WashU ICPC Notebook

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Z Function

```
* Edmonds-Karp
        Bipartite matching
        Min. path cover
        Topological sorting
        Strongly connected components
        Cutvertices, cutedges och 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 programmering
        Knapsack
        Coin change
        Longest common subsequence
        Longest increasing subsequence
        Number of paths in a dag
        Shortest path in a dag
        Dynprog over intervals
        Dynprog over subsets
        Dynprog over probabilities
        Dynprog over trees
        3^n set cover
        Divide and conquer
        Knuth optimization
        Convex hull optimizations
        RMQ (sparse table a.k.a 2^k-jumps)
        Bitonic cycle
        Log partitioning (loop over most restricted)
        Computation of binomial coefficients
        Pigeon-hole principle
        Inclusion/exclusion
        Catalan number
        Pick's theorem (Area = interior + boundary/2 - 1)
Number theory
        Integer parts
        Divisibility
        Euclidean algorithm
        Modular arithmetic
        * Modular multiplication
        * Modular inverses
        * Modular exponentiation by squaring
        Chinese remainder theorem
        Fermat's small 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
        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 polynom 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
```

2 Data Structures

2.1 Fenwick Tree (Binary Index Tree)

```
// Fenwick Tree / Binary Indexed Tree
11 bit[N];
void add(int p, int v) {
  for (p += 2; p < N; p += p & -p) bit[p] += v;
}

11 query(int p) {
    11 r = 0;
    for (p += 2; p; p -= p & -p) r += bit[p];
    return r;
}</pre>
```

2.2 Segment Tree

```
// Segment Tree (Range Query and Range Update)
// Update and Query - O(log n)
int n, v[N], lz[4*N], st[4*N];
void build(int p = 1, int l = 1, int r = n) {
  if (l == r) { st[p] = v[l]; return; }
  build(2*p, 1, (1+r)/2);
  build (2*p+1, (1+r)/2+1, r);
  st[p] = min(st[2*p], st[2*p+1]); // RMQ \rightarrow min/max, RSQ \rightarrow +
void push(int p, int 1, int r) {
  if (lz[p]) {
    st[p] = lz[p];
     // RMQ -> update: = 1z[p],
                                                increment: += lz[p]
     // RSQ -> update: = (r-1+1)*lz[p], increment: += (r-1+1)*lz[p]
     if(1!=r) lz[2*p] = lz[2*p+1] = lz[p]; // update: =, increment +=
     lz[p] = 0;
int query(int i, int j, int p = 1, int 1 = 1, int r = n) {
  \begin{array}{ll} push\,(p,\ 1,\ r)\,;\\ \mbox{if}\ (1>\ j\ \mbox{or}\ r<\ i)\ \mbox{return INF;}\ //\ \mbox{\it RMQ}\ \to\ \mbox{\it INF,}\ \mbox{\it RSQ}\ \to\ 0 \end{array}
```

2.3 Segment Tree (from Lyon)

```
//Seament Tree
#include <iostream>
#define N (1<<18)
using namespace std;
//int\ find(vector\ < int>\&\ C,\ int\ x)\ \{return\ (C[x]==x)\ ?\ x\ :\ C[x]=find(C,\ C[x]);\}\ C++
//int find(int x) {return (C[x]==x) ?x:C[x]=find(C[x]);}
typedef pair<int, int> ii;
ii arb[N]={{0,0}};
int n,m,a,b,v;
char type;
void update(int node,int l,int r,int a,int b,int val,int p)
        if(a<=1 && r<=b)
                arb[node].first=val;
                arb[node].second=p;
        else
                int mid=(1+r)/2;
                if(a<=mid)</pre>
                         update(node*2,1,mid,a,b,val,p);
                if(b>mid)
                         update(2*node+1,mid+1,r,a,b,val,p);
pair<int.int> search(int node.int l.int r.int a)
        if(a==1 && a==r)
                return arb[node];
        else
                int mid=(1+r)/2;
                if(a<=mid)
                         cur=search(2*node,1,mid,a);
                else
                         cur=search(2*node+1,mid+1,r,a);
                if (cur.second<arb[node].second)
                         return arb[node];
                else
                         return cur;
```

2.4 Sparse Table

```
const int N;
const int M; //log2(N)
int sparse[N][M];

void build() {
   for(int i = 0; i < n; i++)
        sparse[i][0] = v[i];

   for(int j = 1; j < M; j++)
        for(int i = 0; i < n; i++)
        sparse[i][j] =
        i + (1 << j - 1) < n
        ? min(sparse[i][j - 1], sparse[i + (1 << j - 1)][j - 1])
        : sparse[i][j - 1];</pre>
```

```
int query(int a, int b) {
  int pot = 32 - _builtin_clz(b - a) - 1;
  return min(sparse[a][pot], sparse[b - (1 << pot) + 1][pot]);
}</pre>
```

2.5 Treap (Cartesian Tree)

```
// Treap (probabilistic BST)
// O(logn) operations (supports lazy propagation)
mt19937 64 llrand(random device()());
struct node {
  int cnt, rev;
  int mn, mx, mindiff; // value-based treap only!
  node* 1;
 node* r;
 node(int x) : val(x), cnt(1), rev(0), mn(x), mx(x), mindiff(INF), pri(llrand()), 1(0), r(0) {}
struct treap {
 node* root;
  treap() : root(0) {}
  "treap() { clear(); }
  int cnt(node* t) { return t ? t->cnt : 0; }
  int mn (node* t) { return t ? t->mn : INF; }
  int mx (node* t) { return t ? t->mx : -INF; }
  int mindiff(node* t) { return t ? t->mindiff : INF; }
  void clear() { del(root); }
  void del(node* t) {
    if (!t) return;
    del(t->1); del(t->r);
    delete t;
    t = 0:
  void push(node* t) {
    if (!t or !t->rev) return;
    swap(t->1, t->r);
    if (t->1) t->1->rev ^= 1;
    if (t->r) t->r->rev ^= 1;
    t->rev = 0;
  void update(node*& t) {
    if (!t) return;
    t \rightarrow cnt = cnt(t \rightarrow 1) + cnt(t \rightarrow r) + 1:
    t->mn = min(t->val, min(mn(t->l), mn(t->r)));
    t\rightarrow mx = max(t\rightarrow val, max(mx(t\rightarrow l), mx(t\rightarrow r)));
    t-> mindiff = min(mn(t->r) - t->val, min(t->val - mx(t->l), min(mindiff(t->l), mindiff(t->r))));
  node* merge(node* 1, node* r) {
    push(1); push(r);
    if (!1 or !r) t = 1 ? 1 : r;
    else if (1->pri > r->pri) 1->r = merge(1->r, r), t = 1;
    else r->1 = merge(1, r->1), t = r;
    update(t):
    return t;
  // pos: amount of nodes in the left subtree or
  // the smallest position of the right subtree in a 0-indexed array
  pair<node*, node*> split(node* t, int pos) {
    if (!t) return {0, 0};
    push(t);
    if (cnt(t->1) < pos) {
     auto x = split(t->r, pos-cnt(t->1)-1);
      t->r = x.st;
      update(t);
      return { t, x.nd };
    auto x = split(t->1, pos);
    t->1 = x.nd;
    update(t);
    return { x.st, t };
```

```
// Position-based treap
// used when the values are just additional data
// the positions are known when it's built, after that you
// query to get the values at specific positions
// 0-indexed array!
void insert(int pos, int val) {
 push (root);
 node* x = new node(val);
 auto t = split(root, pos);
 root = merge(merge(t.st, x), t.nd);
void erase(int pos) {
 auto t1 = split(root, pos);
 auto t2 = split(t1.nd, 1);
 delete t2.st;
 root = merge(t1.st, t2.nd);
int get_val(int pos) { return get_val(root, pos); }
int get_val(node* t, int pos) {
 push(t);
 if (cnt(t->1) == pos) return t->val;
 if (cnt(t->1) < pos) return get_val(t->r, pos-cnt(t->1)-1);
return get_val(t->1, pos);
// Value-based treap
// used when the values needs to be ordered
int order(node* t, int val) {
 if (!t) return 0;
 push(t);
 if (t->val < val) return cnt(t->1) + 1 + order(t->r, val);
 return order(t->1, val);
bool has(node* t, int val) {
 if (!t) return 0;
 push(t);
 if (t->val == val) return 1;
 return has((t->val > val ? t->1 : t->r), val);
void insert(int val) {
 if (has(root, val)) return; // avoid repeated values
 push(root);
 node* x = new node(val);
 auto t = split(root, order(root, val));
 root = merge(merge(t.st, x), t.nd);
void erase(int val) {
 if (!has(root, val)) return;
 auto t1 = split(root, order(root, val));
 auto t2 = split(t1.nd, 1);
 delete t2.st;
 root = merge(t1.st, t2.nd);
// Get the maximum difference between values
int querymax(int i, int j) {
 if (i == j) return -1;
 auto t1 = split(root, j+1);
 auto t2 = split(t1.st, i);
 int ans = mx(t2.nd) - mn(t2.nd);
 root = merge(merge(t2.st, t2.nd), t1.nd);
 return ans;
// Get the minimum difference between values
int querymin(int i, int j) {
 if (i == j) return -1;
 auto t2 = split(root, j+1);
 auto t1 = split(t2.st, i);
 int ans = mindiff(t1.nd);
 root = merge(merge(t1.st, t1.nd), t2.nd);
 return ans:
void reverse(int 1, int r) {
 auto t2 = split(root, r+1);
 auto t1 = split(t2.st, 1);
 t1.nd->rev = 1;
```

```
root = merge(merge(t1.st, t1.nd), t2.nd);
}
void print() { print(root); printf("\n"); }
void print(node+ t) {
   if (!t) return;
   push(t);
   print(t->1);
   print("%d ", t->val);
   print(t->r);
};
};
```

2.6 Cartesian Tree (from Lyon)

```
struct Tr
        Tr *1, *r;
        int key,pr,cnt,val,rev;
        long long sum;
        Tr(int new_key,int new_pr,int new_val)
                rev=0;
                key=new_key;
                cnt=1:
                1=r=NULL:
                pr=new pr;
                val=new val;
                sum=new_val;
};
#define T Tr*
T R=NULL:
int cnt(T t)
        if(!t) return 0:
        return t->cnt;
void upd_cnt(T &t)
        if(t) t->ent=ent(t->1)+ent(t->r)+1;
long long sum(T t)
        if(!t) return 0;
        return t->sum;
void upd sum(T &t)
    if(t) t->sum=sum(t->1)+sum(t->r)+t->val;
void push (T &t)
        if(t && t->rev)
                swap(t->1,t->r);
                upd_sum(t);
                if(t->1) t->1->rev^=1;
                if(t->r) t->r->rev^=1;
void split(T t,T &l,T &r,int key,int add)
                return void(l=r=NULL);
        push(t);
        upd_cnt(t);
        int current_key=add+cnt(t->1)+1;
        if(key<=current_key)</pre>
               split(t->1,1,t->1,key,add),r=t;
                split(t->r,t->r,r,key,current_key),l=t;
        upd_cnt(t);
        upd_sum(t);
void merge(T &t,T 1,T r)
```

```
push(1);
        push(r);
        if(!1 || !r)
                t=1?1:r;
        else if(l->pr>r->pr)
                merge(1->r,1->r,r), t=1;
                merge(r->1,1,r->1), t=r;
        upd_cnt(t);
        upd_sum(t);
void insert(T &t,T it,int add)
        push(t);
        if(!t)
                 upd_cnt(t);
        upd_sum(t);
        if(it->pr > t->pr)
                 split(t,it->1,it->r,it->key,add),t=it;
        else if(it->key>add+cnt(t->1)+1)
                 insert(t->r, it, add+cnt(t->1)+1);
        else
                insert(t->1,it,add);
        upd sum(t):
        upd_cnt(t);
void print(T t)
        if(!t) return;
        print(t->1);
        cout<<t->val<<" ":
        print(t->r);
void reverse(int left,int right)
        Tr *t1, *t2, *t3;
        t1=t2=t3=NULL:
        split(R,t1,t2,left,0);
        split(t2,t2,t3,right-left+2,0);
        t2->rev^=1;
        merge(R,t1,t2);
        merge(R,R,t3);
void get_sum(int left,int right)
        Tr *t1.*t2.*t3:
        t1=t2=t3=NULL:
        split(R,t1,t2,left,0);
        split(t2,t2,t3,right-left+2,0);
        cout << t2->sum << "\n";
        merge(R, t1, t2);
        merge(R,R,t3);
int n,m, q,a,b;
void example()
        ios_base::sync_with_stdio(0);
        cin.tie(0);
        freopen("reverse.in", "r", stdin);
freopen("reverse.out", "w", stdout);
        srand(time(0));
        cin>>n>>m;
        for (int i=1; i <=n; ++i)</pre>
                 T it=new Tr(i,rand()+1,a);
                 insert(R,it,0);
        for (int i=0; i < m; ++i)</pre>
                 cin>>q>>a>>b;
                 if(g) reverse(a.b);
                 else get_sum(a,b);
        return 0:
```

