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adapted from KACTL and MIT NULL 2022-11-14

1 Contest

const int MOD = 1e9+7;

const db PI = acos((db)-1);

mt19937 rng(0); // or mt19937\_64

### TemplateShort .bashrc hash troubleshoot

1

```
2 Mathematics
3 Data Structures
                                                             \mathbf{2}
   Number Theory
                                                             \mathbf{3}
5 Combinatorial
  Numerical
   Graphs
8 Geometry
9 Strings
10 Various
Contest (1)
TemplateShort.cpp
                                                   77aa31, 44 lines
#include <bits/stdc++.h>
using namespace std;
using 11 = long long;
using db = long double; // or double if tight TL
using str = string;
using pi = pair<int,int>;
#define mp make_pair
#define f first
#define s second
#define tcT template<class T
tcT> using V = vector<T>;
tcT, size_t SZ> using AR = array<T,SZ>;
using vi = V<int>;
using vb = V<bool>;
using vpi = V<pi>;
#define sz(x) int((x).size())
#define all(x) begin(x), end(x)
#define sor(x) sort(all(x))
#define rsz resize
#define pb push_back
#define ft front()
#define bk back()
#define FOR(i,a,b) for (int i = (a); i < (b); ++i)
#define F0R(i,a) FOR(i,0,a)
#define ROF(i,a,b) for (int i = (b)-1; i \ge (a); --i)
#define R0F(i,a) ROF(i,0,a)
#define rep(a) FOR( ,a)
#define each(a,x) for (auto& a: x)
```

```
tcT> bool ckmin(T& a, const T& b) {
 return b < a ? a = b, 1 : 0; } // set a = min(a,b)
tcT> bool ckmax(T& a, const T& b) {
 return a < b ? a = b, 1 : 0; } // set a = max(a,b)
int main() { cin.tie(0)->sync_with_stdio(0); }
.bashrc
alias clr="printf '\33c'"
co() { g++ -std=c++17 -O2 -Wall -Wextra -Wshadow -Wconversion -
  →o $1 $1.cpp; }
run() { co $1 && ./$1; }
hash.sh
# Hash file ignoring whitespace and comments. Verifies that
# code was correctly typed. Usage: 'sh hash.sh < A.cpp'
cpp -dD -P -fpreprocessed|tr -d '[:space:]'|md5sum|cut -c-6
troubleshoot.txt
General:
Write down most of your thoughts, even if you're not sure
whether they're useful.
Give your variables (and files) meaningful names.
Stay organized and don't leave papers all over the place!
You should know what your code is doing ...
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?
Remove debug output.
Make sure to submit the right file.
Wrong answer:
Print your solution! Print debug output as well.
Read the full problem statement again.
Have you understood the problem correctly?
Are you sure your algorithm works?
Try writing a slow (but correct) solution.
Can your algorithm handle the whole range of input?
Did you consider corner cases (ex. n=1)?
Is your output format correct? (including whitespace)
Are you clearing all data structures between test cases?
Any uninitialized variables?
Any undefined behavior (array out of bounds)?
Any overflows or NaNs (or shifting 11 by >=64 bits)?
Confusing N and M, i and j, etc.?
Confusing ++i and i++?
Return vs continue vs break?
Are you sure the STL functions you use work as you think?
Add some assertions, maybe resubmit.
Create some test cases 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.
Rewrite your solution from the start or let a teammate do it.
Geometry:
Work with ints if possible.
Correctly account for numbers close to (but not) zero. Related:
for functions like acos make sure absolute val of input is not
```

(slightly) greater than one.

```
Correctly deal with vertices that are collinear, concyclic,
coplanar (in 3D), etc.
Subtracting a point from every other (but not itself)?
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's your complexity? Large TL does not mean that something
simple (like NlogN) isn't intended.
Are you copying a lot of unnecessary data? (References)
Avoid vector, map. (use arrays/unordered_map)
How big is the input and output? (consider FastIO)
What do your teammates think about your algorithm?
Calling count() on multiset?
```

What is the max amount of memory your algorithm should need?

