
    for(auto val : a) dp[val] += 1;
    for(ll b=0; b<LOGMAX; b++) {</pre>
        for(11 mask=0; mask<(1LL<<LOGMAX); mask++) {</pre>
             if (mask & (1LL << b)) {</pre>
                 dp[mask] += dp[mask ^ (1LL << b)];
                 dp2[mask] += dp2[mask ^ (1LL << b)];</pre>
    for(auto val : a) {
        cout << dp[val] << " ";
        cout << dp2[full ^ val] << " ";</pre>
        cout << n - dp[full^val] << endl;</pre>
```

Game theory (4)

4.1 Classic Game

- There are n piles (heaps), each one with x_i stones.
- Each turn, a players must remove t stones (non-zero) from a pile, turning x_i into y_i.
- The game ends when it's impossible to make any more moves and the player without moves left lose.

4.2 Bouton's Theorem

Let s be the xor-sum value of all the piles sizes, a state s=0 is a losing position and a state s!=0 is a winning position

All wining positions will have at least one valid move to turn the game into a losing position.

All losing positions will only have moves that turns the game into winning positions (except the base case when there are no piles left and the player already lost)

4.3 DAG Representation

Consider all game positions or states of the game as $\mathbf{Vertices}$ of a graph

Valid moves are the transition between states, therefore, the directed ${\bf Edges}$ of the graph

If a state has no outgoing edges, it's a dead end and a losing state (degenerated state).

If a state has only edges to winning states, therefore it is a losing state.

if a state has at least one edge that is a losing state, it is a winning state.

4.4 Sprague-Grundy Theorem

Let's consider a state u of a two-player impartial game and let v_i be the states reachable from it.

To this state, we can assign a fully equivalent game of Nim with one pile of size x. The number x is called the **Grundy value or nim-value or nimber** of the state u.

If all transitions lead to a *winning state*, the current state must be a *losing state* with nimber 0.

If at least one transition lead to a *losing state*, the current state must be a *winning state* with nimber *i*, 0.

The **MEX** operator satisfies both condition above and can be used to calculate the nim-value of a state:

 $nimber_u = MEX \text{ of all } nimber_{vi}$

Viewing the game as a DAG, we can gradually calculate the Grundy values starting from vertices without outgoing edges (nimber=0).

Note that the MEX operator **garantees** that all nim-values smaller than the considered nimber can be reached, which is essentially the nim game with a single heap with pile size = nimber.

There are only two operations that are used when considering a Sprague-Grundy game:

4.4.1 Composition

XOR operator to compose sub-games into a single composite game

When a game is played with multiple sub-games (as nim is played with multiple piles), you are actually choosing one sub-game and making a valid move there (choosing a pile and subtracting a value from it).

The final result/winner will depend on all the sub-games played. Because you need to play all games. To compute the final result, one can simply consider the XOR of the nimbers of all sub-games.

4.4.2 Decomposition

MEX operator to compute the nimber of a state that has multiple transitions to other states

A state with nimber x can be transitioned (decomposed) into all states with nimber y < x

Nevertheless a state may reach several states, only a single one will be used during the game. This shows the difference between **states** and **sub-games**: All sub-games must be played by the players, but the states of a sub-game may be ignored.

To compute the mex of a set efficiently:

mex.cpp

Description: Compute MEX efficiently by keeping track of the frequency of all existent elements and also the missing ones

Time: $\mathcal{O}(\log N)$ per addition/removal, $\mathcal{O}(\bar{1})$ to get mex value, $\mathcal{O}(N\log(N))$ to initialize

d6f2b9, 27 lines

```
struct MEX {
   map<11, 11> freq;
   set<11> missing;
    // initialize set with values up to {max_valid_value}
         inclusive
   MEX(1l max_valid_value) { // <math>O(n log(n))}
        for(ll i=0; i<=max_valid_value; i++)</pre>
            missing.insert(i);
   ll get() { // O(1)
       if (missing.empty()) return 0;
       return *missing.begin();
   void remove(ll val) { // O(log(n))
        freg[val]--;
       if (freq[val] == 0)
            missing.insert(val);
   void add(ll val) { // O(log(n))
       freq[val]++;
       if (missing.count(val))
            missing.erase(val);
};
```

4.5 Variations and Extensions

4.5.1 Nim with Increases

Consider a modification of the classical nim game: a player can now add stones to a chosen pile instead of removing.

Note that this extra rule needs to have a restriction to keep the game acyclic (finite game).

Lemma: This move is not used in a winnig strategy and can be ignored.

Proof: If a player adds t stones in a pile, the next player just needs to remove t stones from this pile.

Considering that the game is finite and this ends sooner or later.

Example: If the set of possible outcomes for a state is 0, 1, 2, 7, 8, 9. The nimber is 3, because the MEX is 3, which is the smallest nim-value you can't transition into and also you can transition to all smaller nim-values.

Note that 7, 8, 9 transitions can be ignored, because you can simply revert the play by subtracting the same amount.

4.6 Misère Game

In this version, the player who takes the last object loses. To consider this version, simply swap the winning and losing player of the normal version.

4.7 Staircase Nim

4.7.1 Description

In Staircase Nim, there is a staircase with n steps, indexed from 0 to n-1. In each step, there are zero or more coins. Two players play in turns. In his/her move, a player can choose a step (i>0) and move one or more coins to step below it (i-1). The player who is unable to make a move lose the game. That means the game ends when all the coins are in step 0.

4.7.2 Strategy

We can divide the steps into two types, odd steps, and even steps.

Now let's think what will happen if a player A move x coins from an even step(non-zero) to an odd step. Player B can always move these same x coins to another even position and **the state of odd positions aren't affected**

But if player A moves a coin from an odd step to an even step, similar logic won't work. Due to the degenerated case, there is a situation when x coins are moved from stair 1 to 0, and player B can't move these coins from stair 0 to -1 (not a valid move).

From this argument, we can agree that coins in even steps are useless, they don't interfere to decide if a game state is winning or losing.

Therefore, the staircase nim can be visualized as a simple nim game with only the odd steps.

When stones are sent from an odd step to an even step, it is the same as removing stones from a pile in a classic nim game.

And when stones are sent from even steps to odd ones, it is the same as the increasing variation described before.

UnB point line manhattanMST 7

4.8 Grundy's Game

Initially there is only one pile with x stones. Each turn, a player must divide a pile into two non-zero piles with different sizes. The player who can't do any more moves loses.

4.8.1 Degenerate (Base) States

```
x = 1 \text{ (nim-val} = 0) \text{ (losing)}
```

```
x = 2 \text{ (nim-val} = 0) \text{ (losing)}
```

4.8.2 Other States

nim-val = MEX (all transitions)

Examples

x = 3:

```
\{2, 1\} \rightarrow (0) \text{ xor } (0) \rightarrow 0

nim-val = MEX(\{0\}) = 1
```

x = 4:

```
{3, 1} \rightarrow (1) \text{ xor } (0) \rightarrow 1

nim-val = MEX({1}) = 0
```

x = 5:

```
\{4, 1\} \rightarrow (0) \text{ xor } (0) \rightarrow 0
\{3, 2\} \rightarrow (1) \text{ xor } (0) \rightarrow 1
\text{nim-val} = \text{MEX}(\{0, 1\}) = 2
```

x = 6:

```
\{5, 1\} \rightarrow (2) \text{ xor } (0) \rightarrow 2
\{4, 2\} \rightarrow (0) \text{ xor } (0) \rightarrow 0
\text{nim-val} = \text{MEX}(\{0, 2\}) = 1
```

Important observation: All nimbers for $(n \ge 2000)$ are non-zero. (missing proof here and testing for values above 1e6).

4.9 Insta-Winning States

Classic nim game: if all piles become 0, you lose. (no more moves)

Modified nim game: if any pile becomes 0, you lose.

To adapt to this version of nim game, we create insta-winning states, which represents states that have a transition to any empty pile (will instantly win). Insta-winning states must have an specific nimber so they don't conflict with other nimbers when computing. A possible solution is nimber=INF, because no other nimber will be high enough to cause conflict.

Because of this adaptation, we can now ignore states with empty piles, and consider them with (nullvalue = -1). And the (nimber = 0) now represents the states that only have transitions to insta-winning states.

After this, beside winning states and losing states, we have added two new categories of states (insta-winning and empty-pile). Notice that:

```
empty-pile <- insta-winning <- nimber(0)</pre>
```

Therefore, we have returned to the classical nim game and can proceed normally.

OBS: Empty piles (wasn't empty before) (nimber = -1) is different from Non-existent piles (never existed) (nimber = 0)

Usage Example:

https://codeforces.com/gym/101908/problem/B

4.10 References

```
https://cp-algorithms.com/game_theory/
sprague-grundy-nim.html
```

https://codeforces.com/blog/entry/66040

https://brilliant.org/wiki/nim/

return ! (*this == o);

Geometry (5)

5.1 Point Struct

point.cpp

Description: Point struct for point operations, supports floating points and integers

Time: $\mathcal{O}(1)$

```
7e11ab, 43 lines
```

```
const 1d EPS = 1e-9;
// T can be int, long long, float, double, long double
template < class T > bool eq(T a, T b) {
   if (is_integral<T>::value) return a == b;
   else return abs(a-b) <= EPS;</pre>
template<class T> struct P {
   T x, y;
   ll id; // (optional)
   P(T xx=0, T yy=0): x(xx), y(yy) {}
   P operator + (P const& o) const { return { x+o.x, y+o.y }; }
   P operator - (P const& o) const { return { x-o.x, y-o.y }; }
   P operator *(T const& t) const { return { x*t, y*t }; }
   P operator / (T const& t) const { return { x/t, y/t }; }
   T operator *(P const& o) const { return x*o.x + y*o.y; }
   T operator ^(P const& o) const { return x*o.y - y*o.x; }
   bool operator <(P const& o) const { // enables sorting, set
        return (eq(x, o.x) ? y < o.y : x < o.x);
   bool operator == (P const& o) const {
       return eq(x, o.x) and eq(y, o.y);
   bool operator !=(P const& o) const {
```

```
friend istream& operator >>(istream& in, P &p) {
    return in >> p.x >> p.y;
}

friend ostream& operator <<(ostream& out, P const& p) {
    return out << p.x << ' ' << p.y;
}

;

using point = P<11>;
// using point = P<ld>;
```

5.2 Line Struct

line.cpp

Description: Line struct for line operations

Time: $\mathcal{O}\left(1\right)$

5f33bd, 29 lines

```
template<class T> struct L {
    point p1, p2;
    T a, b, c; // ax+by+c = 0;
    // y-y1 = ((y2-y1)/(x2-x1))(x-x1)
    L(point pp1=0, point pp2=0) : p1(pp1), p2(pp2) {
       a = p1.y - p2.y;
       b = p2.x - p1.x;
       c = p1 ^ p2;
   T eval(point p) {
        return a*p.x + b*p.y + c;
    bool inside(point p) { // reta
        return eq(eval(p), T(0));
   point normal() {
        return point(a, b);
    bool insideSeg(point p) { // segmento [p1, p2]
       return ((p1-p) ^ (p2-p)) == 0 and ((p1-p) * (p2-p))
            <= 0 );
using line = L<11>;
// using line = L<ld>;
```

5.3 Manhattan Minimum Spanning Tree

Also called the rectilinear or L1 Minimum Spanning Tree problem.

manhattanMST.cpp

Description: Given \widetilde{N} points, returns up to 4*N edges, which are guaranteed to contain a minimum spanning tree for the graph with edge weights w(p, q) = -p.x - q.x - + -p.y - q.y -. Edges are in the form (distance, src, dst). Use a standard MST algorithm on the result to find the final MST. **Time:** $\mathcal{O}(N \log N)$

```
// requires point struct, at least the constructor and operator

vector<array<11, 3>> manhattanMST(vector<point> ps) {
   vector<11> id(size(ps));
   iota(id.begin(), id.end(), 0);
```

```
vector<array<11, 3>> edges;
for(11 k=0; k<4; k++) {
   sort(id.begin(), id.end(), [&](ll i, ll j) {
       return (ps[i]-ps[j]).x < (ps[j]-ps[i]).y;
   });
   map<11, 11> sweep;
   for (11 i : id) {
       for (auto it = sweep.lower_bound(-ps[i].y); it !=
           sweep.end(); sweep.erase(it++)) {
          11 j = it -> ss;
          point d = ps[i] - ps[j];
          if (d.y > d.x) break;
          edges.pb(\{d.y + d.x, i, j\});
       sweep[-ps[i].y] = i;
   x, p.y);
return edges;
```

Graph (6)

6.1 Fundamentals

```
dfs.cpp
```

Description: Simple DFS template for anonymous function Time: $\mathcal{O}(V+E)$

af867f, 19 lines

```
int32 t main() { sws;
    // compute cardinality of each subtree
    vector<vll> q(n);
    vector<ll> card(n);
    vector<bool> vis(n); // redundant here
    function<11 (11, 11)> dfs = [&](11 u, 11 p) -> 11 {
       if (vis[u]) return;
       vis[u] = 1;
        card[u] += 1;
        for (auto v : g[u]) if (v != p) {
           card[u] += dfs(v, u);
        return card[u];
   };
    dfs(1, -1);
}
```

bfs.cpp

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

Description: Simple BFS template

7bed46, 34 lines vector<vll> q(n); vector<ll> d(n); vector<bool> vis(n); void bfs(ll src, ll sink) { queue<11> q; q.push(src); d[src] = 0;vis[src] = 1;while(!q.empty()) { auto u = q.front(); q.pop();