2.7 Trie

```
// Trie <O(|S|), O(|S|)>
int trie[N][26], trien = 1;
int add(int u, char c) {
    c-='a';
    if (trie[u][c]) return trie[u][c];
    return trie[u][c] = ++trien;
}
//to add a string s in the trie
int u = 1;
for(char c : s) u = add(u, c);
```

2.8 Union Find (Disjoint Set Union/Merge Find)

2.9 Union-Find Set (from Lyon)

```
struct Edge // MST
        int a, b, d;
        bool operator < (const Edge &E) const{
        return this->d <E.d;
int ranks[M]; int c[N];
int Find(int x)
        while (y!=c[y])
        while (x!=c[x])
                int aux=c[x];
                c[x]=y;
                x=aux;
        return y;
void Union(int x, int y)
        if(ranks[x]>ranks[y])
                c[x]=y;
        else
        if(ranks[x] == ranks[y])
                ranks[y]++;
```

3 Dynamic Programming

3.1 Divide and Conquer Optimization

```
/**********************************
* DIVIDE AND CONQUER OPTIMIZATION ( dp[i][k] = min j < k \{dp[j][k-1] + C(j,i)\} )
* Description: searches for bounds to optimal point using the monotocity condition*
* Condition: L[i][k] <= L[i+1][k]
* Time Complexity: O(K*N^2) becomes O(K*N*logN)
* Notation: dp[i][k]: optimal solution using k positions, until position i
            L[i][k]: optimal point, smallest j which minimizes dp[i][k]
            C(i,j): cost for splitting range [j,i] to j and i
const int N = 1e3+5;
11 dp[N][N];
//Cost for using i and i
11 C(11 i, 11 j);
void compute(11 1, 11 r, 11 k, 11 opt1, 11 optr){
     // stop condition
    if(1 > r) return;
    11 \text{ mid} = (1+r)/2;
   //best : cost, pos
    pair<11,11> best = {LINF,-1};
    //searchs best: lower bound to right, upper bound to left
   for(11 i = opt1; i <= min(mid, optr); i++){</pre>
       best = min(best, \{dp[i][k-1] + C(i, mid), i\});
    dp[mid][k] = best.first;
    11 opt = best second;
    compute(1, mid-1, k, opt1, opt);
    compute(mid + 1, r, k, opt, optr);
//Iterate over k to calculate
ll solve(){
  //dimensions \ of \ dp[N][K]
  int n, k;
  //Initialize DP
  for(11 i = 1; i <= n; i++) {</pre>
    //dp[i,1] = cost from 0 to i
    dp[i][1] = C(0, i);
  for(11 1 = 2; 1 <= k; 1++) {
   compute(1, n, 1, 1, n);
  /*+ Iterate over i to get min{dp[i][k]}, don't forget cost from n to i
   for (11 i=1; i<=n; i++) {
       11 rest = :
        ans = min(ans, dp[i][k] + rest);
```

3.2 Knuth Optimization

```
// Knuth DP Optimization - O(n^3) -> O(n^2)
//
// 1) dp[i][j] = min i<k<j ( dp[i][k] + dp[k][j] ) + C[i][j]
// 2) dp[i][j] = min k<i ( dp[k][j-1] + C[k][i] )
//
// Condition: A[i][j-1] <= A[i][j] <= A[i+1][j]
// A[i][j] is the smallest k that gives an optimal answer to dp[i][j]
//
// reference (pt-br): https://algorithmmarch.wordpress.com/2016/08/12/a-otimizacao-de-pds-e-o-garcom-da-maratona/
//
// 1) dp[i][j] = min i<k<j ( dp[i][k] + dp[k][j] ) + C[i][j]
int n;
int dp[N][N], a[N][N];
// declare the cost function
int cost(int i, int j) {
    // ...
}

void knuth() {
    // calculate base cases
    memset(dp, 63, sizeof(dp));
    for (int i = 1; i <= n; i++) dp[i][i] = 0;</pre>
```

```
// set initial a[i][j]
  for (int i = 1; i <= n; i++) a[i][i] = i;
  for (int j = 2; j \le n; ++j)
    for (int i = j; i >= 1; --i) {
  for (int k = a[i][j-1]; k <= a[i+1][j]; ++k) {</pre>
        11 v = dp[i][k] + dp[k][j] + cost(i, j);
        // store the minimum answer for d[i][k]
         // in case of maximum, use v > dp[i][k]
        if (v < dp[i][j])
          a[i][j] = k, dp[i][j] = v;
      //+ Iterate over i to get min{dp[i][j]} for each j, don't forget cost from n to
// 2) dp[i][j] = min k < i { <math>dp[k][j-1] + C[k][i] }
int dp[N][J], a[N][J];
// declare the cost function
int cost(int i, int j) {
void knuth() {
 // calculate base cases
  memset(dp, 63, sizeof(dp));
  for (int i = 1; i <= n; i++) dp[i][1] = // ...
  // set initial a[i][j]
  for (int i = 1; i <= n; i++) a[i][1] = 1, a[n+1][i] = n;
  for (int j = 2; j \le maxj; j++)
    for (int i = n; i >= 1; i--){
  for (int k = a[i][j-1]; k <= a[i+1][j]; k++) {</pre>
        11 v = dp[k][j-1] + cost(k, i);
        // store the minimum answer for d[i][k]
        // in case of maximum, use v > dp[i][k]
        if (v < dp[i][j])
          a[i][j] = k, dp[i][j] = v;
      //+ Iterate over i to get min{dp[i][j]} for each j, don't forget cost from n to
```

3.3 Longest Increasing Subsequence

```
// Longest Increasing Subsequence - O(nlogn)
//
// dp(i) = max j<i { dp(j) | a[j] < a[i] } + 1
//
// int dp[N], v[N], n, lis;
memset(dp, 63, sizeof dp);
for (int i = 0; i < n; ++i) {
// increasing: lower_bound
// non-decreasing: upper_bound
int j = lower_bound(dp, dp + lis, v[i]) - dp;
dp[j] = min(dp[j], v[i]);
lis = max(lis, j + 1);
}</pre>
```

3.4 Sum Over Subsets DP

```
// O(N + 2^N)
// A[i] = initial values
// Calculate F[i] = Sum of A[j] for j subset of i
for(int i = 0; i < (1 << N); i++)
        F[i] = A[i];
for(int i = 0; i < N; i++)
        for(int j = 0; j < (1 << N); j++)
        if(j & (1 << i))
             F[j] += F[j ^ (1 << i)];</pre>
```

4 Geometry

4.1 Convex Hull Algorithm

```
bool compare(PT a,PT b) { return a.y<b.y || (a.y==b.y && a.x<b.x); }</pre>
double cross(PT o,PT a, PT b)
        return (a.x-o.x) * (b.y-o.y) - (a.y-o.y) * (b.x-o.x);
vector<PT> ConvexHull(vector<PT> p) {    int n=p.size();    int k=0;
    vector<PT> h(2*n);
    sort(p.begin(),p.end(),compare);
         //build lower hull
        for (int i=0; i < n; ++i)
                 while (k>=2 \&\& cross(h[k-2],h[k-1],p[i]) <=0) k--;
                h[k++]=p[i];
        //build top hull
        for (int i=n-2, t=k+1; i>=0; --i)
                while (k>=t \&\& cross(h[k-2],h[k-1],p[i]) <=0) k--;
                h[k++]=p[i];
        h.resize(k);
        return h;
```

4.2 Delaunay Triangulation

```
Stanford notebook
Delaunay Algorithm Does not handle degenerate cases
Running time: O(n^4)
INPUT: x[] = x-coordinates
y[] = y-coordinates
OUTPUT: triples = a vector containing m triples
(indices corresponding to triangle vertices)
typedef double T;
struct triple {
        int i, j, k;
        triple(int i, int j, int k) : i(i), j(j), k(k) {}
};
vector<triple> delaunayTriangulation(vector<T>& x, vector<T>& y)
        int n = x.size();
        vector<T> z(n);
        vector<triple> ret;
    for (int i = 0; i < n; i++)
         z[i] = x[i] * x[i] + y[i] * y[i];
    for (int i = 0; i < n-2; i++)
        for (int j = i+1; j < n; j++)
                for (int k = i+1; k < n; k++)
                         if ( == k) continue:
                double z_n = (x[j]-x[i])*(y[k]-y[i]) - (x[k]-x[i])*(y[j]-y[i]);
                bool flag = zn < 0;
                for (int m = 0; flag && m < n; m++)</pre>
                         flag = flag && ((x[m]-x[i])*xn + (y[m]-y[i])*yn + (z[m]-z[i])*zn <= 0);
                if (flag) ret.push_back(triple(i, j, k));
        }
    return ret:
int main() {
        T xs[]={0, 0, 1, 0.9};
        T xs[]={0, 1, 0, 0.9};
T ys[]={0, 1, 0, 0.9};
vector<T> x(&xs[0], &xs[4]), y(&ys[0], &ys[4]);
vector<triple> tri = delaunayTriangulation(x, y);
```