Are you clearing all data structures between test cases?

# Mathematics (2)

If using pointers try BumpAllocator.

Memory limit exceeded:

# 2.1 Trigonometry

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

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

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

$$\sin v + \sin w = 2\sin \frac{v+w}{2}\cos \frac{v-w}{2}$$

$$\cos v + \cos w = 2\cos \frac{v+w}{2}\cos \frac{v-w}{2}$$

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

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

### 2.2 Geometry

### 2.2.1 Triangles

Side lengths: a, b, cSemiperimeter:  $s = \frac{a+b+c}{2}$ Area:  $A = \sqrt{s(s-a)(s-b)(s-c)}$ 

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

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

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

Length of bisector (divides angles in two):

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

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

Law of tangents:  $\frac{a+b}{a-b} = \frac{\tan \frac{\alpha+\beta}{2}}{\tan \frac{\alpha-\beta}{2}}$ 

### Derivatives/Integrals

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

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

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

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

Integration by parts:

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

# 2.4 Sums/Series

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

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

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

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

# Data Structures (3)

### 3.1 STL

```
MapComparator.h
```

Description: example of function object (functor) for map or set Usage: set<int,cmp> s; map<int,int,cmp> m;

struct cmp{bool operator()(int l,int r)const{return l>r;}};

### HashMap.h

Description: Hash map with similar API as unordered\_map. Initial capacity must be a power of 2 if provided.

Usage: ht<int, int> h({},{},{},{},{1<<16}); Memory:  $\sim 1.5x$ 

Time:  $\sim 3x$  faster

<ext/pb\_ds/assoc\_container.hpp> using namespace gnu pbds; struct chash { const uint64\_t C = 11(4e18\*acos(0))+71; // large odd number const int RANDOM = rng(); 11 operator()(11 x) const { return \_\_builtin\_bswap64((x^  $\hookrightarrow$ RANDOM) \*C); } template<class K, class V> using ht = gp\_hash\_table<K, V, chash>; template<class K, class V> V get(ht<K, V>& u, K x) { auto it = u.find(x); return it == end(u) ? 0 : it->s; }

### OrderStatisticTree.h

**Description:** A set (not multiset!) with support for finding the n'th element, and finding the index of an element. Change null\_type to get a

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

```
<ext/pb_ds/assoc_container.hpp>
using namespace __gnu_pbds;
tcT> using Tree = tree<T, null_type, less<T>,
 rb_tree_tag, tree_order_statistics_node_update>;
#define ook order_of_key
#define fbo find_by_order
void treeExample() {
 Tree<int> t, t2; t.insert(8);
 auto it = t.insert(10).f; assert(it == t.lb(9));
 assert(t.ook(10) == 1 && t.ook(11) == 2 && *t.fbo(0) == 8);
 t.join(t2); // assuming T < T2 or T > T2, merge t2 into t
```

#### LineContainer.h

**Description:** Add lines of the form ax + b, query maximum y-coordinate for any x.

Time:  $\mathcal{O}(\log N)$ using T = 11; const T INF = LLONG\_MAX; // a/b rounded down // ll fdiv(ll a, ll b) { return a/b-((a^b)<0&&a%b); } bool  $_Q = 0;$ struct Line { T a, b; mutable T lst; T eval(T x) const { return a\*x+b; } bool operator<(const Line&o)const{return \_Q?lst<o.lst:a<o.a;}</pre> T last\_gre(const Line& o) const { assert(a <= o.a);</pre> // greatest x s.t. a\*x+b >= o.a\*x+o.breturn lst=(a==o.a?(b>=o.b?INF:-INF):fdiv(b-o.b,o.a-a));} struct LineContainer: multiset<Line> { bool isect(iterator it) { auto n it = next(it);

```
if (n_it == end()) return it->lst = INF, 0;
  return it->last_gre(*n_it) >= n_it->lst; }
void add(T a, T b) {
  auto it = ins(\{a,b,0\}); while (isect(it)) erase(next(it));
  if (it == begin()) return;
  if (isect(--it)) erase(next(it)), isect(it);
  while (it != begin()) {
    --it; if (it->lst < next(it)->lst) break;
    erase(next(it)); isect(it); }
T qmax(T x) { assert(!empty());
  _Q = 1; T res = lb(\{0,0,x\}) \rightarrow eval(x); _Q = 0;
  return res; }
```

#### 3.21D Range Queries

### RMQ.h

Description: 1D range minimum query. If TL is an issue, use arrays instead of vectors and store values instead of indices.