```
// add here a special break condition if needed, ex:
    if (u == sink) break;
    for(auto v : q[u]) {
        // each v is added to queue only once
        // due to checking visited inside for (auto v : g[u
        // and setting vis[v] = 1 before pushing to queue
        if (!vis[v]) {
            vis[v] = 1;
            d[v] = d[u] + 1;
            q.push(v);
        else { // already added to queue, but there may be
             a shorter path
            d[v] = min(d[v], d[u] + 1);
}
```

6.2 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, $\mathcal{O}(V^2E)$; Unit Capacity, $\mathcal{O}((V+E)\sqrt{E})$; 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]++){
            ll e = g[s][px[s]];
            auto &v = edges[e], &rev = edges[e^1];
            if ( lvl[v.to] != lvl[s]+1 || v.flow >= v.cap)
                 continue;
            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) {
```

```
at = 0;
    qu[qt++] = source;
    lvl[source] = 1;
    vis[source] = ++pass;
    for(ll i=0; i<qt; i++) {</pre>
        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;
ll flow(ll source, ll sink) { // max_flow
    reset_flow();
    11 \text{ ans} = 0;
    while(bfs(source, sink))
        ans += run(source, sink, LLINF);
    return ans:
void addEdge(11 u, 11 v, 11 c, 11 rc = 0) { // c = capacity
     , rc = retro-capacity;
    Edge e = \{u, v, 0, c\};
    edges.pb(e);
    g[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;
    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] <</pre>
              pass and cap > 0)
            cuts.pb({from, to});
    return cuts;
```

dinitz.cpp

};

Description: This second version may be slower due to dynamic allocation, queue, etc but it's more readable, more memory efficient

Time: General Case, $\mathcal{O}(V^2E)$; Unit Capacity, $\mathcal{O}((V+E)\sqrt{E})$; Bipartite and unit capacity, $\mathcal{O}\left((V+E)\sqrt{V}\right)$ 49775d, 88 lines

```
struct Dinitz {
    struct Edge { //u \rightarrow v
```

UnB min-vertex-cover

```
11 u, v, cap, flow=0; // u is redundant, but nice for
         some problems
};
vector<Edge> edges;
vector<vector<ll>> q;
vector<ll> level, ptr;
// n need to be big enough for all nodes, including src/
     sink
ll n, src, sink;
Dinitz(11 nn, 11 s = -1, 11 t = -1) : n(nn+10) {
    src = (s == -1 ? n-2 : s);
    sink = (t == -1 ? n-1 : t);
    g.resize(n);
void addEdge(ll u, ll v, ll cap, ll rcap = 0) { // rcap =
     retrocapacity for bidiretional edges
    g[u].push_back( (ll)edges.size() );
    edges.push_back({u, v, cap});
    g[v].push_back((ll)edges.size());
    edges.push_back({v, u, rcap});
bool bfs() {
    level.assign(n, -1); // not vis
    level[src] = 0;
    queue<11> q;
    q.push(src);
    while (!q.empty()) {
        11 u = q.front(); q.pop();
        for (auto eid : g[u]) {
            auto e = edges[eid];
            if (e.flow >= e.cap or level[e.v] != -1)
                 continue:
            level[e.v] = level[u] + 1;
            q.push(e.v);
    return level[sink] != -1;
ll dfs(ll u, ll f) {
    if (f == 0 or u == sink) return f;
    for (ll &i = ptr[u]; i < (ll)g[u].size(); i++) {</pre>
        ll eid = q[u][i];
        auto &e = edges[eid];
        if(e.flow >= e.cap or level[u]+1 != level[e.v])
             continue;
        11 newf = dfs(e.v, min(f, e.cap - e.flow));
        if (newf == 0) continue;
        e.flow += newf;
        edges[eid^1].flow -= newf;
        return newf;
    return 0:
11 \text{ max flow} = 0;
11 flow(bool reset_flow = true) {
    if (reset_flow) {
        \max flow = 0;
        for(11 u=0; u<n; u++) {</pre>
            for(auto eid : g[u]) {
                auto &e = edges[eid];
                e.flow = 0;
```

```
while (bfs()) {
           ptr.assign(n, 0);
            while (ll newf = dfs(src, INF))
               max_flow += newf;
       return max flow;
    // minimum cut set cost = minimum cost = max flow
    // minimum cut set is the minimum 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 [u, v, cap, flow]: edges) {
            if (level[u] != -1 and level[v] == -1) {
                cuts.pb({u, v});
       return cuts;
};
```

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.

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

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

6.2.1 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).

6.2.2 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.

6.2.3 Minimum Vertex Cover

The **Konig's Theorem** describes an equivalence between the maximum matching problem and the minimum vertex cover problem in bipartite graphs.

Therefore, the value for the maximum flow in a bipartite graph is the same value as the number of nodes in a minimum vertex cover.

To retrieve the set of vertices of the minimum vertex cover:

- Give orientation to the edges, matched edges start from the right side of the graph to the left side, and free edges start from the left side of the graph to the right side.
- Run DFS from unmatched nodes of the left side, in this traversal some nodes will become visited, others will stay unvisited.
- The MVC nodes are the visited nodes from the right side, and unvisited nodes from the left side.

```
MVC = Visited_{Right} \cup Unvisited_{Left}
```

min-vertex-cover.cpp

Description: computes the min vertex cover for a bipartite graph matched with dinitz

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

963b5e, 55 lines

```
10
```

```
vector<vector<ll>> q(n);
set<ll> left, right; // unique
vector<bool> matched(n);
for(auto e : dinitz.edges) {
   if (e.u == dinitz.src or e.u == dinitz.sink) continue;
   if (e.v == dinitz.src or e.v == dinitz.sink) continue;
   if (e.cap > 0) { // not retro edge
        left.insert(e.u);
        right.insert(e.v);
        if (e.flow == e.cap) {
            // orient matched edges from right to left
            g[e.v].pb(e.u);
            matched[e.u] = 1;
            matched[e.v] = 1;
        else {
            // orient non-matched edges from left to right
            g[e.u].pb(e.v);
   }
};
vector<bool> vis(n, 0);
function < void (11) > dfs = [&](11 u) {
   vis[u] = 1;
    for(auto v : g[u])
        if (!vis[v])
            dfs(v);
};
for(auto 1 : left) if (!matched[1]) {
    dfs(1):
vector<11> ans:
for(auto 1 : left) if (!vis[1]) {
   ans.pb(1);
for(auto r : right) if (vis[r]) {
    ans.pb(r);
```

6.2.4 reNfakimum Independent Selecated by an offset

A **Independent Set** is a subset of nodes, in which all pairs u, v in the subset are not adjacent (There is no direct edge between nodes u and v).

A Maximum Independent Set is a $Independent\ Set$ with maximum cardinality;

The Maximum Independent Set is complementar to the Minimum Vertex Cover.

 $MaxIS = all_{Vertices} \setminus MVC$

Therefore, to acquire the Maximum Independent Set, run the MVC algorithm and subtract them from the set of vertices and it will end up with the maxIS.