4.3 Various Geometry Functions

return false;

```
// Stanford Notebook
double INF = 1e100:
double EPS = 1e-12;
struct PT {
        double x, y;
        PT() {}
        PT(double x, double y) : x(x), y(y) {}
        PT(const PT &p) : x(p.x), y(p.y)
        PT operator + (const PT &p) const { return PT(x+p.x, y+p.y);
        PT operator - (const PT &p) const { return PT(x-p.x, y-p.y); }
        PT operator * (double c)
                                       const { return PT(x*c, y*c );
        PT operator / (double c)
                                       const { return PT(x/c, y/c ); }
double dot (PT p, PT \underline{q}) { return p.x*q.x+p.y*q.y; }
double dist2(PT p, PT q) { return dot(p-q,p-q); }
double cross(PT p, PT q) { return p.x*q.y-p.y*q.x; }
// rotate a point CCW or CW around the origin
PT RotateCCW90(PT p) { return PT(-p.y,p.x); }
PT RotateCW90(PT p) { return PT(p.y,-p.x); }
PT RotateCCW(PT p, double t) { return PT(p.x*cos(t)-p.y*sin(t), p.x*sin(t)+p.y*cos(t)); }
// project point c onto line through a and b // assuming a != b
PT ProjectPointLine(PT a, PT b, PT c) { return a + (b-a)*dot(c-a, b-a)/dot(b-a, b-a); } // project point c onto line segment through a and b
PT ProjectPointSegment(PT a, PT b, PT c)
        double r = dot(b-a,b-a):
        if (fabs(r) < EPS) return a;</pre>
        r = dot(c-a, b-a)/r;
        if (r < 0) return a;</pre>
        if (r > 1) return b;
        return a + (b-a) *r;
// compute distance from c to segment between a and b
double DistancePointSegment(PT a, PT b, PT c)
        return sqrt(dist2(c, ProjectPointSegment(a, b, c)));
// compute distance between point (x,y,z) and plane ax+by+cz=d
double DistancePointPlane(double x, double y, double z, double a, double b, double c, double d)
    return fabs(a*x+b*y+c*z-d)/sqrt(a*a+b*b+c*c);
// determine if lines from a to b and c to d are parallel or collinear
bool LinesParallel(PT a, PT b, PT c, PT d) { return fabs(cross(b-a, c-d)) < EPS; }</pre>
bool LinesCollinear(PT a, PT b, PT c, PT d)
        return LinesParallel(a, b, c, d)
        && fabs(cross(a-b, a-c)) < EPS
        && fabs(cross(c-d, c-a)) < EPS:
// determine if line segment from a to b intersects with
 // line segment from c to d
bool SegmentsIntersect (PT a, PT b, PT c, PT d)
        if (LinesCollinear(a, b, c, d))
                 if (dist2(a, c) < EPS || dist2(a, d) < EPS ||</pre>
                 dist2(b, c) < EPS || dist2(b, d) < EPS) return true;
                 if (dot(c-a, c-b) > 0 && dot(d-a, d-b) > 0 && dot(c-b, d-b) > 0)
                         return false:
                 return true;
        if (cross(d-a, b-a) * cross(c-a, b-a) > 0)
                return false;
        if (cross(a-c, d-c) * cross(b-c, d-c) > 0)
```

```
// compute intersection of line passing through a and b
// with line passing through c and d, assuming that unique
// intersection exists; for segment intersection, check if
// segments intersect first
PT ComputeLineIntersection(PT a, PT b, PT c, PT d)
        b=b-a; d=c-d; c=c-a;
        assert (dot (b, b) > EPS && dot (d, d) > EPS);
        return a + b*cross(c, d)/cross(b, d);
// compute center of circle given three points
PT ComputeCircleCenter(PT a, PT b, PT c)
        b = (a+b)/2;
        c = (a + c) / 2;
        return ComputeLineIntersection(b, b+RotateCW90(a-b), c, c+RotateCW90(a-c));
// determine if point is in a possibly non-convex polygon (by William
// Randolph Franklin); returns 1 for strictly interior points, 0 for
// strictly exterior points, and 0 or 1 for the remaining points.
// Note that it is possible to convert this into an *exact* test using
// integer arithmetic by taking care of the division appropriately
// (making sure to deal with signs properly) and then by writing exact
// tests for checking point on polygon boundary
bool PointInPolygon(const vector<PT> &p, PT q)
        bool c = 0;
        for (int i = 0; i < p.size(); i++)</pre>
                int j = (i+1)%p.size();
                if ((p[i].y <= q.y && q.y < p[j].y ||</pre>
                p[j].y \le q.y && q.y < p[i].y) &&
                q.x < p[i].x + (p[j].x - p[i].x) * (q.y - p[i].y) / (p[j].y - p[i].y))
                c = !c;
        return c:
// determine if point is on the boundary of a polygon
bool PointOnPolygon(const vector<PT> &p, PT q)
        for (int i = 0; i < p.size(); i++)</pre>
                if (dist2(ProjectPointSegment(p[i], p[(i+1)%p.size()], q), q) < EPS)</pre>
                        return true:
        return false:
// compute intersection of line through points a and b with
// circle centered at c with radius r > 0
 vector<PT> CircleLineIntersection (PT a, PT b, PT c, double r)
        vector<PT> ret;
        b = b-a; a = a-c;
        double A = dot(b, b);
        double B = dot(a, b);
        double C = dot(a, a) - r*r;
        double D = B*B - A*C;
        if (D < -EPS) return ret;</pre>
        ret.push_back(c+a+b*(-B+sqrt(D+EPS))/A);
        if (D > EPS)
                ret.push_back(c+a+b*(-B-sqrt(D))/A);
        return ret;
// compute intersection of circle centered at a with radius r
// with circle centered at b with radius R
vector<PT> CircleCircleIntersection(PT a, PT b, double r, double R)
         vector<PT> ret;
        double d = sqrt(dist2(a, b));
        if (d > r+R | | d+min(r, R) < max(r, R)) return ret;</pre>
        double x = (d*d-R*R+r*r)/(2*d);
        double y = sqrt(r*r-x*x);
        PT v = (b-a)/d;
        ret.push_back(a+v*x + RotateCCW90(v)*y);
        if (y > 0) ret.push_back(a+v*x - RotateCCW90(v)*y);
        return ret:
// This code computes the area or centroid of a (possibly nonconvex)
// polygon, assuming that the coordinates are listed in a clockwise or
// counterclockwise fashion. Note that the centroid is often known as
// the "center of gravity" or "center of mass".
double ComputeSignedArea(const vector<PT> &p)
```

return true;

```
double area = 0;
        for(int i = 0; i < p.size(); i++)</pre>
                int j = (i+1) % p.size();
                area += p[i].x*p[j].y - p[j].x*p[i].y;
        return area / 2.0;
double ComputeArea(const vector<PT> &p) {         return fabs(ComputeSignedArea(p)); }
PT ComputeCentroid(const vector<PT> &p)
        PT c(0,0);
        double scale = 6.0 * ComputeSignedArea(p);
        for (int i = 0; i < p.size(); i++)
                int j = (i+1) % p.size();
                c = c + (p[i]+p[j])*(p[i].x*p[j].y - p[j].x*p[i].y);
        return c / scale;
// tests whether or not a given polygon (in CW or CCW order) is simple
bool IsSimple(const vector<PT> &p)
        for (int i = 0; i < p.size(); i++)</pre>
                for (int k = i+1; k < p.size(); k++)</pre>
                         int j = (i+1) % p.size();
                         int 1 = (k+1) % p.size();
                         if (i == 1 \mid \mid j == k) continue;
                         if (SegmentsIntersect(p[i], p[j], p[k], p[l]))
                                 return false;
        return true;
```

5 Graphs

5.1 BFS

```
* BFS (BREADTH-FIRST SEARCH)
* Time complexity: O(V+E)
* Usage: bfs(node)
* Notation: s: starting node
        adj[i]: adjacency list for node i
         vis[i]: visited state for node i (0 or 1)
const int N = 1e5+10; // Maximum number of nodes
int dist[N], par[N];
vector <int> adj[N];
queue <int> q;
void bfs (int s) {
   memset(dist, 63, sizeof(dist));
   dist[s] = 0;
   q.push(s);
   while (!q.empty()) {
      int u = q.front(); q.pop();
      for (auto v : adj[u]) if (dist[v] > dist[u] + 1) {
    par[v] = u;
         dist[v] = dist[u] + 1;
         q.push(v);
```

5.2 DFS

```
vis[i]: visited state for node i (0 or 1)

const int N = le5+l0;
int vis[N];
vector<int> adj[N];

void dfs(int u) {
    vis[u] = 1;
    for (int v : adj[u]) {
        if (!vis[v]) {
            dfs(v);
        }
        // vis[u] = 0;
        // Uncomment the line above if you need to
        // traverse only one path at a time (backtracking)
}
```

5.3 Dijkstra's Algorithm

```
//Dijkstra Algorithm
int t,n,m,s,e;
vector<ii> edges[N];//pair<NodeEnd,dist>
int distances[N]; // =INF=0x3f3f3f3f
int parent[N]; // =-1
int Dijkstra()
         vector<ii>> :: iterator it;
        priority_queue< ii, vector<ii>, greater<ii> > pq;
        distances[s]=0;
        pq.push(ii(distances[s],s));
         while(!pq.empty())
                ii p = pq.top();
                 pq.pop();
                int d=p.first;
                int a=p.second;
                for(it=edges[a].begin();it!=edges[a].end();++it)
                         if(distances[it->first]>distances[a]+it->second)
                             distances[it->first]=distances[a]+it->second;
                             parent[it->first]=a;
                             pq.push(ii(distances[it->first],it->first));
        return distances[e];
```

5.4 Shortest Path (Dijkstra)

```
* DIJKSTRA'S ALGORITHM (SHORTEST PATH TO A VERTEX)
* Time complexity: O((V+E)logE)
* Usage: dist[node]
* Notation: m:
            (a, b, w): edge between a and b with weight w
                          starting node
            par[v]:
                         parent node of u, used to rebuild the shortest path
vector<int> adj[N], adjw[N];
int dist[N]:
memset(dist, 63, sizeof(dist));
priority_queue<pii> pq;
pq.push (mp(0,0));
while (!pq.empty()) {
  int u = pq.top().nd;
  int d = -pq.top().st;
  pq.pop();
  if (d > dist[u]) continue;
  for (int i = 0; i < adj[u].size(); ++i) {</pre>
   int v = adj[u][i];
int w = adjw[u][i];
    if (dist[u] + w < dist[v])</pre>
     dist[v] = dist[u]+w, pq.push(mp(-dist[v], v));
```

5.5 Shortest Path (Bellman-Ford)

```
* BELLMAN-FORD ALGORITHM (SHORTEST PATH TO A VERTEX - WITH NEGATIVE COST)
* Time complexity: O(VE)
* Usage: dist[node]
* Notation: m:
                       number of edges
          n:
                       number of vertices
          (a, b, w): edge between a and b with weight w
                      starting node
          s:
const int N = 1e4+10; // Maximum number of nodes
vector<int> adj[N], adjw[N];
int dist[N], v, w;
memset(dist, 63, sizeof(dist));
dist[0] = 0;
for (int i = 0; i < n-1; ++i)
 for (int u = 0; u < n; ++u)

for (int j = 0; j < adj[u].size(); ++j)

v = adj[u][j], w = adjw[u][j],
     dist[v] = min(dist[v], dist[u]+w);
```

5.6 Topological Sort (Kahn)

```
* KAHN'S ALGORITHM (TOPOLOGICAL SORTING)
* Time complexity: O(V+E)
* Notation: adj[i]: adjacency matrix for node i
                   number of vertices
          e:
                   number of edges
          a, b: edge between a and b
          inc:
                  number of incoming arcs/edges
                   queue with the independent vertices
          tsort: final topo sort, i.e. possible order to traverse graph
*******************************
vector <int> adj[N];
int inc[N]; // number of incoming arcs/edges
// undirected graph: inc[v] <= 1
// directed graph: inc[v] == 0
queue<int> q;
for (int i = 1; i <= n; ++i) if (inc[i] <= 1) q.push(i);
while (!g.emptv()) {
 int u = q.front(); q.pop();
  for (int v : adj[u])
   if (inc[v] > 1 and --inc[v] <= 1)</pre>
     q.push(v);
```

5.7 Minimum Spanning Tree (Kruskal)

5.8 Minimum Spanning Tree (Prim)

```
// Prim - MST O(ElogE)
vi adj[N], adjw[N];
int vis[N];

priority_queue<pii>> pq;
pq.push(mp(0, 0));

while (!pq.empty()) {
   int u = pq.top().nd;
   pq.pop();
   if (vis[u]) continue;
   vis[u]=1;
   for (int i = 0; i < adj[u].size(); ++i) {
     int v = adj[u][i];
     int w = adjw[u][i];
     if (!vis[v]) pq.push(mp(-w, v));
   }
}</pre>
```

5.9 Max Flow (Dinic's Algorithm)

```
// Stanford Notebook
typedef long long LL;
struct Edge
        int from, to, cap, flow, index;
        Edge(int from, int to, int cap, int flow, int index) :
        from(from), to(to), cap(cap), flow(flow), index(index) {}
        LL rcap() { return cap - flow; }
};
struct Dinic
        int N:
        vector<vector<Edge> > G;
        vector<vector<Edge *> > Lf;
        vector<int> layer;
        vector<int> Q;
        Dinic(int N) : N(N), G(N), Q(N) {}
        void AddEdge(int from, int to, int cap)
                if (from == to) return;
                G[from].push\_back(Edge(from, to, cap, 0, G[to].size()));
                G[to].push_back(Edge(to, from, 0, 0, G[from].size() - 1));
        LL BlockingFlow(int s, int t)
                laver.clear():
                layer.resize(N, -1);
                laver[s] = 0;
                Lf.clear(); Lf.resize(N);
        int head = 0, tail = 0;
               Q[tail++] = s;
        while (head < tail)</pre>
            int x = Q[head++];
                        for (int i = 0; i < G[x].size(); i++)
                                Edge &e = G[x][i]; if (e.rcap() <= 0) continue;</pre>
                                if (layer[e.to] == -1)
                                         layer[e.to] = layer[e.from] + 1;
                                        Q[tail++] = e.to;
                                if (layer[e.to] > layer[e.from])
                                        Lf[e.from].push_back(&e);
        if (layer[t] == -1) return 0;
        LL totflow = 0;
               vector<Edge *> P;
                while (!Lf[s].empty())
                        int curr = P.empty() ? s : P.back()->to;
                        if (curr == t)
                                // Augment
```

```
LL amt = P.front()->rcap();
                                   for (int i = 0; i < P.size(); ++i)</pre>
                                            amt = min(amt, P[i]->rcap());
                                   totflow += amt;
                                   for (int i = P.size() - 1; i >= 0; --i)
                                            G[P[i]\rightarrow to][P[i]\rightarrow index].flow -= amt;
                                            if (P[i]->rcap() <= 0)</pre>
                                                     Lf[P[i]->from].pop_back();
                                                     P.resize(i);
                           else if (Lf[curr].empty())
                                    // Retreat
                                   P.pop_back();
                                   for (int i = 0; i < N; ++i)
                                            for (int j = 0; j < Lf[i].size(); ++j)
    if (Lf[i][j]->to == curr)
                                                              Lf[i].erase(Lf[i].begin() + j);
                           else
                                    // Advance
                                   P.push_back(Lf[curr].back());
                  return totflow;
        LL GetMaxFlow(int s, int t)
                  LL totflow = 0;
                  while (LL flow = BlockingFlow(s, t))
                          totflow += flow;
                  return totflow;
};
```