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

```
Time: \mathcal{O}(1)
                                                        a3f881, 19 lines
tcT> struct RMO {
 int level(int x) { return 31-__builtin_clz(x); }
 V<T> v; V<vi> jmp;
  int cmb(int a, int b) {
    return v[a] == v[b]?min(a,b):(v[a] < v[b]?a:b);}
  void init(const V<T>& _v) {
    v = v; jmp = \{vi(sz(v))\};
    iota(all(jmp[0]),0);
```

for (int j = 1; 1 << j <= sz(v); ++j) { jmp.pb(vi(sz(v) - (1 << j) +1));FOR(i,sz(jmp[j])) jmp[j][i] = cmb(jmp[j-1][i],jmp[j-1][i+(1<<(j-1))]);int index(int 1, int r) { assert( $1 \le r$ ); int d = level(r-l+1); return cmb(jmp[d][1],jmp[d][r-(1<<d)+1]); } T query(int 1, int r) { return v[index(1,r)]; }

SegmentTree.h

Description: 1D point update and range query where cmb is any associative operation. seg[1] = guery(0, N-1).

Time:  $\mathcal{O}(\log N)$ 1630f3, 18 lines

```
tcT> struct SegTree { // cmb(ID,b) = b
 const T ID{}; T cmb(T a, T b) { return a+b; }
 int n; V<T> seg;
 void init(int _n) { // upd, query also work if n = _n
   for (n = 1; n < _n; ) n *= 2;
   seq.assign(2*n,ID);}
 void pull(int p) { seg[p] = cmb(seg[2*p], seg[2*p+1]); }
 void upd(int p, T val) { // set val at position p
   seg[p += n] = val; for (p /= 2; p; p /= 2) pull(p); }
 T query(int 1, int r) { // zero-indexed, inclusive
   T ra = ID, rb = ID;
    for (1 += n, r += n+1; 1 < r; 1 /= 2, r /= 2) {
     if (1&1) ra = cmb(ra, seg[1++]);
      if (r\&1) rb = cmb(seq[--r],rb);
   return cmb(ra,rb);
};
```

2672f9, 24 lines

```
LazySegmentTree.h
```

Description: 1D range increment and sum query.

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

78a06d, 26 lines

```
tcT, int SZ> struct LazySeg {
  static_assert(pct(SZ) == 1); // SZ must be power of 2
  const T ID{}; T cmb(T a, T b) { return a+b; }
  T seg[2*SZ], lazy[2*SZ];
  LazySeg() { FOR(i,2*SZ) seg[i] = lazy[i] = ID; }
  void push(int ind, int L, int R) {
    seg[ind] += (R-L+1)*lazy[ind]; // dependent on operation
   if (L != R) F0R(i,2) lazy[2*ind+i] += lazy[ind];
   lazy[ind] = 0;
  } // recalc values for current node
  void pull(int ind) {seg[ind]=cmb(seg[2*ind], seg[2*ind+1]);}
  void build() { ROF(i,1,SZ) pull(i); }
  void upd(int lo,int hi,T inc,int ind=1,int L=0, int R=SZ-1) {
    push(ind, L, R); if (hi < L || R < lo) return;
    if (lo <= L && R <= hi) {
     lazy[ind] = inc; push(ind, L, R); return; }
    int M = (L+R)/2; upd(lo,hi,inc,2*ind,L,M);
    upd(lo,hi,inc,2*ind+1,M+1,R); pull(ind);
  T query(int lo, int hi, int ind=1, int L=0, int R=SZ-1) {
    push(ind,L,R); if (lo > R || L > hi) return ID;
    if (lo <= L && R <= hi) return seg[ind];</pre>
    int M = (L+R)/2; return cmb(query(lo,hi,2*ind,L,M),
      query(lo,hi,2*ind+1,M+1,R));
};
```