6.3 Minimum Cost Matching

6.3.1 Minimum Cost with Dinitz

min-cost-dinitz.cpp

Description: change bfs to spfa to attribute a weight for the edges **Time:** SPFA is $\mathcal{O}\left(E\right)$ at average and $\mathcal{O}\left(VE\right)$ in the worst case $_{51a5b,\ 88\ lines}$

```
struct Dinitz {
   struct Edge { // u \rightarrow v
       11 u, v, cost, cap, flow=0;
   vector<Edge> edges;
   vector<vector<ll>> q;
   vector<11> dist, ptr; // uses dist instead of level
    // n need to be big enough for all nodes, including src/
   ll n, src, sink;
   Dinitz(11 nn, 11 s = -1, 11 t = -1) : n(nn+10) {
       src = (s == -1 ? n-2 : s);
       sink = (t == -1 ? n-1 : t);
       q.resize(n);
   void addEdge(ll u, ll v, ll cost, ll cap, ll rcap = 0) { //
         rcap = retrocapacity for bidiretional edges
       g[u].push_back( (ll)edges.size() );
       edges.push_back({u, v, cost, cap});
       g[v].push back( (ll)edges.size() );
       edges.push_back({v, u, -cost, rcap});
   bool spfa() {
       dist.assign(n, INF);
       vector<bool> inqueue(n, false);
       queue<11> q; q.push(src);
       dist[src] = 0;
       inqueue[src] = true;
       while (!q.empty()) {
           11 u = q.front(); q.pop();
           inqueue[u] = false;
            for (auto eid : g[u]) {
               auto const& e = edges[eid];
               if (e.flow >= e.cap) continue;
               if (dist[e.u] + e.cost < dist[e.v]) {</pre>
                    dist[e.v] = dist[e.u] + e.cost;
                    if (!inqueue[e.v]) {
                        q.push(e.v);
                        inqueue[e.v] = true;
        return dist[sink] != INF;
   11 min_cost = 0;
   ll dfs(ll u, ll f) {
       if (f == 0 or u == sink) return f;
       for (ll &i = ptr[u]; i < (ll)q[u].size();) {
           ll \ eid = g[u][i++];
```

```
auto &e = edges[eid];
        if(e.flow >= e.cap or (dist[e.u] + e.cost) != dist[
             e.vl) continue;
        11 newf = dfs(e.v, min(f, e.cap - e.flow));
        if (newf == 0) continue;
        e.flow += newf;
        edges[eid^1].flow -= newf;
        min_cost += e.cost * newf;
        return newf;
    return 0;
11 \text{ max flow} = 0;
pair<11, 11> flow(bool reset_flow_cost = true) {
    if (reset_flow_cost) {
        max_flow = 0;
        min_cost = 0;
        for(11 u=0; u<n; u++) {
            for(auto eid : g[u]) {
                auto &e = edges[eid];
                e.flow = 0;
    while (spfa()) {
        ptr.assign(n, 0);
        while (ll newf = dfs(src, INF))
            max_flow += newf;
```

6.3.2 Hungarian
Solves the Assignment Problem:

There are several standard formulations of the assignment problem (all of which are essentially equivalent). Here are some of them:

There are n jobs and n workers. Each worker specifies the amount of money they expect for a particular job. Each worker can be assigned to only one job. The objective is to assign jobs to workers in a way that minimizes the total cost.

Given an $n \times n$ matrix A, the task is to select one number from each row such that exactly one number is chosen from each column, and the sum of the selected numbers is minimized.

Given an $n \times n$ matrix A, the task is to find a permutation p of length n such that the value $\sum A[i][p[i]]$ is minimized.

Consider a complete bipartite graph with n vertices per part, where each edge is assigned a weight. The objective is to find a perfect matching with the minimum total weight.

hungarian dijkstra extendedDijkstra bellman-ford

It is important to note that all the above scenarios are "square" problems, meaning both dimensions are always equal to n. In practice, similar "rectangular" formulations are often encountered, where n is not equal to m, and the task is to select $\min(n,m)$ elements. However, it can be observed that a "rectangular" problem can always be transformed into a "square" problem by adding rows or columns with zero or infinite values, respectively.

We also note that by analogy with the search for a minimum solution, one can also pose the problem of finding a maximum solution. However, these two problems are equivalent to each other: it is enough to multiply all the weights by -1.

hungarian.cpp

Description: Solves the assignment problem

Time: $\mathcal{O}\left(n^3\right)$

```
06d970, 72 lines
```

```
// Hungaro
// Resolve o problema de assignment (matriz n x n)
// Colocar os valores da matriz em 'a' (pode < 0)
// assignment() retorna um par com o valor do
// assignment minimo, e a coluna escolhida por cada linha
// 0-idx
// O(n^3)
template<typename T> struct Hungarian {
 int n:
 vector<vector<T>> a;
 vector<T> u, v;
 vector<int> p, way;
 T inf;
  Hungarian(int n_): n(n_{-}), u(n+1), v(n+1), p(n+1), way(n+1)
   a = vector<vector<T>>(n, vector<T>(n));
   inf = numeric_limits<T>::max();
 pair<T, vector<int>> assignment() {
   for (int i = 1; i <= n; i++) {
     p[0] = i;
     int j0 = 0;
     vector<T> minv(n+1, inf);
     vector<int> used(n+1, 0);
       used[j0] = true;
       int i0 = p[j0], j1 = -1;
       T delta = inf;
        for (int j = 1; j \le n; j++) if (!used[j]) {
         T cur = a[i0-1][j-1] - u[i0] - v[j];
         if (cur < minv[j]) minv[j] = cur, way[j] = j0;</pre>
         if (minv[j] < delta) delta = minv[j], j1 = j;</pre>
        for (int j = 0; j \le n; j++)
         if (used[j]) u[p[j]] += delta, v[j] -= delta;
         else minv[j] -= delta;
        j0 = j1;
      } while (p[j0] != 0);
       int j1 = way[j0];
       p[j0] = p[j1];
       j0 = j1;
     } while (j0);
```

6.4 Coloring

6.4.1 k-Coloring

TODO: Add this blog

https://codeforces.com/blog/entry/57496 https://en.wikipedia.org/wiki/Graph_coloring https://open.kattis.com/problems/coloring

6.5 Shortest Paths

For weighted directed graphs

6.5.1 Dijkstra

Single Source and there ${\bf cannot}$ be any negative weighted edges.

dijkstra.cpp

Description: By keeping track of the distances sorted using an priority queue of candidates. if an edge can reduce the current min distance, insert into the priority queue. ONLY when the vertice is dequeued and its cost is $\langle = d|u|$, it is in fact a part of a shortest path

By inverting the sorting order, Dijkstra can be modified for the opposite operation: *longest paths*.