5.10 Max Flow (Edmonds-Karp Algorithm)

```
Stanford Notebook
{\it MinCostMaxFlow} (adjacency matrix, Edmonds and Karp 1972)
This implementation keeps track of forward and reverse edges separately (so you can set cap[i][j] != cap[j][i]). For a regular max flow, set all edge costs to 0.
Running time, O(|V|^2) cost per augmentation max flow: O(|V|^3) augmentations min cost max flow: O(|V|^4 * MAX\_EDGE\_COST) augmentations
INPUT: - graph, constructed using AddEdge()
             - source
             - sink
OUTPUT: - (maximum flow value, minimum cost value)
                  - To obtain the actual flow, look at positive values only.
typedef vector<int> VI;
typedef vector<VI> VVI;
typedef long long L;
typedef vector<L> VL;
typedef vector<VL> VVL;
typedef pair<int, int> PII;
typedef vector<PII> VPII;
const L INF = numeric_limits<L>::max() / 4;
struct MinCostMaxFlow {
         int N;
         VVL cap, flow, cost;
         VI found;
         VL dist, pi, width;
         VPII dad;
     MinCostMaxFlow(int N) :
     N(N), cap(N, VL(N)), flow(N, VL(N)), cost(N, VL(N)),
     found(N), dist(N), pi(N), width(N), dad(N) {}
     void AddEdge(int from, int to, L cap, L cost)
         this->cap[from][to] = cap;
         this->cost[from][to] = cost;
```

```
void Relax(int s, int k, L cap, L cost, int dir)
    L \text{ val} = \text{dist}[s] + \text{pi}[s] - \text{pi}[k] + \text{cost};
    if (cap && val < dist[k])</pre>
             dist[k] = val;
             dad[k] = make_pair(s, dir);
             width[k] = min(cap, width[s]);
L Dijkstra(int s, int t)
    fill(found.begin(), found.end(), false);
fill(dist.begin(), dist.end(), INF);
    fill(width.begin(), width.end(), 0);
    dist[s] = 0;
    width[s] = INF;
    while (s != -1)
             int best = -1:
             found[s] = true;
             for (int k = 0; k < N; k++)
                      if (found[k]) continue;
                      Relax(s, k, cap[s][k] - flow[s][k], cost[s][k], 1);
                      Relax(s, k, flow[k][s], -cost[k][s], -1);
                      if (best == -1 || dist[k] < dist[best]) best = k;</pre>
         s = best:
    for (int k = 0; k < N; k++)
            pi[k] = min(pi[k] + dist[k], INF);
    return width[t];
pair<L, L> GetMaxFlow(int s, int t)
    L totflow = 0, totcost = 0;
while (L amt = Dijkstra(s, t))
             totflow += amt;
             for (int x = t; x != s; x = dad[x].first)
                      if (dad[x].second == 1)
                               flow[dad[x].first][x] += amt;
                               totcost += amt * cost[dad[x].first][x];
                      else
                               flow[x][dad[x].first] -= amt;
                               totcost -= amt * cost[x][dad[x].first];
    return make_pair(totflow, totcost);
```

5.11 Min Cost Max Flow

};

```
// HSE TNF = 1691
/******************************
* MIN COST MAX FLOW (MINIMUM COST TO ACHIEVE MAXIMUM FLOW)
* Description: Given a graph which represents a flow network where every edge has *
* a capacity and a cost per unit, find the minimum cost to establish the maximum *
* possible flow from s to t.
* Note: When adding edge (a, b), it is a directed edge!
* Usage: min_cost_max_flow()
       add_edge(from, to, cost, capacity)
* Notation: flw: max flow
         cst: min cost to achieve flw
* Testcase:
* add_edge(src, 1, 0, 1); add_edge(1, snk, 0, 1); add_edge(2, 3, 1, INF);
* add_edge(src, 2, 0, 1); add_edge(2, snk, 0, 1); add_edge(3, 4, 1, INF);
* add_edge(src, 2, 0, 1); add_edge(3, snk, 0, 1);
* add_edge(src, 2, 0, 1); add_edge(4, snk, 0, 1); => flw = 4, cst = 3
// w: weight or cost, c : capacity
struct edge {int v, f, w, c; };
```

```
int n, flw_lmt=INF, src, snk, flw, cst, p[N], d[N], et[N];
vector<edge> e;
vector<int> g[N];
void add_edge(int u, int v, int w, int c) {
  int k = e.size();
  g[u].push_back(k);
  g[v].push_back(k+1);
  e.push_back({ v, 0, w, c });
 e.push_back({ u, 0, -w, 0 });
void clear() {
  flw lmt = INF;
  for(int i=0; i<=n; ++i) g[i].clear();</pre>
  e.clear();
void min_cost_max_flow() {
  flw = 0, cst = 0;
  while (flw < flw_lmt) {</pre>
   memset(et, 0, (n+1) * sizeof(int));
    memset(d, 63, (n+1) * sizeof(int));
    deque<int> q:
    q.push_back(src), d[src] = 0;
    while (!q.empty()) {
     int u = q.front(); q.pop_front();
      et[u] = 2;
      for(int i : g[u]) {
        edge &dir = e[i];
        int v = dir.v;
        if (dir.f < dir.c and d[u] + dir.w < d[v]) {
          d[v] = d[u] + dir.w;
         if (et[v] == 0) q.push_back(v);
          else if (et[v] == 2) q.push_front(v);
         et[v] = 1;
         p[v] = i;
   if (d[snk] > INF) break;
    int inc = flw_lmt - flw;
    for (int u=snk; u != src; u = e[p[u]^1].v) {
      edge &dir = e[p[u]];
      inc = min(inc, dir.c - dir.f);
    for (int u=snk; u != src; u = e[p[u]^1].v) {
     edge &dir = e[p[u]], &rev = e[p[u]^1];
      dir.f += inc:
     rev.f -= inc:
     cst += inc * dir.w:
    if (!inc) break;
    flw += inc;
```

5.12 Eulerian Path

```
struct Edge:
typedef list<Edge>::iterator iter;
struct Edge
       int next vertex:
       iter reverse edge:
    Edge(int next_vertex) :next_vertex(next_vertex) { }
};
const int max_vertices = ;
int num_vertices;
list<Edge> adj[max_vertices]; // adjacency list
vector<int> path;
void find_path(int v)
       while(adj[v].size() > 0)
               int vn = adj[v].front().next vertex;
       adj[vn].erase(adj[v].front().reverse edge);
       adj[v].pop_front();
       find_path(vn);
```

```
}
path.push_back(v);
}

void add_edge(int a, int b)
{
    adj[a].push_front(Edge(b));
    iter ita = adj[a].begin();
    adj[b].push_front(Edge(a));
    iter itb = adj[b].begin();
    ita->reverse_edge = itb;
    itb->reverse_edge = ita;
}
```

5.13 Hopcroft-Karp Algorithm

```
#include <vector>
vector<int> g[N];
int r[N], l[N], n, m, e, a, b;
bool dfs(int v)
        if(vis[v]) return false;
        vis[v] = true;
        for(int u=0; u<g[v].size(); ++u)</pre>
                 if(!r[g[v][u]])
                          1[v]=g[v][u];
r[g[v][u]]=v;
                          return true:
        for (int u=0; u<q[v].size(); ++u)</pre>
                 if(dfs(r[g[v][u]]))
                          1[v]=g[v][u];
                          r[g[v][u]]=v;
                          return true;
        return false:
void hoperoft karp()
        bool change = true;
        while (change)
                 fill(vis, vis+n+1, false);
                 for(int i=1; i<=n; ++i)</pre>
                         if(!l[i])
                                   change |= dfs(i);
```

5.14 2-SAT Kosaraju

```
/***********************************
* 2-SAT (TELL WHETHER A SERIES OF STATEMENTS CAN OR CANNOT BE FEASIBLE AT THE SAME TIME)
* Time complexity: O(V+E)
* Usage: n
               -> number of variables, 1-indexed
       p = v(i) -> picks the "true" state for variable i
       p = nv(i) -> picks the "false" state for variable i, i.e. ~i
       add(p, q) \rightarrow add clause (p \ v \ q) (which also means p \rightarrow q, which also means q \rightarrow p)
       run2sat() -> true if possible, false if impossible
               -> tells if i has to be true or false for that solution
int n, vis[2*N], ord[2*N], ordn, cnt, cmp[2*N], val[N];
vector<int> adj[2*N], adjt[2*N];
// for a variable u with idx i
// u is 2*i and !u is 2*i+1
// (a v b) == !a -> b ^ !b -> a
int v(int x) { return 2*x; }
```

```
int nv(int x) { return 2*x+1; }
// add clause (a v b)
void add(int a, int b) {
  adj[a^1].push_back(b);
  adj[b^1].push_back(a);
  adjt[b].push_back(a^1);
  adjt[a].push_back(b^1);
void dfs(int x) {
  vis[x] = 1;
 for(auto v : adj[x]) if(!vis[v]) dfs(v);
ord[ordn++] = x;
void dfst(int x) {
  cmp[x] = cnt, vis[x] = 0;
  for(auto v : adjt[x]) if(vis[v]) dfst(v);
bool run2sat(){
  for(int i = 1; i <= n; i++) {</pre>
    if(!vis[v(i)]) dfs(v(i));
    if(!vis[nv(i)]) dfs(nv(i));
  for(int i = ordn-1; i >= 0; i--)
   if(vis[ord[i]]) cnt++, dfst(ord[i]);
  for(int i = 1; i <= n; i ++) {
   if(cmp[v(i)] == cmp[nv(i)]) return false;</pre>
    val[i] = cmp[v(i)] > cmp[nv(i)];
  return true;
int main () {
    for (int i = 1; i <= n; i++) {
        if (val[i]); // i-th variable is true
                       // i-th variable is false
```

5.15 Articulation points and bridges

```
// Articulation points and Bridges O(V+E)
// An articulation (cut vertex), if removed, disconnects the graph
int par[N], art[N], low[N], num[N], ch[N], cnt;

void articulation(int u) {
    low[u] = num[u] = ++cnt;
    for (int v : adj[u]) {
        if (!num[v]) {
            par[v] = u; ch[u]++;
            articulation(v);
        if (low[v] >= num[u]) art[u] = 1;
            if (low[v] >= num[u]) { /* u-v bridge */ }
        low[u] = min(low[u], low[v]);
        }
    else if (v != par[u]) low[u] = min(low[u], num[v]);
    }
}
for (int i = 0; i < n; ++i) if (!num[i])
    articulation(i), art[i] = ch[i]>1;
```

5.16 Hungarian

```
// Hungarian - O(m*n^2)
// Assignment Problem

int n, m;
int pu[N], pv[N], cost[N] [M];
int pairV[N], way[M], minv[M], used[M];

void hungarian() {
  for(int i = 1, j0 = 0; i <= n; i++) {
    pairV[0] = i;
    memset(minv, 63, sizeof minv);
    memset(used, 0, sizeof used);
    do {
        used[j0] = 1;
        int i0 = pairV[j0], delta = INF, j1;
        for(int j = 1; j <= m; j++) {
            if (used[j1)) continue;
        }
}</pre>
```

```
int cur = cost[i0][j] - pu[i0] - pv[j];
    if(cur < minv[j]) minv[j] = cur, way[j] = j0;
    if(minv[j] < delta) delta = minv[j], j1 = j;
}

for(int j = 0; j <= m; j++) {
    if(used[j]) pu[pairv[j]] += delta, pv[j] -= delta;
    else minv[j] -= delta;
    }
    j0 = j1;
    } while(pairv[j0]);

do {
    int j1 = way[j0];
    pairv[j0] = pairv[j1];
    j0 = j1;
    } while(j0);
}
}
// in main
// for(int j = 1; j <= m; j++)
// if(pairv[j]) ans += cost[pairv[j]][j];</pre>
```