### PSeg.h

**Description:** Persistent min segtree with lazy updates, no propagation. If making d a vector then save the results of upd and build in local variables first to avoid issues when vector resizes in C++14 or lower.

```
Memory: \mathcal{O}(N + Q \log N)
                                                      8f37fa, 46 lines
tcT, int SZ> struct pseq {
  static const int LIM = 2e7;
  struct node {
   int 1, r; T val = 0, lazy = 0;
   void inc(T x) { lazy += x; }
   T get() { return val+lazy; }
  node d[LIM]; int nex = 0;
  int copy(int c) { d[nex] = d[c]; return nex++; }
  T cmb(T a, T b) { return min(a,b); }
  void pull(int c) { d[c].val =
   cmb(d[d[c].1].get(), d[d[c].r].get()); }
  //// MAIN FUNCTIONS
  T query(int c, int lo, int hi, int L, int R) {
   if (lo <= L && R <= hi) return d[c].get();</pre>
   if (R < lo || hi < L) return MOD;
   int M = (L+R)/2;
   return d[c].lazy+cmb(query(d[c].l,lo,hi,L,M),
              query(d[c].r,lo,hi,M+1,R));
  int upd(int c, int lo, int hi, T v, int L, int R) {
   if (R < lo || hi < L) return c;
    int x = copy(c);
   if (lo <= L && R <= hi) { d[x].inc(v); return x; }</pre>
    int M = (L+R)/2;
    d[x].1 = upd(d[x].1, lo, hi, v, L, M);
   d[x].r = upd(d[x].r,lo,hi,v,M+1,R);
   pull(x); return x;
  int build(const V<T>& arr, int L, int R) {
   int c = nex++;
   if (L == R) {
```

```
if (L < sz(arr)) d[c].val = arr[L];</pre>
    return c;
  int M = (L+R)/2;
  d[c].l = build(arr, L, M), d[c].r = build(arr, M+1, R);
  pull(c); return c;
vi loc; //// PUBLIC
void upd(int lo, int hi, T v) {
  loc.pb(upd(loc.bk,lo,hi,v,0,SZ-1)); }
T query(int ti, int lo, int hi) {
  return query(loc[ti],lo,hi,0,SZ-1); }
void build(const V<T>&arr) {loc.pb(build(arr, 0, SZ-1));}
```

### Treap.h

tnode(int val) {

if (!t) return {t,t};

if (getsz(t->c[0]) >= sz) {

return {p.f,calc(t)};

auto p = splitsz(t->c[0],sz); t->c[0] = p.s;

auto p=splitsz(t->c[1],sz-getsz(t->c[0])-1); t->c[1]=p.f;

pri = rng(); sum = val = \_val;