Furthermore, Dijkstra be extended to keep track of more information, such as:

- how many minimum-price routes are there? (modulo $10^9 + 7$)
- what is the minimum number of flights in a minimum-price route?
- what is the maximum number of flights in a minimum-price route?

extendedDijkstra.cpp

Description: Also counts the numbers of shortest paths, the minimum and maximum number of edges transversed in any shortest path.

```
Time: \mathcal{O}((V+E)\log V)
                                                      f93094, 32 lines
priority_queue<pl1, vector<pl1>, greater<pl1>> pq; // min\ pq
vector<vector<pll>>> q(MAX);
vector<11> d(MAX, INF), ways(MAX, 0), mx(MAX, -INF), mn(MAX,
    INF);
//INF = INT64\_MAX
void dijkstra(ll start) {
    pq.push({0, start});
    ways[start] = 1;
    d[start] = mn[start] = mx[start] = 0;
    while( !pq.empty() ){
       auto [p1, u] = pq.top(); pq.pop();
        if (p1 > d[u]) continue;
        for(auto [v, p2] : g[u]){
            // reset info, shorter path found, previous ones
                 are discarted
            if (d[u] + p2 < d[v]) {
                d[v] = d[u] + p2;
                ways[v] = ways[u];
                mx[v] = mx[u]+1;
                mn[v] = mn[u]+1;
                pq.push({d[v], v});
            // same distance, different path, update info
            else if (d[u] + p2 == d[v]) {
                ways[v] = (ways[v] + ways[u]) % MOD;
                mn[v] = min(mn[v], mn[u]+1);
                mx[v] = max(mx[v], mx[u]+1);
```

6.5.2 Bellman-Ford

Single Source and it **supports** negative edges

Conjecture: After at most n-1 (Vertices-1) iterations, all shortest paths will be found.

bellman-ford.cpp

Description: n-1 iterations is sufficient to find all shortest paths

bellman-ford-cycle floyd-warshall bridges bridgeTree

```
Time: \mathcal{O}(V*E) \rightarrow \mathcal{O}(N^2)
                                                          d749f1, 15 lines
using T = array<11, 3>;
vector<T> edges;
vector<ll> d(MAX, INF);
//INF = 0x3f3f3f3f3f3f3f3f3f, to avoid overflow
void BellmanFord(ll src, ll n) {
    d[src] = 0;
    for(ll i=0; i<n-1; i++) { // n-1 iterations
         for(auto [u, v, w] : edges) {
             if (d[u] + w < d[v]) {
                 d[v] = d[u] + w;
```

By iterating once more, one can check if the last iteration reduced once again any distance. If so, it means that there must be a negative cycle, because the shortest distance should have been found before elseway.

To retrieve the negative cycle itself, one can keep track of the last vertice that reaches a considered vertice

bellman-ford-cycle.cpp

Description: By using the property that n-1 iterations is sufficient to find all shortest paths in a graph that doesn't have negative cycles. Iterate n times and retrieve the path using a vector of parents

Time: $\mathcal{O}(V*E) \rightarrow \mathcal{O}(N^2)$

0506b5, 35 lines

using T = array<11, 3>;vector<T> edges; vector<ll> d(MAX, INF), p(MAX, -1); vector<11> cycle; //INF = 0x3f3f3f3f3f3f3f3f3f3f, to avoid overflow void BellmanFordCycle(ll src, ll n) { d[src] = 0;11 x = -1; // possible node inside a negative cycle for(11 i=0; i<n; i++) { // n iterations x = -1;for(auto [u, v, w] : edges) { if (d[u] + w < d[v]) { d[v] = d[u] + w;p[v] = u;x = v;// set x to a node, contained in a cycle in p[] for (11 i=0; i < n; i++) x = p[x]; 11 tmp = x;cycle.pb(tmp); tmp = p[tmp];while (tmp != x);cycle.pb(x); reverse(cycle.begin(), cycle.end());

```
6.5.3 Floyd Warshall
```

All-Pair Shortest Paths

floyd-warshall.cpp

Description: By using an auxiliar vertice, check if a smaller path exists between a pair (u, v) of vertices, if so, update minimum distance. Time: $\mathcal{O}(V^3)$

```
fa5f60, 15 lines
//N < sqr3(1e8) = 460
11 N = 200:
// d[u][v] = INF (no edge)
vector<vll> d(N+1, vll(N+1, INF));
void floydWarshall() { // O(N^3)
    for(ll i=1; i<=N; i++) d[i][i] = 0;
    for(ll aux=1; aux<=N; aux++)</pre>
        for(11 u=1; u<=N; u++)
            for(11 v=1; v<=N; v++)</pre>
                if (d[u][aux] < INF and d[v][aux] < INF)
                     d[u][v] = min(d[u][v], d[u][aux] + d[v][aux]
                          1);
```

Undirected Graph

Bridges and Articulation Points are concepts for undirected graphs!

6.6.1 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]; 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: $\mathcal{O}(V+E)$

```
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);
```

6.6.2 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.cpp

87e0d3, 25 lines

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> qc(MAX);
11 \text{ 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]);
        else {
            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(11 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] : g[u]) {
        if (comp[u] != comp[v]) {
            gc[comp[u]].pb(comp[v]);
        }
    }
}</pre>
```

6.6.3 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> q(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 // is AP[i] >= 1 {i is a Articulation Point} void dfs(ll u, ll p = -1) { low[u] = tin[u] = timer++; for (auto v : g[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 and isAP[u]) isAP[u]--;void findAP(ll n) { for(ll i=1; i<=n; i++) if (!tin[i])</pre>

6.6.4 Block Cut Tree

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

 $2\text{-}\mathrm{vertex}$ 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

 $\bf Description:$ 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> q(MAX), tree, blocks;
vector<vector<pll>>> blockEdges;
stack<ll> st; // st for vertices.
stack<pl1> st2; // st2 for edges
vector<ll> low, tin, comp, isAP;
11 \text{ 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<pl1>(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
    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);
```

6.6.5 Strong Orientation

Ofstrong, enjectation of dunundirected graph is anside ignment biridirection to the direction we include the direction we include the direction we include the direction of the direction to the effect, so we can't make the graph strongly connected.

Now consider a DFS through a bridgeless connected graph. Clearly, we will visit each vertex. And since there are no bridges, we can remove any DFS tree edge and still be able to go from below the edge to above the edge by using a path that contains at least one back edge. From this follows that from any vertex we can go to the root of the DFS tree. Also, from the root of the DFS tree we can visit any vertex we choose. We found a strong orientation!

In other words, to strongly orient a bridgeless connected graph, run a DFS on it and let the DFS tree edges point away from the DFS root and all other edges from the descendant to the ancestor in the DFS tree.

The result that bridgeless connected graphs are exactly the graphs that have strong orientations is called **Robbins' Acyclic Graph Orientation** theorem.

Problem: Given an undirected graph, your task is to choose a direction for each edge so that the resulting directed graph is acyclic.