5.17 Hungarian Navarro

```
// Hungarian - O(n^2 * m)
template<br/>bool is max = false, class T = int, bool is zero indexed = false>
struct Hungarian {
  bool swap_coord = false;
  int lines, cols;
  vector<int> pairV, way;
  vector<bool> used:
  vector<T> pu, pv, minv;
  vector<vector<T>> cost;
  Hungarian(int _n, int _m)
    if (_n > _m) {
      swap(n. m):
      swap_coord = true;
    lines = _n + 1, cols = _m + 1;
    cost.resize(lines);
    for (auto& line : cost) line.assign(cols, 0);
  void clear() {
    pairV.assign(cols, 0);
    way.assign(cols, 0);
    pv.assign(cols, 0);
    pu.assign(lines, 0);
  void update(int i, int j, T val) {
    if (is_zero_indexed) i++, j++;
    if (is_max) val = -val;
    if (swap_coord) swap(i, j);
    assert(i < lines);</pre>
    assert(j < cols);
    cost[i][j] = val;
  T run() {
    T _INF = numeric_limits<T>::max();
    for (int i = 1, j0 = 0; i < lines; i++) {
      pairV[0] = i;
       minv.assign(cols, _INF);
       used.assign(cols, 0);
         used[j0] = 1;
         int i0 = pairV[j0], j1;
         T delta = _INF;
for (int j = 1; j < cols; j++) {
          if (used[j]) continue;
T cur = cost[i0][j] - pu[i0] - pv[j];
if (cur < minv[j]) minv[j] = cur, way[j] = j0;
if (minv[j] < delta) delta = minv[j], j1 = j;</pre>
```

```
for (int j = 0; j < cols; j++) {
                              if (used[j]) pu[pairV[j]] += delta, pv[j] -= delta;
                                else minv[j] -= delta;
                            j0 = j1;
                    } while (pairV[j0]);
                   do {
                         int j1 = way[j0];
pairV[j0] = pairV[j1];
                   } while (j0);
             for (int j = 1; j < cols; j++) if (pairV[j]) ans += cost[pairV[j]][j];</pre>
              if (is_max) ans = -ans;
             if (is_zero_indexed) {
                   for (int j = 0; j + 1 < cols; j++) pairV[j] = pairV[j + 1], pairV[j]--;</pre>
                   pairV[cols - 1] = -1;
             if (swap_coord) {
                   vector<int> pairV_sub(lines, 0);
                   swap(pairV, pairV_sub);
             return ans:
1:
template <bool is_max = false, bool is_zero_indexed = false>
struct HungarianMult : public Hungarian<is_max, long double, is_zero_indexed> {
      using super = Hungarian<is_max, long double, is_zero_indexed>;
      HungarianMult(int _n, int _m) : super(_n, _m) {}
       \begin{tabular}{ll} \beg
             super::update(i, j, log2(x));
};
```

5.18 Max Bipartite Cardinality Matching (Kuhn)

```
* KUHN'S ALGORITHM (FIND GREATEST NUMBER OF MATCHINGS - BIPARTITE GRAPH)
* Time complexity: O(VE)
* Notation: ans:
                 number of matchings
         b[j]:
                 matching edge b[j] <-> j
         adj[i]: adjacency list for node i
         vis:
                 visited nodes
                 counter to help reuse vis list
*************************
// TIP: If too slow, shuffle nodes and try again.
int x, vis[N], b[N], ans;
bool match (int u) {
 if (vis[u] == x) return 0;
 vis[u] = x;
 for (int v : adj[u])
   if (!b[v] or match(b[v])) return b[v]=u;
 return 0;
for (int i = 1; i <= n; ++i) ++x, ans += match(i);</pre>
// Maximum Independent Set on bipartite graph
MIS + MCBM = V
// Minimum Vertex Cover on bipartite graph
MVC = MCBM
```

5.19 Lowest Common Ancestor

```
// floor of the binary logarithm of n
int 1b (unsigned int n)
        if(n==0) return -1;
        if (n >= 1<<16) { n >>= 16; p += 16; }
        if (n >= 1<< 8) { n >>= 8; p += 8;
        if (n >= 1<< 4) { n >>= 4; p += 4;
        if (n >= 1<< 2) { n >>= 2; p += 2;
        if (n >= 1<< 1) {
        return p;
void DFS(int i, int 1)
        L[i] = 1;
        for(int j = 0; j < children[i].size(); j++)</pre>
                 DFS(children[i][j], 1+1);
int LCA(int p, int q) {
             nsure node p is at least as deep as node q
        if(L[p] < L[q]) swap(p, q);
         // "binary search" for the ancestor of node p situated on the same level as q
        for(int i = log_num_nodes; i >= 0; i--)
                if(L[p] - (1<<i) >= L[q])
         p = A[p][i];
        if(p == q) return p;
          // "binary search" for the LCA
        for(int i = log_num_nodes; i >= 0; i--)
                 if(A[p][i] != -1 && A[p][i] != A[q][i])
                         p = A[p][i];
                         q = A[q][i];
    return A[p][0];
int main(int argc,char* argv[])
         // read num nodes, the total number of nodes
        log_num_nodes=lb(num_nodes);
        for(int i = 0; i < num_nodes; i++)</pre>
                 // read p, the parent of node i or -1 if node i is the root
        A[i][0] = p;
        if(p != -1) children[p].push_back(i);
        else root = i;
    // precompute A using dynamic programming
    for(int j = 1; j <= log_num_nodes; j++)
    for(int i = 0; i < num_nodes; i++)
        if(A[i][j-1] != -1)        A[i][j] = A[A[i][j-1]][j-1];</pre>
                else A[i][j] = -1;
    // precompute L
        DFS (root, 0);
        return 0:
```

5.20 Strongly Connected Components

```
void fill_backward(int x)
        v[x]=false;
        group_num[x]=group_cnt;
        for(i=spr[x];i;i=er[i].nxt)
                if(v[er[i].e])
                        fill backward(er[i].e);
void add_edge(int v1, int v2) //add edge v1->v2
        e[++E].e=v2:
        e[E].nxt=sp [v1];
        sp[v1]=E;
        er[E].e=v1;
        er[E].nxt=spr[v2];
        spr[v2]=E;
void SCC()
        int i:
        stk[0]=0;
        memset(v, false, sizeof(v));
        for(i=1;i<=V;i++)
               if(!v[i])
                        fill_forward(i);
        group_cnt=0;
        for(i=stk[0];i>=1;i--)
                if(v[stk[i]]) {group_cnt++; fill_backward(stk[i]);}
```

5.21 Strongly Connected Components (Kosaraju)

```
* KOSARAJU'S ALGORITHM (GET EVERY STRONGLY CONNECTED COMPONENTS (SCC))
* Description: Given a directed graph, the algorithm generates a list of every
* strongly connected components. A SCC is a set of points in which you can reach
* every point regardless of where you start from. For instance, cycles can be
* a SCC themselves or part of a greater SCC.
* This algorithm starts with a DFS and generates an array called "ord" which
* stores vertices according to the finish times (i.e. when it reaches "return").
* Then, it makes a reversed DFS according to "ord" list. The set of points
* visited by the reversed DFS defines a new SCC.
\star One of the uses of getting all SCC is that you can generate a new DAG (Directed \star
* Acyclic Graph), easier to work with, in which each SCC being a "supernode" of
* the DAG.
* Time complexity: O(V+E)
* Notation: adj[i]: adjacency list for node i
           adjt[i]: reversed adjacency list for node i
                     array of vertices according to their finish time
            ord:
            ordn:
                     ord counter
            scc[i]: supernode assigned to i
           scc cnt: amount of supernodes in the graph
const int N = 2e5 + 5;
vector<int> adj[N], adjt[N];
int n, ordn, scc_cnt, vis[N], ord[N], scc[N];
//Directed Version
void dfs(int u) {
  vis[u] = 1;
  for (auto v : adj[u]) if (!vis[v]) dfs(v);
 ord[ordn++] = u;
void dfst(int u) {
 scc[u] = scc\_cnt, vis[u] = 0;
  for (auto v : adjt[u]) if (vis[v]) dfst(v);
// add edge: u -> v
void add_edge(int u, int v) {
  adj[u].push_back(v);
  adjt[v].push_back(u);
//Undirected version:
 int par[N];
  void dfs(int u) {
   vis[u] = 1;
```

```
for (auto v : adj[u]) if(!vis[v]) par[v] = u, dfs(v);
    ord[ordn++] = u;
}

void dfst(int u) {
    scc[u] = scc_cnt, vis[u] = 0;
    for (auto v : adj[u]) if(vis[v] and u != par[v]) dfst(v);
}

// add edge: u -> v
    void add_edge(int u, int v) {
        adj[u].push_back(v);
        adj[v].push_back(u);
}

*/

// run kosaraju
void kosaraju() {
    for (int i = 1; i <= n; ++i) if (!vis[i]) dfs(i);
    for (int i = ordn - 1; i >= 0; --i) if (vis[ord[i]]) scc_cnt++, dfst(ord[i]);
}
```

5.22 Tarjan

```
// Tarjan for SCC and Edge Biconnected Componentes - O(n + m)
vector<int> adj[N];
stack<int> st:
bool inSt[N];
int id[N], cmp[N];
int cnt, cmpCnt;
void clear(){
 memset(id, 0, sizeof id);
  cnt = cmpCnt = 0;
int tarjan(int n) {
  id[n] = low = ++cnt;
  st.push(n), inSt[n] = true;
  \quad \textbf{for} \, (\textbf{auto} \ x \ : \ \texttt{adj} \, [n] \,) \, \{
    if(id[x] and inSt[x]) low = min(low, id[x]);
    else if(!id[x]) {
      int lowx = tarjan(x);
      if(inSt[x])
         low = min(low, lowx);
  if(low == id[n]){
    while(st.size()){
      int x = st.top();
inSt[x] = false;
      cmp[x] = cmpCnt;
       st.pop();
      if(x == n) break;
    cmpCnt++:
  return low;
```

5.23 Zero One BFS

```
// 0-1 BFS - O(V+E)
const int N = 1e5 + 5;
int dist[N];
vector<pii> adj[N];
deque<pii> dq;

void zero_one_bfs (int x){
    cl(dist, 63);
    dist[x] = 0;
    dc,push_back({x, 0});
    while(!dq.empty()){
        int u = dq.front().st;
        int ud = dq.front().nd;
```

```
dq.pop_front();
    if(dist[u] < ud) continue;
    for(auto x : adj[u]) {
        int v = x.st;
        int w = x.nd;
        if(dist[u] + w < dist[v]) {
            dist[v] = dist[u] + w;
            if(w) dq.push_back({v, dist[v]});
            else dq.push_front({v, dist[v]});
        }
    }
}</pre>
```

6 Math with Numbers

6.1 Extended Euclid's Algorithm

```
#include "GcdLcm.h"

// returns d = gcd(a,b); find x, y such that d = ax + by
int extended_euclid(int a, int b, int &x, int &y) {
    int current_x = y = 0;
    int current_y = x = 1;
    while(b)
    {
        int q = a/b;
        int t = b;
        b = a&b;
        a = t;
        t = current_x; current_x = x-q*current_x; x = t;
        t = current_y; current_y = y-q*current_y; y = t;
    }
    return a;
}
```

6.2 Fast Prime Number Sieve

6.3 GCD and LCM

```
// return a%b
int mod(int a, int b)
{
     return ((a%b)+b)%b;
}
// computes gcd(a,b)
```

```
int gcd(int a, int b)
{
        int tmp;
        while(b)
        {
            a %= b;
            tmp = a;
            a = b;
            b = tmp;
        }
        return a;
}
// computes lcm(a,b)
int lcm(int a, int b)
{
        return a/gcd(a,b)*b;
}
```