Description: Easy BBST. Use split and merge to implement insert and delete.

```
Time: \mathcal{O}(\log N)
                                                        bdb758, 65 lines
using pt = struct tnode*;
struct tnode {
 int pri, val; pt c[2]; // essential
 int sz; 11 sum; // for range queries
 bool flip = 0; // lazy update
```

```
sz = 1; c[0] = c[1] = nullptr;
 ~tnode() { FOR(i,2) delete c[i]; }
int getsz(pt x) { return x?x->sz:0; }
11 getsum(pt x) { return x?x->sum:0; }
pt prop(pt x) { // lazy propagation
 if (!x || !x->flip) return x;
  swap (x->c[0], x->c[1]);
 x->flip = 0; FOR(i,2) if (x->c[i]) x->c[i]->flip ^= 1;
 return x:
pt calc(pt x) {
 pt a = x->c[0], b = x->c[1];
 assert(!x->flip); prop(a), prop(b);
 x->sz = 1+getsz(a)+getsz(b);
 x->sum = x->val+getsum(a)+getsum(b);
 return x;
void tour(pt x, vi& v) { // print values of nodes,
 if (!x) return; // inorder traversal
 prop(x); tour(x->c[0],v); v.pb(x->val); tour(x->c[1],v);
pair<pt, pt> split(pt t, int v) { // >= v goes to the right
 if (!t) return {t,t};
 prop(t);
 if (t->val >= v) {
    auto p = split(t->c[0], v); t->c[0] = p.s;
    return {p.f,calc(t)};
 } else {
    auto p = split(t->c[1], v); t->c[1] = p.f;
    return {calc(t),p.s};
pair<pt,pt> splitsz(pt t, int sz) { // sz nodes go to left
```

```
return {calc(t),p.s};
pt merge(pt 1, pt r) { // keys in 1 < keys in r
 if (!1 || !r) return 1?:r;
 prop(l), prop(r); pt t;
 if (1->pri > r->pri) 1->c[1] = merge(1->c[1],r), t = 1;
  else r - c[0] = merge(1, r - c[0]), t = r;
  return calc(t);
pt ins(pt x, int v) { // insert v
 auto a = split(x,v), b = split(a.s,v+1);
 return merge(a.f, merge(new tnode(v),b.s)); }
pt del(pt x, int v) { // delete v
 auto a = split(x,v), b = split(a.s,v+1);
  return merge(a.f,b.s); }
```

# Number Theory (4)

### 4.1 Modular Arithmetic

### ModIntShort.h

**Description:** Modular arithmetic. Assumes MOD is prime.

```
Usage: mi a = MOD+5; inv(a); // 400000003
```

```
template<int MOD, int RT> struct mint {
 static const int mod = MOD;
 static constexpr mint rt() { return RT; } // primitive root
 explicit operator int() const { return v; }
 mint():v(0) {}
 mint(ll _v):v(int(_v%MOD)) \{ v += (v<0)*MOD; \}
 mint& operator+=(mint o) {
   if ((v += o.v) >= MOD) v -= MOD;
   return *this; }
 mint& operator-=(mint o) {
   if ((v -= o.v) < 0) v += MOD;
   return *this; }
 mint& operator*=(mint o) {
   v = int((11)v*o.v%MOD); return *this; }
 friend mint pow(mint a, ll p) { assert(p >= 0);
   return p==0?1:pow(a*a,p/2)*(p&1?a:1); }
 friend mint inv(mint a) { assert(a.v != 0); return pow(a, MOD
```

#### ModFact.h

using vmi = V<mi>;

using mi = mint < (int) 1e9 + 7, 5>;

**Description:** Combinations modulo a prime MOD. Assumes  $SZ \leq MOD$ . Usage: Factorials F; F.init(10); F.C(6, 4); // 15 Time:  $\mathcal{O}\left(SZ\right)$ 

friend mint operator+(mint a, mint b) { return a += b; }

friend mint operator-(mint a, mint b) { return a -= b; }

friend mint operator\*(mint a, mint b) { return a \*= b; }

```
"ModInt.h"
                                                         63a7f9, 13 lines
struct Factorials {
 vmi invs, fac, ifac;
 void init(int N) { // idempotent
    invs.rsz(N), fac.rsz(N), ifac.rsz(N);
    invs[1] = fac[0] = ifac[0] = 1;
    FOR (i, 2, N) invs[i] = mi(-(11) MOD/i*(int) invs[MOD%i]);
    FOR(i,1,N) fac[i] = fac[i-1]*i, ifac[i] = ifac[i-1]*invs[i
       \hookrightarrow1;
 mi C(int a, int b) {
```

```
if (a < b || b < 0) return 0;
return fac[a]*ifac[b]*ifac[a-b];
}
};</pre>
```

### ModMulLL.h

**Description:** Multiply two 64-bit integers mod another if 128-bit is not available. modMul is equivalent to (u1) (\_int128(a) \*b\$mod). Works for  $0 \le a, b < mod < 2^{63}$ .

```
using ul = uint64_t;
ul modMul(ul a, ul b, const ul mod) {
    ll ret = a*b-mod*(ul) ((db)a*b/mod);
    return ret+((ret<0)-(ret>=(l1)mod))*mod; }
ul modPow(ul a, ul b, const ul mod) {
    if (b == 0) return 1;
    ul res = modPow(a,b/2,mod); res = modMul(res,res,mod);
    return b&1 ? modMul(res,a,mod) : res;
}
```