Solution: Do a dfs tree, every span-edge is oriented according to the dfs transversal, and every back-edge is oriented contrary to 6.6.6 Minimum Spanning Tree the dfs transversal A minimum spanning tree (MST) or minimum weight spanning

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(ll 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>
```

6.7 Directed Graph

6.7.1 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 lime vector < vol < display < vol < vol
```

6.7.2 Kosaraju

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

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)

struct Kosaraju {
    11 n;
    vector<vll> g, gi, gc;
    vector<bool> vis;
    vector<ll> 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 : qi[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 : q[u])
                if (comp[u] != comp[v])
                    gc[comp[u]].pb(comp[v]);
};
```

6.7.3 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

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

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

```
Time: 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> g, gi; // g = graph; gi = transposed \ graph (all edges are inverted )
```

TwoSat(ll n) { // number of variables (add +1 faor 1-idx)

```
N = 2 * n;
    g.assign(N, vll());
    qi.assiqn(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
void dfs(ll u) {
   vis[u] = 1;
    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 : qi[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);
    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;
    g[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 6.8

lca.cpp

Description: Solves LCA for trees

```
Time: \mathcal{O}(N \log(N)) to build, \mathcal{O}(\log(N)) per query
                                                       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++;
        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 : g[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];
   ll lca(ll a, ll b, ll 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<11> depth;
    vector<vll> up, mx, mn;
    BinaryLifting(vector<vpll> &g_)
    : q(q), n(q.size() + 1) { // 1-idx}
        depth.assign(n, 0);
        while((1 << logN) < n) logN++;
        up.assign(n, vll(logN, 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]][i-1]);
            up[u][i] = up[up[u][i-1]][i-1];
        for (auto [v, w] : q[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(ll 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};
```

```
}
;
```

Mathematics (7)

7.1 Modular Arithmetic

modular.cpp

Description: mint struct for modular arithmetic operations **Time:** $\mathcal{O}(1)$ for most operations, $\mathcal{O}(log(n))$ for division and exponentiation

```
// supports operations between int/ll and mint,
// and it will return a mint object independently of the order
    of operations
template <11 P> struct Z {
   ll val:
    Z(11 a = 0) {
       val = a % P;
       if (val < 0) val += P;</pre>
    Z& operator += (Z rhs) {
       val += rhs.val;
       if (val >= P) val -= P;
        return *this;
   friend Z operator +(Z lhs, Z rhs) { return lhs += rhs; }
    Z& operator -= (Z rhs) {
       val += P - rhs.val;
       if (val >= P) val -= P;
       return *this;
    friend Z operator -(Z lhs, Z rhs) { return lhs -= rhs; }
    Z& operator *=(Z rhs) {
       val = (val * rhs.val) % P;
       return *this;
    friend Z operator *(Z lhs, Z rhs) { return lhs *= rhs; }
   Z \text{ fexp}(Z x, 11 i)  {
       if (i == 0) return 1;
       if (i == 1) return x;
       Z m = fexp(x, i/2);
       if (i & 1) return m * x;
       else return m;
   Z& operator /=(Z rhs) {
        return *this *= fexp(rhs, P-2);
    friend Z operator / (Z lhs, Z rhs) { return lhs /= rhs; }
   bool operator == (Z rhs) { return val == rhs.val; }
   bool operator !=(Z rhs) { return val != rhs.val; }
    friend ostream& operator << (ostream& out, Z a) { return out
         << a.val; }
    friend istream& operator >> (istream& in, Z& a) {
       ll x; in >> x;
       a = Z(x);
        return in;
```

```
};
using mint = Z<MOD>;
```

7.1.1 Lucas's Theorem

$$\binom{n}{m} \equiv \prod_{i=0}^{k} \binom{n_i}{m_i} \pmod{p}$$

For p prime. n_i and m_i are the coefficients of the representations of n and m in base p.

Example:

11 (in base p=3) =
$$1 \cdot 3^2 + 0 \cdot 3^1 + 2 \cdot 3^0$$

 $\Rightarrow n_2 = 1, n_1 = 0, n_0 = 2$

7.2 Combinatorics

$$\binom{n}{m} = \frac{n!}{m! \cdot (n-m)!}, \qquad 0 <= m <= n$$

$$0, \qquad otherwise$$

7.2.1 Factorial

7.2.2 Combinatorial Struct

combinatorics.cpp

 $\bf Description:$ basic operations for combinatorics problems under a certain modulo

Time: $\mathcal{O}(n)$ to construct, $\mathcal{O}(1)$ operations

d4da1d, 86 lines

```
// remeber to import mint struct !!
struct Combinatorics {
    vector<mint> fact, ifact;
    Combinatorics(ll n) : fact(n+1), ifact(n+1) { // inclusive
        fact[0] = 1;
        for (ll i=1; i<=n; i++) fact[i] = fact[i-1] * i;
        ifact[n] = 1 / fact[n];
        for (ll i=n; i>0; i--) ifact[i-1] = ifact[i] * i;
    // "Combinacao / Binomio de Newton"
    // n objects to place in k spaces
    // the order doesn't matter, so we consider the re-
         orderings
    // = n! / (k! * (n-k)!)
    mint combination(ll n, ll k) {
        if (k < 0 \text{ or } n < k) return 0;
        return fact[n] * ifact[k] * ifact[n-k];
    // "Permutacao"
```

```
// n objects to place in n spaces
    // = n!
   mint permutation(ll n) {
       if (n < 0) return 0;
       return fact[n];
       "Permutacao com repeticao"
    // n objects to place in n spaces
    // some objects are equal
    // therefore, we consider the possible re-orderings
    // = n! / (k1! k2! k3!)
    mint permutationRepetition(ll n, vector<ll> vec) {
       if (n < 0) return 0;
       mint ans = fact[n];
       for(auto val : vec) ans *= ifact[val];
       return ans;
    // "Arranjo Simples"
    // n objects to place in k spaces (k < n)
    // n * (n-1) * \dots * (n-k+1)
    // = n! / (n-k)!
    mint arrangement (ll n, ll k) {
       if (n < 0) return 0;
        return fact[n] * ifact[n-k];
    // "Pontos e Virgulas"
    // n stars to distribute among
    // k distint groups, that can contain 0, 1 or more stars
    // separated by k-1 bars
    // = (n+k-1)! / (n! * (k-1)!)
    mint starsBars(ll n, ll k) {
       if (k == 0) {
           if (n == 0) return 1;
            else return 0;
        return combination(n + k - 1, k - 1);
    // a derangement is a permutation of the elements of a set
    // in which no element appears in its original position
    // In other words, a derangement is a permutation that has
         no fixed points.
    // derangement(n) = subfactorial(n) = !n
    //!n = (n-1) * (!(n-1) + !(n-2)), for n >= 2
    // !1 = 0, !0 = 1
    vector<mint> subfact;
    void computeSubfactorials(ll n) {
        subfact.assign(n+1, 0);
       subfact[0] = 1;
       subfact[1] = 0;
        for(11 i=2; i<=n; i++) {
            subfact[i] = (i-1) * (subfact[i-1] + subfact[i-2]);
    // remeber to compute subfactorials first !!
   mint derangement(ll n) {
       if (n < 0) return 0;
        return subfact[n];
};
Combinatorics op (MAX); // MAX = inclusive max_value for fact[]
```

17

Let G be a group that acts on a set X. The Burnside Lemma states that the number of distinct orbits is equal to the average number of points fixed by an element of G.

$$T = \frac{1}{|G|} \sum_{g \in G} |\text{fix}(g)|$$

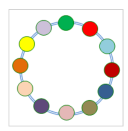
Where a orbit orb(x) is defined as

$$\mathrm{orb}(x) = \{ y \in X : \exists g \in G \ gx = y \}$$

and fix(g) is the set of elements in X fixed by g

$$fix(g) = \{x \in X : gx = x\}$$

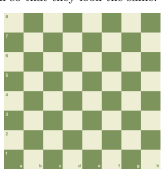
Example1: With k distinct types of beads how many distinct necklaces of size n can be made? Considering that two necklaces are equal if the rotation of one gives the other.



$$\frac{1}{n} \sum_{i=1}^{n} k^{\gcd(i,n)}$$

Example2: Count the number of different $n \times n$ grids whose each square is black or white.