6.4 Euler Phi/Totient

```
// Euler phi (totient)
int ind = 0, pf = primes[0], ans = n;
while (ll1*pf*pf <= n) {
    if (n%pf==0) ans -= ans/pf;
    while (n%pf==0) n /= pf;
    pf = primes[++ind];
}
if (n != 1) ans -= ans/n;
// IME2014
int phi[N];
void totient() {
    for (int i = 1; i < N; ++i) phi[i]=i;
    for (int i = 2; i < N; i+=2) phi[i]>>=1;
    for (int i = 2; i < N; i+=2) phi[j]==j) {
        phi[j]--;
        for (int i = 2*j; i < N; i+=j) phi[i]=phi[i]/j*(j-1);
    }
}</pre>
```

7 Math with Matrices?

7.1 Modular Linear Equation Solver

```
// Stanford Notebook
#include <vector>
#include "ExtendedEuclid.h"
// find all solutions to ax = b \pmod{n}
vector<int> modular_linear_equation_solver(int a, int b, int n)
        int x, v;
        vector<int> sol;
        int d = extended_euclid(a, n, x, y);
        if(!(b%d)) {
                x = mod(x%(b/d), n);
                for (int i=0; i < d; ++i)
                        sol.push\_back(mod(x + i*(n/d),n));
        return sol:
// computes b such tath ab = 1 \pmod{n}, returs -1 on failure
int mod_inverse(int a, int n)
        int x, y;
        int d = extended_euclid(a, n, x, y);
        if(d>1) return -1;
        return mod(x,n);
// computes x and y such that ax + by =c; on failure, x = y = -1
void linear_diophantine(int a, int b, int c, int &x, int &y)
        int d = gcd(a,b);
        if (c%d) \bar{x} = y = -1;
        else
                x = c/d * mod_inverse(a/d,b/d);
                y = (c-a*x)/b;
```

7.2 Fast Fourier Transform

```
struct cpx
         cpx(double aa): a(aa){}
         cpx(double aa, double bb): a(aa), b(bb){}
        double a:
        double b;
        double modsg(void) const{ return a*a+b*b; }
        cpx bar(void) const {return cpx(a,-b);}
1:
cpx b[N+100],c[N+100],B[N+100],C[N+100];
int a[N+100], int x[N+100];
double coss[N+100], sins[N+100];
int n,m,p;
cpx operator + (cpx a, cpx b) {
                                    return cpx(a.a+b.a,a.b+b.b); }
cpx operator *(cpx a,cpx b) {
                                    return cpx(a.a*b.a-a.b*b.b,a.a*b.b+a.b*b.a); }
cpx operator / (cpx a,cpx b) {       cpx r = a*b.bar();       return cpx(r.a/b.modsq(),r.b/b.modsq()); }
cpx EXP(int i, int dir) { return cpx(coss[i], sins[i]*dir); }
const double two_pi = 4 * acos(0);
void FFT(cpx *in,cpx *out,int step,int size,int dir)
        if(size<1) return;</pre>
        if(size==1)
                  out[0]=in[0];
                  return:
        FFT(in,out,step*2,size/2,dir);
        FFT (in+step, out+size/2, step*2, size/2, dir);
        for(int i=0; i < size/2; ++i)</pre>
                  cpx even=out[i];
                 cpx odd=out[i+size/2];
                  out[i] = even+EXP(i*step,dir)*odd;
                  out[i+size/2]=even+EXP((i+size/2)*step,dir)*odd;
void exemple
        for(int i=0;i<=N;++i)</pre>
                  coss[i]=cos(two_pi*i/N);
                  sins[i]=sin(two_pi*i/N);
        while (scanf ("%d", &n) ==1)
                  fill(x,x+N+100,0);
                  fill(a,a+N+100,0);
                  for (int i=0; i< n; ++i)
                           scanf("%d",&p);
                          x[p]=1;
                  for(int i=0;i<N+100;++i)
                          b[i]=cpx(x[i],0);
                  scanf("%d", &m);
                  for(int i=0;i<m;++i)</pre>
                          scanf("%d",&a[i]);
                  FFT(b,B,1,N,1);
                  \quad \textbf{for}\,(\,\textbf{int}\  \  \, \textbf{i}\!=\!0\,;\,\textbf{i}\!<\!N\,;\,+\!+\!\textbf{i}\,)
                          C[i]=B[i]*B[i];
                 FFT(C,c,1,N,-1);
                  for (int i=0; i<N; ++i)
                         c[i]=c[i]/N;
                 int cnt=0;
for(int i=0;i<16;++i)</pre>
                         cout<<c[il.a<<" ":
                  for(int i=0;i<m;++i)</pre>
                         if(c[a[i]].a>0.5 || x[a[i]])
                                   cnt++;
                  printf("%d\n",cnt);
```

7.3 Gauss-Jordan Elimination

```
Stanford notebook
GaussJordan Algorithm (elimination with full pivoting)
(1) solving systems of linear equations (AX=B)
(2) inverting matrices (AX=I)
(3) computing determinants of square matrices
Running time: O(n^3)
INPUT: a[][] = an nxn matrix
             b[][] = an nxm matrix
               = an nxm matrix (stored in b[][])
                 A^{-1} = an nxn matrix (stored in a[][])
                 returns determinant of a[][]
const double EPS = 1e-10;
typedef vector<int> VI;
typedef double T;
typedef vector<T> VT;
typedef vector<VT> VVT;
T GaussJordan (VVT &a, VVT &b)
       const int n = a.size();
        const int m = b[0].size();
       VI irow(n), icol(n), ipiv(n);
       T det = 1;
    for (int i = 0; i < n; i++)
       for (int k = 0; k < n; k++)
                                       if (!ipiv[k])
                                               if (pj == -1 || fabs(a[j][k]) > fabs(a[pj][pk])) { pj
                                                     = j; pk = k; }
               if (fabs(a[pj][pk]) < EPS)</pre>
                        cerr << "Matrix is singular." << endl;</pre>
                        exit(0);
        ipiv[pk]++;
       swap(a[pj], a[pk]);
       swap(b[pj], b[pk]);
if (pj != pk) det *= -1;
irow[i] = pj;
icol[i] = pk;
       T c = 1.0 / a[pk][pk];
       det *= a[pk][pk];
       a[pk][pk] = 1.0;
        for (int p = 0; p < n; p++)
               a[pk][p] *= c;
        for (int p = 0; p < m; p++)
               b[pk][p] *= c;
        for (int p = 0; p < n; p++)
               if (p != pk)
                       c = a[p][pk];
a[p][pk] = 0;
                       for (int p = n-1; p >= 0; p--)
               if (irow[p] != icol[p])
                       for (int k = 0; k < n; k++)
                               swap(a[k][irow[p]], a[k][icol[p]]);
       return det;
```

```
int main()
        const int n = 4;
        const int m = 2;
        double A[n][n] = \{ \{1,2,3,4\}, \{1,0,1,0\}, \{5,3,2,4\}, \{6,1,4,6\} \};
        double B[n][m] = \{ \{1,2\}, \{4,3\}, \{5,6\}, \{8,7\} \};
        VVT a(n), b(n);
        for (int i = 0; i < n; i++)</pre>
                a[i] = VT(A[i], A[i] + n);

b[i] = VT(B[i], B[i] + m);
        double det = GaussJordan(a, b);
        // expected: 60
        cout << "Determinant: " << det << endl;</pre>
        // expected: -0.233333 0.166667 0.133333 0.0666667
                      0.166667 0.166667 0.333333 -0.333333
                      0.233333 0.833333 -0.133333 -0.0666667
        // 0.05 -0.75 -0.1 0.2 cout << "Inverse: " << endl;
        for (int i = 0; i < n; i++)</pre>
                cout << endl:
        // expected: 1.63333 1.3
                     -0.166667 0.5
                      2.36667 1.7
                      -1.85 -1.35
        cout << "Solution: " << endl;
        for (int i = 0; i < n; i++)
                 for (int j = 0; j < m; j++)
                       cout << b[i][j] << ' ';
                 cout << endl;
```

8 Strings

8.1 Aho-Corasick Algorithm

```
Implementation - Benoit Chabod
Aho Corasick algorithm
*/
struct node
        map<char, int> g;
        vector<short> out;
        node(int fail = -1): f(fail) {}
};
vector<node> nodes;
void add_str(const string & s, int num)
        int cur = 0:
       int n = s.size();
        for(int i = 0; i < n; i++)
                auto it = nodes[cur].g.find(s[i]);
                if(it == nodes[cur].g.end())
                        nodes[cur].q[s[i]] = nodes.size();
                        cur = nodes.size();
                        nodes.push_back(node());
                else
                        cur = it->second:
        nodes[cur].out.push_back(num);
void init_fail()
```

```
int cur = 0;
        queue<int> q;
         q.push(cur);
    while( !q.empty() )
         cur = q.front();
        map<char, int>::iterator it;
        for(it = nodes[cur].g.begin(); it != nodes[cur].g.end(); it++)
                 int child = it->second:
                 int pfail = nodes[cur].f;
char ch = it->first;
                 map<char, int>::iterator f;
                 while( pfail != -1 && ((f = nodes[pfail].g.find(ch)) == nodes[pfail].g.end()) )
                           pfail = nodes[pfail].f;
                 nodes[child].f = (pfail == -1)? 0 : f -> second;
                 pfail = nodes[child].f;
                  nodes[child].out.insert(nodes[child].out.end(),nodes[pfail].out.begin(),nodes[pfail].
                 q.push(child);
         q.pop();
// Usage
void usage()
         nodes.push_back(node());
         for [each word] add_str(word,i)
                 init_fail();
         for [each letter]
                  map<char, int>::iterator f;
                 \textbf{while} ( \text{ cur } != -1 \text{ \&\& } ((f = \text{nodes}[\text{cur}].g.\text{find}(\text{letter})) == \text{nodes}[\text{cur}].g.\text{end}()) )
                          cur = nodes[cur].f;
                 if( cur == -1 )
                           cur = 0:
                           continue:
        cur = f->second;
        for(auto v : nodes[cur].out)
                 // Word v was found
```

8.2 Knuth-Morris-Pratt Algorithm

```
/*
KMP/Pi function
Note: cin >> (s+1) (the operations in the pi-function start at 1)
void preKmp()
         k=kmpNext[1]=0;
         for (int i=2;i<=n;++i)</pre>
                  \label{eq:while} \mbox{while} \mbox{ (k && p[k+1]!=p[i]) k=kmpNext[k];}
                  if(p[k+1]==p[i])
                           k++;
                  kmpNext[i]=k;
         int k=0;
         for (int i=1;i<=m;++i)</pre>
                  while (k \&\& p[k+1]!=s[i])
                           k=kmpNext[k];
                  if(p[k+1]==s[i])
                           k++;
                  if(k==n)
                           // here we have a match
```

```
k=kmpNext[k];
}
```

8.3 Suffix Array

```
//Suffix Array
struct SuffixArray
       const int L;
       string s;
       vector<vector<int> > P;
       vector<pair<int,int>,int> > M;
       SuffixArray(const string &s) : L(s.length()), s(s), P(1, vector<int>(L, 0)), M(L)
              P.push_back(vector<int>(L, 0));
                     for (int i = 0; i < L; i++)
                            M[i] = make_pair(make_pair(P[level-1][i], i + skip < L ? P[level-1][i</pre>
                                  + skip] : -1000), i);
                     sort(M.begin(), M.end());
                     level] [M[i-1].second] : i;
   vector<int> GetSuffixArray()
       return P.back();
        // returns the length of the longest common prefix of s[i...L-1] and s[j...L-1]
       int LongestCommonPrefix(int i, int j)
              int len = 0;
              if (i == j) return L - i;
              for (int k = P.size() - 1; k \ge 0 && i < L && j < L; k--)
                     if (P[k][i] == P[k][j])
                         i += 1 << k:
                         i += 1 << k;
                         len += 1 << k;
              return len;
};
int main()
       // bobocel is the O'th suffix
       // obocel is the 5'th suffix
           bocel is the 1'st suffix
            ocel is the 6'th suffix
            cel is the 2'nd suffix
             el is the 3'rd suffix
              1 is the 4'th suffix
       SuffixArray suffix("bobocel");
       vector<int> v = suffix.GetSuffixArray();
       // Expected output: 0 5 1 6 2 3 4
       for (int i = 0; i < v.size(); i++)</pre>
              cout << v[i] << " ";
       cout << endl;
       cout << suffix.LongestCommonPrefix(0, 2) << endl;</pre>
```