### FastMod.h

**Description:** Barrett reduction computes a%b about 4 times faster than usual where b>1 is constant but not known at compile time. Division by b is replaced by multiplication by m and shifting right 64 bits.

```
using ul = uint64_t; using L = __uint128_t;
struct FastMod {
  ul b, m; FastMod(ul b) : b(b), m(-1ULL / b) {}
  ul reduce(ul a) {
    ul q = (ul)((__uint128_t(m) * a) >> 64), r = a - q * b;
    return r - (r >= b) * b; }
};
```

### ModSqrt.h

**Description:** To nelli-Shanks algorithm for square roots mod a prime. -1 if doesn't exist.

```
Usage: sqrt(mi((11)1e10)); // 100000 Time: O(\log^2(MOD))
```

#### ModSum.h

**Description:** Counts # of lattice points (x, y) in the triangle  $1 \le x, 1 \le y, ax + by \le s \pmod{2^{64}}$  and related quantities.

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

23cbf6, 20 lines

```
using ul = uint64_t;
ul sum2(ul n) { return n/2*((n-1)|1); } // sum(0..n-1)
// \return | {(x,y) | 1 <= x, 1 <= y, a*x+b*y <= S}|
// = sum_{i=1}^{qs} (S-a*i)/b
ul triSum(ul a, ul b, ul s) { assert(a > 0 && b > 0);
ul qs = s/a, rs = s%a; // ans = sum_{i=0}^{qs-1}(i*a+rs)/b
ul ad = a/b*sum2(qs)+rs/b*qs; a %= b, rs %= b;
return ad+(a?triSum(b,a,a*qs+rs):0); // reduce if a >= b
} // then swap x and y axes and recurse
```

```
// \return sum_{x=0}^{n-1} (a*x+b)/m
// = |{(x,y) | 0 < m*y <= a*x+b < a*n+b}|
// assuming a*n+b does not overflow
ul divSum(ul n, ul a, ul b, ul m) { assert(m > 0);
ul extra = b/m*n; b %= m;
return extra+(a?triSum(m,a,a*n+b):0); }
// \return sum_{x=0}^{n-1} (a*x+b) %m
ul modSum(ul n, ll a, ll b, ul m) { assert(m > 0);
a = (a%m+m)%m, b = (b%m+m)%m;
return a*sum2(n)+b*n-m*divSum(n,a,b,m); }
```

### 4.2 Primality

### **4.2.1** Primes

p=962592769 is such that  $2^{21} \mid p-1$ , which may be useful. For hashing use 970592641 (31-bit number), 31443539979727 (45-bit), 3006703054056749 (52-bit). There are 78498 primes less than  $1\,000\,000$ .

Primitive roots exist modulo any prime power  $p^a$ , except for p=2, a>2, and there are  $\phi(\phi(p^a))$  many. For p=2, a>2, the group  $\mathbb{Z}_{2^a}^{\times}$  is instead isomorphic to  $\mathbb{Z}_2 \times \mathbb{Z}_{2^{a-2}}$ .

### 4.2.2 Divisors

```
\sum_{d|n} d = O(n \log \log n).
```

The number of divisors of n is at most around 100 for n < 5e4, 500 for n < 1e7, 2000 for n < 1e10, 200 000 for n < 1e19.