Two grids are considered to be different if it is not possible to rotate one of them so that they look the same.



$$G(Rotations) = 0^{\circ}, 90^{\circ}, 180^{\circ}, 270^{\circ}$$

$$f(rotation) = \\ 0^{\circ} : 2^{(n^{2})} \\ 90^{\circ} / 270^{\circ} : 2^{\frac{n^{2}}{4}}, \qquad n_{even} \\ 2^{\frac{n^{2}-1}{4}} \cdot 2, \qquad n_{odd} \\ 180^{\circ} : 2^{\frac{n^{2}}{2}}, \qquad n_{even} \\ 2^{\frac{n^{2}-1}{2}} \cdot 2, \qquad n_{odd}$$

$$ans = \frac{1}{4}(f(0^{\circ}) + f(90^{\circ}) + f(180^{\circ}) + f(270^{\circ}))$$

7.2.4 Interesting Recursion

$$f(a,b) = f(a-1,b) + f(a,b-1)$$

$$\implies f(a,b) = \frac{(a+b)!}{a!b!} = \binom{a+b}{a}$$

Proof:

$$f(a,b) = \frac{(a+b)!}{a!b!}$$

$$\implies f(a-1,b) = \frac{(a-1+b)!}{(a-1)!b!}, f(a,b-1) = \frac{(a+b-1)!}{a!(b-1)!}$$

$$\implies f(a-1,b) + f(a,b-1) = \frac{(a-1+b)!}{(a-1)!b!} + \frac{(a+b-1)!}{a!(b-1)!}$$

$$\implies f(a,b) = (a+b-1)! \cdot \left(\frac{1}{(a-1)!(b)!} + \frac{1}{(a)!(b-1)!}\right)$$

$$\implies f(a,b) = (a+b-1)! \cdot \left(\frac{a+b}{a!b!}\right)$$

$$\implies f(a,b) = \frac{(a+b)!}{a!b!} = \binom{a+b}{a}$$

\mathbf{FFT}

FFT can be used to turn a polynomial multiplication complexity to $O(N \log N)$.

A **convulution** is easily computed by inverting one of the vector and doing the polynomial multiplication normally.

fft-simple.cpp

Description: Computes the product between two polynomials using fft Time: $\mathcal{O}(N \log N)$

```
// #define ld long double
// const ld PI = acos(-1);
struct num{
    ld a {0.0}, b {0.0};
    num(){}
    num(ld na) : a{na}{}
    num(ld na, ld nb) : a{na}, b{nb} {}
    const num operator+(const num &c) const{
        return num(a + c.a, b + c.b);
    const num operator-(const num &c) const{
        return num(a - c.a, b - c.b);
    const num operator*(const num &c) const{
        return num(a*c.a - b*c.b, a*c.b + b*c.a);
    const num operator/(const int &c) const{
         return num(a/c, b/c);
};
void fft(vector<num> &a, bool invert) {
    int n = (int)a.size();
    for (int i=1, j=0; i<n; i++) {</pre>
        int bit = n >> 1:
        for(; j&bit; bit>>=1)
            j^=bit;
        j^=bit;
        if(i<j)
            swap(a[i], a[j]);
    for(int len = 2; len <= n; len <<= 1){</pre>
        ld ang = 2 * PI / len * (invert ? -1 : 1);
        num wlen(cos(ang), sin(ang));
        for(int i=0;i<n;i+=len){</pre>
            num w(1);
            for (int j=0; j<len/2; j++) {
                 num u = a[i+j], v = a[i+j+len/2] * w;
                 a[i+j] = u + v;
                a[i+j+len/2] = u - v;
vector<1l> multiply(vector<int> const& a, vector<int> const& b)
    vector<num> fa(a.begin(), a.end());
    vector<num> fb(b.begin(), b.end());
    int n = 1;
    while(n < int(a.size() + b.size()) )</pre>
        n <<= 1;
    fa.resize(n);
    fb.resize(n);
    fft(fa, false);
    fft(fb, false);
    for (int i=0; i<n; i++)</pre>
        fa[i] = fa[i] * fb[i];
    fft(fa, true);
    vector<ll> result(n);
```

eratosthenes linear-sieve extended-euclid hashing

for(int i=0;i<n;i++)</pre> result[i] = (11) round(fa[i].a); while(result.back() == 0) result.pop back(); return result;

Number theory (8)

8.1 Sieves

These sieves are used to find all primes up to an upper bound N, which is usually 10⁷

8.1.1 Eratosthenes

Eratosthenes uses less memory than the linear sieve and is almost as fast

eratosthenes.cpp

Description: Optimized sieve of eratosthenes

Time: $\mathcal{O}(N \log \log N)$

8d74e5, 15 lines

```
// O(N \log^2(N)) \rightarrow Teorema de Merten
vector<11> primes {2, 3};
bitset<MAX> sieve; // {sieve[i] == 1} if i is prime
// MAX can be ~1e7
void eratostenes(ll n){
    sieve.set();
    for(11 i=5, step=2; i<=n; i+=step, step = 6 - step){
       if(sieve[i]){ // i is prime
            primes.pb(i);
            for(11 j= i*i; j<=n; j += 2*i) // sieving all odd
                 multiples of i >= i*i
                sieve[j] = false;
```

8.1.2 Linear Sieve

Due to the lp vector, one can compute the factorization of any number very quickly!

Can check primality with lp[i] == i

Uses more memory, because lp is a vector of int or ll and not bits.

Proof of time complexity:

We need to prove that the algorithm sets all values lp[] correctly, and that every value will be set exactly once. Hence, the algorithm will have linear runtime, since all the remaining actions of the algorithm, obviously, work for O(n).

Notice that every number i has exactly one representation in form:

$$i = lp[i] \cdot x,$$

where lp[i] is the minimal prime factor of i, and the number x doesn't have any prime factors less than lp[i], i.e.

$$lp[i] \le lp[x].$$

Now, let's compare this with the actions of our algorithm: in fact, for every x it goes through all prime numbers it could be multiplied by, i.e. all prime numbers up to lp[x] inclusive, in order to get the numbers in the form given above.

Hence, the algorithm will go through every composite number exactly once, setting the correct values lp[] there. Q.E.D.

linear-sieve.cpp

Time: $\mathcal{O}(N)$

Description: Linear Sieve that iterates every value once (prime) or twice

```
vector<ll> primes, lp(MAX);
// lp[i] = smallest prime divisor of i
void linearSieve(ll n) {
   for (11 i=2; i <= n; i++) {
       if (lp[i] == 0) { // i is prime
           lp[i] = i; // \{lp[i] = i\} for prime numbers
           primes.pb(i);
       // visit every composite number that has primes[j] as
       for (11 j = 0; i * primes[j] <= n; j++) {
           lp[i * primes[j]] = primes[j];
           if (primes[j] == lp[i])
               break;
```

Extended Euclid

Solves the ax + by = gcd(a, b) equation.