8.4 Manacher (Longest Palindromic Substring)

```
// Manacher (Longest Palindromic String) - O(n)
int lps[2*N+5];
char s[N];
int manacher() {
  int n = strlen(s);
```

```
string p (2*n+3, '#');
p[0] = '^';
for (int i = 0; i < n; i++) p[2*(i+1)] = s[i];
p[2*n+2] = '$';

int k = 0, r = 0, m = 0;
int l = p.length();
for (int i = 1; i < 1; i++) {
  int o = 2*k - i;
  lps[i] = (r > i) ? min(r-i, lps[o]) : 0;
  while (p[i + 1 + lps[i]] == p[i - 1 - lps[i]]) lps[i]++;
  if (i + lps[i] > r) k = i, r = i + lps[i];
  m = max(m, lps[i]);
} return m;
```

8.5 Z Function

```
// Z-Function - O(n)
// Array where ith element is greatest number of characters starting at i mathing the prefix
vector<int> zfunction(const string& s){
    vector<int> z (s.size());
    for (int i = 1, 1 = 0, r = 0, n = s.size(); i < n; i++){
        if (i < r) z[i] = min(z[i-1], r - i + 1);
        while (i + z[i] < n and s[z[i]] == s[z[i] + i]) z[i]++;
        if (i + z[i] - 1 > r) 1 = i, r = i + z[i] - 1;
    }
    return z;
}
```

9 Python Stuff...

10 Geometry

10.1 Convex Hull

```
def convex_hull(points):
    """Computes the convex hull of a set of 2D points.
    Input: an iterable sequence of (x,\ y) pairs representing the points.
    Output: a list of vertices of the convex hull in counter-clockwise order,
      starting from the vertex with the lexicographically smallest coordinates.
    Implements Andrew's monotone chain algorithm. O(n log n) complexity.
    # Sort the points lexicographically (tuples are compared lexicographically).
    # Remove duplicates to detect the case we have just one unique point.
    points = sorted(set(points))
    # Boring case: no points or a single point, possibly repeated multiple times.
    if len(points) <= 1:</pre>
        return points
    \# 2D cross product of OA and OB vectors, i.e. z-component of their 3D cross product.
    # Returns a positive value, if OAB makes a counter-clockwise turn,
# negative for clockwise turn, and zero if the points are collinear.
    def cross(o, a, b):
        return (a[0] - o[0]) * (b[1] - o[1]) - (a[1] - o[1]) * (b[0] - o[0])
    # Build lower hull
    lower = []
    for p in points:
        while len(lower) >= 2 and cross(lower[-2], lower[-1], p) <= 0:</pre>
            lower.pop()
        lower.append(p)
    # Build upper hull
     upper = []
    for p in reversed(points):
        while len(upper) >= 2 and cross(upper[-2], upper[-1], p) <= 0:</pre>
            upper.pop()
        upper.append(p)
    # Concatenation of the lower and upper hulls gives the convex hull.
    # Last point of each list is omitted because it is repeated at the beginning of the other list.
    return lower[:-1] + upper[:-1]
```

```
# Example: convex hull of a 10-by-10 grid.
assert convex_hull([(i//10, i%10) for i in range(100)]) == [(0, 0), (9, 0), (9, 9), (0, 9)]
```

11 Dynamic Programming

11.1 Max Sum Subarray (Kadane's Algorithm)

```
def maxSubArraySum(a,size):
    max_so_far = 0
    max_ending_here = 0
    for i in range(0, size):
        max_ending_here = max_ending_here + a[i]
    if max_ending_here < 0:
        max_ending_here = 0
    elif (max_so_far < max_ending_here):
        max_so_far = max_ending_here
    return max_so_far</pre>
```

11.2 Longest Common Subsequence

```
def lcs(X , Y):
    # find the length of the strings
    m = len(X)
    n = len(Y)
    # declaring the array for storing the dp values
    L = [[None] * (n+1) for i in xrange(m+1)]
    """Following steps build L[m+1][n+1] in bottom up fashion
    Note: L[i][j] contains length of LCS of X[0..i-1] and Y[0..j-1]"""
    for i in range(m+1):
        for j in range (n+1):
            if i == 0 or j == 0:
                L[i][j] = 0
            elif X[i-1] == Y[j-1]:
   L[i][j] = L[i-1][j-1]+1
             else:
                 L[i][j] = max(L[i-1][j], L[i][j-1])
    # L[m][n] contains the length of LCS of X[0..n-1] & Y[0..m-1]
    return L[m][n]
```

11.3 Levenshtein Distance

```
def levenshtein(s1, s2):
    if len(s1) < len(s2):
        return levenshtein(s2, s1)
    \# len(s1) >= len(s2)
    if len(s2) == 0:
        return len(s1)
    previous_row = range(len(s2) + 1)
    for i, c1 in enumerate(s1):
        current\_row = [i + 1]
        for j, c2 in enumerate(s2):
            insertions = previous_row[j + 1] + 1 # j+1 instead of j since previous_row and current_row
                  are one character longer
            deletions = current_row[j] + 1
                                                 # than s2
            substitutions = previous_row[j] + (c1 != c2)
            current_row.append(min(insertions, deletions, substitutions))
        previous_row = current_row
    return previous_row[-1]
```

11.4 Longest Increasing Subsequence

```
def lis(arr):
    n = len(arr)
# Declare the list (array) for LIS and initialize LIS
# values for all indexes
lis = [l]*n
```

```
# Compute optimized LIS values in bottom up manner
for i in range (1 , n):
    for j in range(0 , i):
        if arr[i] > arr[j] and lis[i] < lis[j] + 1 :
        lis[i] = lis[j] + 1

# Initialize maximum to 0 to get the maximum of all
# LIS
maximum = 0
# Pick maximum of all LIS values
for i in range(n):
    maximum = max(maximum , lis[i])
return maximum</pre>
```

12 Graphs/Trees

12.1 Graph structure example for our DFS and BFS algorithms

12.2 Breadth-First Search

```
def bfs(graph, start):
    visited, queue = set(), [start]
    while queue:
        vertex = queue.pop(0)
        if vertex not in visited:
            visited.add(vertex)
            queue.extend(graph[vertex] - visited)
    return visited
bfs(graph, 'A') # {'B', 'C', 'A', 'F', 'D', 'E'}
```

12.3 Breadth-First Search Paths

12.4 Breadth-First Search Shortest Path

```
def shortest_path(graph, start, goal):
    try:
        return next(bfs_paths(graph, start, goal))
    except StopIteration:
        return None
shortest_path(graph, 'A', 'F') # ['A', 'C', 'F']
```

12.5 Depth-First Search

```
def dfs(graph, start):
    visited, stack = set(), [start]
    while stack:
        vertex = stack.pop()
        if vertex not in visited:
            visited.add(vertex)
            stack.extend(graph[vertex] - visited)
    return visited

dfs(graph, 'A') # {'E', 'D', 'F', 'A', 'C', 'B'}
```

12.6 Depth-First Search Paths

12.7 Dijkstra's Algorithm

```
from collections import defaultdict
from heapq import *
def dijkstra(edges, f, t):
   g = defaultdict(list)
   for 1, r, c in edges:
        g[1].append((c,r))
   q, seen = [(0,f,())], set()
   while q:
        (cost, v1, path) = heappop(q)
        if v1 not in seen:
             seen.add(v1)
             path = (v1, path)
            if v1 == t: return (cost, path)
for c, v2 in g.get(v1, ()):
                 if v2 not in seen:
                     heappush (q, (cost+c, v2, path))
   return float("inf")
#Code example
edges = [("A", "B", 7), ("A", "D", 5), ("B", "C", 8), ("B", "D", 9), ("B", "E", 7), ("C", "E", 5)]
print dijkstra(edges, "A", "E") #(14, ('E', ('B', ('A', ()))))
```

12.8 Kruskal's Algorithm (including Merge-Find set)

```
parent = dict()
rank = dict()
def make set(vertice):
    parent[vertice] = vertice
    rank[vertice] = 0
def find(vertice):
    if parent[vertice] != vertice:
       parent[vertice] = find(parent[vertice])
    return parent[vertice]
def union(vertice1, vertice2):
    root1 = find(vertice1)
    root2 = find(vertice2)
    if root1 != root2:
       if rank[root1] > rank[root2]:
           parent[root2] = root1
       else:
           parent[root1] = root2
       if rank[root1] == rank[root2]: rank[root2] += 1
```

```
def kruskal (graph):
    for vertice in graph['vertices']:
        make_set (vertice)
        minimum_spanning_tree = set()
        edges = list(graph['edges'])
        edges.sort()
        fprint edges
    for edge in edges:
        weight, vertice1, vertice2 = edge
        if find(vertice1) != find(vertice2):
            union(vertice1, vertice2)
            minimum_spanning_tree.add(edge)
    return sorted(minimum spanning_tree)
```

12.9 Bellman-Ford Algorithm

```
# Step 1: For each node prepare the destination and predecessor
def initialize(graph, source):
    d = {} # Stands for destination
    p = {} # Stands for predecessor
    for node in graph:
        d[node] = float('Inf') # We start admitting that the rest of nodes are very very far
    d[source] = 0 # For the source we know how to reach
    return d, p
def relax(node, neighbour, graph, d, p):
    # If the distance between the node and the neighbour is lower than the one I have now
    if d[neighbour] > d[node] + graph[node][neighbour]:
        # Record this lower distance
        d[neighbour] = d[node] + graph[node][neighbour]
        p[neighbour] = node
def bellman_ford(graph, source):
    d, p = initialize(graph, source)
    for i in range(len(graph)-1): #Run this until is converges
        for u in graph:
             for v in graph[u]: #For each neighbour of u
                 relax(u, v, graph, d, p) #Lets relax it
    # Step 3: check for negative-weight cycles
    for u in graph:
        for v in graph[u]:
             assert d[v] \le d[u] + graph[u][v]
    return d. p
def test():
    graph = +
         'a': {'b': -1, 'c': 4},
        'b': {'c': 3, 'd': 2, 'e': 2},
        'c': {},
'd': {'b': 1, 'c': 5},
        'e': {'d': -3}
    d, p = bellman_ford(graph, 'a')
    # d = {'a':0, 'b':-1, 'c':2, 'd':-2, 'e':1},
    # p = {'a':None, 'b':'a', 'c':'b', 'd':'e', 'e':'b'}
```

12.10 Floyd-Warshall Algorithm

```
# Number of vertices in the graph
V = 4

# Define infinity as the large enough value. This value will be
# used for vertices not connected to each other
INF = 99999

# Solves all pair shortest path via Floyd Warshall Algorithm
def floydWarshall(graph):

""" dist[][] will be the output matrix that will finally
have the shortest distances between every pair of vertices """
""" initializing the solution matrix same as input graph matrix
OR we can say that the initial values of shortest distances
are based on shortest paths considerting no
intermedidate vertices """
dist = map(lambda i : map(lambda j : j , i) , graph)

""" Add all vertices one by one to the set of intermediate
vertices.
---> Before start of a iteration, we have shortest distances
```

```
between all pairs of vertices such that the shortest
     distances consider only the vertices in set
    \{0, 1, 2, \ldots k-1\} as intermediate vertices.
        --> After the end of a iteration, vertex no. k is
     added to the set of intermediate vertices and the
    set becomes {0, 1, 2, .. k}
    for k in range (V):
        # pick all vertices as source one by one
        for i in range(V):
            # Pick all vertices as destination for the
            # above picked source
            for j in range(V):
                # If vertex k is on the shortest path from
                # i to j, then update the value of dist[i][j]
                dist[i][j] = min(dist[i][j] ,
                                  dist[i][k]+ dist[k][j]
    printSolution(dist)
            10
       (0) ----> (3)
       (1) ----> (2)
graph = [[0, 5, INF, 10],
             [INF, 0, 3, INF],
             [INF, INF, 0,
             [INF, INF, INF, 0]
floydWarshall(graph) # [[0,5,8,9],[INF,0,3,4],[INF,INF,0,1],[INF,INF,INF,0]]
```

12.11 Max Flow (Ford-Fulkerson Algorithm)

```
from collections import defaultdict
#This class represents a directed graph using adjacency matrix representation
    def __init__(self,graph):
        self.graph = graph # residual graph
self.ROW = len(graph)
        \#self.COL = len(gr[0])
    ""Returns true if there is a path from source 's' to sink 't' in
    residual graph. Also fills parent[] to store the path '''
    def BFS(self,s, t, parent):
        # Mark all the vertices as not visited
        visited =[False] * (self.ROW)
        # Create a queue for BFS
        # Mark the source node as visited and enqueue it
        queue.append(s)
        visited[s] = True
        # Standard BFS Loop
        while queue:
            #Dequeue a vertex from queue and print it
            u = queue.pop(0)
             # Get all adjacent vertices of the dequeued vertex u
            # If a adjacent has not been visited, then mark it
             # visited and enqueue it
            for ind, val in enumerate(self.graph[u]):
                if visited[ind] == False and val > 0 :
                     queue.append(ind)
                     visited[ind] = True
                     parent[ind] = u
        \ensuremath{\text{\#}} If we reached sink in BFS starting from source, then return
        # true, else false
        return True if visited[t] else False
```

```
# Returns the maximum flow from s to t in the given graph
    def FordFulkerson(self, source, sink):
        # This array is filled by BFS and to store path
        parent = [-1] * (self.ROW)
        max_flow = 0 # There is no flow initially
        # Augment the flow while there is path from source to sink
        while self.BFS(source, sink, parent) :
            # Find minimum residual capacity of the edges along the
            # path filled by BFS. Or we can say find the maximum flow
            # through the path found.
            path flow = float("Inf")
              = sink
            while(s != source):
               path_flow = min (path_flow, self.graph[parent[s]][s])
                s = parent[s]
            # Add path flow to overall flow
            max_flow += path_flow
            # update residual capacities of the edges and reverse edges
            # along the path
              = sink
            while (v != source):
               u = parent[v]
                self.graph[u][v] -= path_flow
                self.graph[v][u] += path_flow
                v = parent[v]
        return max flow
# Create a graph given in the above diagram
graph = [[0, 16, 13, 0, 0, 0],
        [0, 0, 10, 12, 0, 0],
        [0, 4, 0, 0, 14, 0],
        [0, 0, 9, 0, 0, 20],
        [0, 0, 0, 7, 0, 4],
        [0, 0, 0, 0, 0, 0]]
g = Graph(graph)
source = 0; sink = 5
print ("The maximum possible flow is %d " % g.FordFulkerson(source, sink))
```

12.12 Segment Tree

```
#encoding:utf-8
class SegmentTree(object):
   def init (self, start, end):
       self.start = start
       self.end = end
       self.max_value = {}
       self.sum_value = {}
        self.len_value = {}
        self._init(start, end)
   def add(self, start, end, weight=1):
        start = max(start, self.start)
        end = min(end, self.end)
        self._add(start, end, weight, self.start, self.end)
       return True
   def guery max(self, start, end):
       return self, query max(start, end, self, start, self, end)
   def query_sum(self, start, end):
        return self._query_sum(start, end, self.start, self.end)
   def query_len(self, start, end):
        return self._query_len(start, end, self.start, self.end)
   def _init(self, start, end):
        self.max_value[(start, end)] = 0
        self.sum_value[(start, end)] = 0
        self.len_value[(start, end)] = 0
       if start < end:</pre>
           mid = start + int((end - start) / 2)
           self. init(start, mid)
           self._init(mid+1, end)
```

```
def _add(self, start, end, weight, in_start, in_end):
    key = (in_start, in_end)
   if in_start == in_end:
       self.max_value[key] += weight
        self.sum_value[key] += weight
        self.len_value[key] = 1 if self.sum_value[key] > 0 else 0
   mid = in_start + int((in_end - in_start) / 2)
   if mid >= end:
       self._add(start, end, weight, in_start, mid)
   elif mid+1 <= start:</pre>
       self._add(start, end, weight, mid+1, in_end)
   else:
        self._add(start, mid, weight, in_start, mid)
        self._add(mid+1, end, weight, mid+1, in_end)
    self.max_value[key] = max(self.max_value[(in_start, mid)], self.max_value[(mid+1, in_end)])
    self.sum_value[key] = self.sum_value[(in_start, mid)] + self.sum_value[(mid+1, in_end)]
    self.len_value[key] = self.len_value[(in_start, mid)] + self.len_value[(mid+1, in_end)]
def _query_max(self, start, end, in_start, in_end):
   if start == in_start and end == in_end:
       ans = self.max_value[(start, end)]
   else.
        mid = in_start + int((in_end - in_start) / 2)
        if mid >= end:
            ans = self._query_max(start, end, in_start, mid)
        elif mid+1 <= start:</pre>
           ans = self._query_max(start, end, mid+1, in_end)
        else:
            ans = max(self._query_max(start, mid, in_start, mid),
                   self._query_max(mid+1, end, mid+1, in_end))
    #print start, end, in_start, in_end, ans
    return ans
def _query_sum(self, start, end, in_start, in_end):
   if start == in_start and end == in_end:
       ans = self.sum_value[(start, end)]
   else:
        mid = in_start + int((in_end - in_start) / 2)
        if mid >= end:
            ans = self._query_sum(start, end, in_start, mid)
        elif mid+1 <= start:
            ans = self._query_sum(start, end, mid+1, in_end)
        else:
            ans = self._query_sum(start, mid, in_start, mid) + self._query_sum(mid+1, end, mid+1,
                  in end)
    return ans
def _query_len(self, start, end, in_start, in_end):
   if start == in_start and end == in_end:
       ans = self.len_value[(start, end)]
    else.
        mid = in_start + int((in_end - in_start) / 2)
        if mid >= end:
            ans = self. guery len(start, end, in start, mid)
        elif mid+1 <= start:</pre>
           ans = self. guery len(start, end, mid+1, in end)
        else:
            ans = self._query_len(start, mid, in_start, mid) + self._query_len(mid+1, end, mid+1,
                  in_end)
    #print start, end, in_start, in_end, ans
```

13 Math

13.1 Prime Number Sieve (generator)

```
from itertools import count
def postponed_sieve():
                                          # postponed sieve, by Will Ness
    yield 2; yield 3; yield 5; yield 7;
                                          # original code David Eppstein,
                                              Alex Martelli, ActiveState Recipe 2002
    ps = postponed_sieve()
                                           # a separate base Primes Supply:
    p = next(ps) and next(ps)
                                          # (3) a Prime to add to dict
                                          # (9) its sQuare
    for c in count (9,2):
                                          # the Candidate
        if c in sieve:
                                       # c's a multiple of some base prime
            s = sieve.pop(c)
                                          i.e. a composite ; or
        elif c < q:
    vield c</pre>
                                      # a prime
             continue
                                      # or the next base prime's square:
        else: # (c==q):
            s=count (q+2*p, 2*p)
                                           (9+6, by 6: 15, 21, 27, 33, ...)
            p=next(ps)
```

13.2 GCD and Euler's Totient Function

```
# Function to return gcd of a and b
def gcd(a, b):
    if a == 0:
        return b
    return gcd(b%a, a)

# A simple method to evaluate Euler Totient Function
def phi(n):
    result = 1
    for i in range(2, n):
        if gcd(i, n) == 1:
            result = result + 1
    return result
```

13.3 Miller-Rabin Primality Test

```
def miller_rabin(n, k):
    # The optimal number of rounds (k) for this test is 40
    # for justification
    if n == 2:
        return True
    if n % 2 == 0:
        return False
       s = 0, n - 1
    while s % 2 == 0:
        r += 1
        s //= 2
    for \_ in xrange(k):
        a = random.randrange(2, n - 1)
        x = pow(a, s, n)
        if x == 1 or x == n - 1:
            continue
        for \underline{\phantom{a}} in xrange (r - 1):
             x = pow(x, 2, n)
            if x == n - 1:
                break
        else.
            return False
    return True
```

13.4 Gauss-Jordan Elimination (Matrix inversion and linear system solving)

```
def gauss_jordan(m, eps = 1.0/(10**10)):
  """Puts given matrix (2D array) into the Reduced Row Echelon Form.
     Returns True if successful, False if 'm' is singular.
     NOTE: make sure all the matrix items support fractions! Int matrix will NOT work!
     Written by Jarno Elonen in April 2005, released into Public Domain"""
  (h, w) = (len(m), len(m[0]))
  for y in range(0,h):
   maxrow = v
   for y2 in range(y+1, h): # Find max pivot
     if abs(m[y2][y]) > abs(m[maxrow][y]):
       maxrow = y2
    (m[y], m[maxrow]) = (m[maxrow], m[y])
   if abs(m[y][y]) <= eps:
   for y2 in range(y+1, h):
                               # Eliminate column y
      c = m[y2][y] / m[y][y]
     for x in range(y, w):
       m[y2][x] = m[y][x] * c
  for y in range(h-1, 0-1, -1): # Backsubstitute
   for y2 in range(0,y):
     for x in range (w-1, y-1, -1):
       m[y2][x] = m[y][x] * m[y2][y] / c
   m[y][y] /= c
   for x in range(h, w):
                               # Normalize row y
```

```
m[y][x] /= c
  return True
def solve(M, b):
  solves M*x = b
  return vector x so that M*x = b
  :param M: a matrix in the form of a list of list
  :param b: a vector in the form of a simple list of scalars """
  m2 = [row[:]+[right] for row, right in zip(M,b) ]
  return [row[-1] for row in m2] if gauss_jordan(m2) else None
def inv(M):
  return the inv of the matrix M
  #clone the matrix and append the identity matrix
  \# [int(i==j) for j in range_M] is nothing but the i(th row of the identity matrix
  m2 = [row[:]+[int(i==j) for j in range(len(M))] for i,row in enumerate(M)]
  # extract the appended matrix (kind of m2[m:,...]
  return [row[len(M[0]):] for row in m2] if gauss_jordan(m2) else None
def zeros( s , zero=0):
    return a matrix of size 'size'
    :param size: a tuple containing dimensions of the matrix
    :param zero: the value to use to fill the matrix (by default it's zero )
    return [zeros(s[1:]) for i in range(s[0])] if not len(s) else zero
```

14 Strings

14.1 Knuth-Morris-Pratt Algorithm (fast pattern matching)

```
def KnuthMorrisPratt(text, pattern):
    ^{\prime\prime\prime}Yields all starting positions of copies of the pattern in the text.
Calling conventions are similar to string.find, but its arguments can be
lists or iterators, not just strings, it returns all matches, not just
the first one, and it does not need the whole text in memory at once.
Whenever it yields, it will have read the text exactly up to and including
the match that caused the yield.'''
    # allow indexing into pattern and protect against change during yield
    pattern = list(pattern)
    # build table of shift amounts
    shifts = [1] * (len(pattern) + 1)
    shift = 1
    for pos in range(len(pattern)):
        while shift <= pos and pattern[pos] != pattern[pos-shift]:</pre>
           shift += shifts[pos-shift]
        shifts[pos+1] = shift
    # do the actual search
    startPos = 0
        while matchLen == len(pattern) or \
             matchLen >= 0 and pattern[matchLen] != c:
            startPos += shifts[matchLen]
            matchLen -= shifts[matchLen]
        matchLen += 1
        if matchLen == len(pattern):
            vield startPos
```

14.2 Rabin-Karp Algorithm (multiple pattern matching)

```
# d is the number of characters in input alphabet
d = 256
# pat -> pattern
# txt -> text
# q -> A prime number

def search(pat, txt, q):
    M = len(pat)
    N = len(txt)
    i = 0
    i = 0
```

```
j+=1
    # if p == t and pat[0...M-1] = txt[i, i+1, ...i+M-1]
    if j==M:
        print "Pattern found at index " + str(i)

# Calculate hash value for next window of text: Remove
# leading digit, add trailing digit
if i < N-M:
        t = (d*(t-ord(txt[i])*h) + ord(txt[i+M]))%q

# We might get negative values of t, converting it to
# positive
    if t < 0:
        t = t+q

# Driver program to test the above function
txt = "GEEKS FOR GEEKS"
pat = 101 # A prime number
search(pat,txt,q)</pre>
```