8.2.1 Inverse Multiplicative

if qcd(a,b) = 1:

then:

$$ax + by \equiv 1$$

also, if you apply \pmod{b} to the equation:

$$ax \pmod{b} + by \pmod{b} \equiv 1 \pmod{b}$$

 $ax \equiv 1 \pmod{b}$

In other words, one can find the inverse multiplicative of any number a in modulo b if gcd(a, b) = 1

8.2.2 Diofantine Equation

$$ax \equiv c \pmod{b}$$

if $q = qcd(a, b, c) \neq 1$, divide everything by q.

After this, if qcd(a,b) = 1, find a^{-1} , then multiply both sides of the Diofantine equation.

```
x \equiv c * a^{-1} \pmod{b}
```

After this, one has simply found x

```
extended-euclid.cpp
```

```
Description: Solves the a * x + b * y = gcd(a, b) equation
Time: \mathcal{O}(\log min(a,b))
```

```
// equation: a*x + b*y = qcd(a, b)
              // input: (a, b)
              // returns gcd of (a, b)
              // also computes &x and &y, which are passed by reference
2124a6, 18 lines
              ll extendedEuclid(ll a, ll b, ll &x, ll &y) {
                  x = 1, y = 0;
                  11 x1 = 0, y1 = 1, a1 = a, b1 = b;
                  while (b1) {
                      11 q = a1 / b1;
                      tie(x, x1) = pll\{x1, x - q * x1\};
                      tie(y, y1) = p11{y1, y - q * y1};
                      tie(a1, b1) = pll\{b1, a1 - q * b1\};
```

Numerical (9)

Strings (10)

return al;

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:

```
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

```
Time: \mathcal{O}(n) to build, \mathcal{O}(1) per query
```

18

60dd70, 16 lines

```
// s[0]*P^n + s[1]*P^n(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_, 11 m)
     : n(s_.size()), s(s_), mod(m), p(n), h(n) {
        for(ll i=0; i<n; i++)</pre>
            p[i] = (i ? P*p[i-1] : 1) % mod;
        for(11 i=0; i<n; i++)
            h[i] = (s[i] + P*(i ? h[i-1] : 0)) % mod;
```

```
ll query(ll 1, ll r) { // [l, r] inclusive (\theta-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<ll> 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;
    Hashing hash(s, mods[0]);
```

10.2 Z-Function

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 ithat 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 = 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
```

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

Description: For each substring starting at position i, compute the maximum match with the original prefix. z[0] = 0

```
14b37c, 12 lines
vector<11> z_function(string &s) { // O(n)
   ll n = (ll) s.length();
   vector<ll> z(n);
   for (11 i=1, 1=0, r=0; i<n; i++) {
        if (i \le r) z[i] = min(r - i + 1, z[i - 1]);
        while (i + z[i] < n \text{ and } s[z[i]] == s[i + z[i]]) z[i]++;
        if (r < i + z[i] - 1) 1 = i, r = i + z[i] - 1;
   return z;
```

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...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 "abcabed" 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
vector<11> kmp(string &s) { // O(n)
   ll n = (ll) s.length();
   vector<ll> pi(n);
    for (ll i = 1; i < n; i++) {
       11 j = pi[i-1];
       while (j > 0 \&\& s[i] != s[j])
           j = pi[j-1];
       if (s[i] == s[j])
           j++;
       pi[i] = j;
   return pi;
```

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:

```
Let the given string be "banana".
0 banana
                                  5 a
1 anana
            Sort the Suffixes
                                  3 ana
```

```
2 nana
                                   1 anana
3 ana
             alphabetically
                                   0 banana
4 na
                                   4 na
                                   2 nana
So the suffix array for "banana" is \{5, 3, 1, 0, 4, 2\}
```

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

```
suffix-array.cpp
```

Description: Creates the Suffix Array

sa[i]] !=

if (ra[sa[n-1]] == n-1) break;

return vector<ll>(sa.begin()+1, sa.end());

ra = nra;

```
Time: \mathcal{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<ll> 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[
        for (ll i = 1, r = 0; i < n; i++) nra[sa[i]] = r += ra[
```

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])

ra[sa[i-1]] or ra[(sa[i]+k)%n] != ra[(sa[i-1]+k)%n]

```
kasai.cpp
```

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

```
913195, 13 lines
vector<ll> kasai(string s, vector<ll> sa) {
    11 n = s.size(), k = 0;
    vector<ll> 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 \text{ and } j+k < n \text{ and } s[i+k] == s[j+k]) k++;
        lcp[ra[i]] = k;
    return lcp;
```

Problems that can be solved:

Numbers of Distinct Substrings:

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

Longest Repeated Substring:

manacher booth random ternary-search

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

Find how many distinct substrings there are for each len in

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

10.5Manacher

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(OLL, 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
string getLongest() {
   11 len = 0, pos = 0;
    for(ll i=0; i<(ll)t.size(); i++) {</pre>
        11 \text{ sz} = \text{cnt}[i]-1;
        if (sz > len) {
            len = sz;
            pos = i;
```

10.6 Booth

};

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.

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

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}(n)
                                                      64184b, 30 lines
// Booth Algorithm
ll least_rotation(string &s) { // O(n)
    ll n = s.length();
    vector<ll> f(2*n, -1);
    11 k = 0;
    for(11 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 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)

11.1 Random Generator

random.cpp

Description: Good randomizer to generate int in a range or shuffle vectors **Time:** $\mathcal{O}(1)$ for randint, $\mathcal{O}(nlog(n))$ for shuffle

```
mt19937 rng(chrono::steady_clock::now().time_since_epoch().
    count());
// or for 64 bits
mt19937_64 rng(chrono::steady_clock::now().time_since_epoch().
// to shuffle a vector
vector<int> vec;
shuffle(vec.begin(), vec.end(), rng);
// to limit the number to the range [l, r]
int randint(int 1, int r) {
    return (rng() % (r-1+1)) + 1;
```

11.2 Read an Fraction Input

```
char c;
11 num, den;
cin >> num >> c >> den;
```

11.3 Ternary Search

ternary-search.cpp

Description: Computes the min/max for a function that is monotonically increasing then decreasing or decreasing then increasing.

Time: $\mathcal{O}(N \log N_3)$

c3a5d7, 48 lines

```
Float and Min Version: Requires EPS (precision usually defined
     in the question text)
ld f(ld d){
    // function here
// for min value
ld ternary search(ld l, ld r){
    while (r - 1 > EPS) {
        // divide into 3 equal parts and eliminate one side
        1d m1 = 1 + (r - 1) / 3;
        1d m2 = r - (r - 1) / 3;
        if (f(m1) < f(m2)) {
            r = m2;
        else {
            1 = m1;
    return f(1); // check here for min/max
```

UnB

Techniques (A)

techniques.txt

160 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 Transform edges into vertices, duplicating the nodes of the 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 Flovd-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) Combinatorics 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 Quadratic 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 Quadtrees KD-trees All segment-segment intersection 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