**Dirichlet Convolution:** Given a function f(x), let

$$(f * g)(x) = \sum_{d|x} g(d)f(x/d).$$

If the partial sums  $s_{f*g}(n)$ ,  $s_g(n)$  can be computed in O(1) and  $s_f(1...n^{2/3})$  can be computed in  $O\left(n^{2/3}\right)$  then all  $s_f\left(\frac{n}{d}\right)$  can as well. Use

$$s_{f*g}(n) = \sum_{d=1}^{n} g(d)s_f(n/d).$$

```
If f(x) = \mu(x) then g(x) = 1, (f * g)(x) = (x == 1), and s_f(n) = 1 - \sum_{i=2}^n s_f(n/i).
```

```
If f(x) = \phi(x) then g(x) = 1, (f * g)(x) = x, and s_f(n) = \frac{n(n+1)}{2} - \sum_{i=2}^n s_f(n/i).
```

### Sieve.h

**Description:** Tests primality up to SZ. Runs faster if only odd indices are stored

```
Time: \mathcal{O}\left(SZ\log\log SZ\right) or \mathcal{O}\left(SZ\right)
```

41c6ed, 20 lines

```
template<int SZ> struct Sieve {
 bitset<SZ> is_prime; vi primes;
 Sieve() {
   is_prime.set(); is_prime[0] = is_prime[1] = 0;
   for (int i = 4; i < SZ; i += 2) is_prime[i] = 0;
   for (int i = 3; i*i < SZ; i += 2) if (is_prime[i])
     for (int j = i*i; j < SZ; j += i*2) is_prime[j] = 0;
   FOR(i,SZ) if (is_prime[i]) primes.pb(i);
 // int sp[SZ]{}; // smallest prime that divides
 // Sieve() { // above is faster
 // FOR(i,2,SZ) {
       if (sp[i] == 0) sp[i] = i, primes.pb(i);
       for (int p: primes) {
         if (p > sp[i] \mid | i*p >= SZ) break;
         sp[i*p] = p;
 // }
 // }
```

### MultiplicativePrefix.h

**Description:**  $\sum_{i=1}^{N} f(i)$  where  $f(i) = \prod \text{val}[e]$  for each  $p^e$  in the factorization of i. Must satisfy val[1] = 1. Generalizes to any multiplicative function with  $f(p) = p^{\text{fixed power}}$ .

#### PrimeCnt.h

**Description:** Counts number of primes up to N. Can also count sum of primes.

```
Time: \mathcal{O}\left(N^{3/4}/\log N\right), 60ms for N=10^{11}, 2.5s for N=10^{13}
11 count_primes(11 N) { // count_primes(1e13) == 346065536839
 if (N <= 1) return 0;
 int sq = (int)sqrt(N);
 vl big_ans((sq+1)/2), small_ans(sq+1);
  FOR(i, 1, sq+1) small_ans[i] = (i-1)/2;
 FOR(i, sz(big_ans)) big_ans[i] = (N/(2*i+1)-1)/2;
  vb skip(sq+1); int prime_cnt = 0;
  for (int p = 3; p \le sq; p += 2) if (!skip[p]) { // primes
    for (int j = p; j \le sq; j += 2*p) skip[j] = 1;
    FOR(j, min((11)sz(big_ans), (N/p/p+1)/2)) {
      11 \text{ prod} = (11)(2*j+1)*p;
      big_ans[j] -= (prod > sq ? small_ans[(double) N/prod]
             : big_ans[prod/2])-prime_cnt;
    for (int j = sq, q = sq/p; q >= p; --q) for (; j >= q*p; --j)
      small_ans[j] -= small_ans[q]-prime_cnt;
    ++prime_cnt;
  return big_ans[0]+1;
```

#### MillerRabin.h

**Description:** Deterministic primality test, works up to  $2^{64}$ . For larger numbers, extend A randomly.

```
"ModMulLL.h"
                                                        89df33, 11 lines
bool prime (ul n) { // not 11!
 if (n < 2 || n % 6 % 4 != 1) return n-2 < 2;
  ul A[] = {2, 325, 9375, 28178, 450775, 9780504, 1795265022},
      s = \underline{\quad} builtin_ctzll(n-1), d = n>>s;
  each(a,A) { // ^ count trailing zeroes
   ul p = modPow(a,d,n), i = s;
    while (p != 1 \&\& p != n-1 \&\& a%n \&\& i--) p = modMul(p,p,n);
    if (p != n-1 && i != s) return 0;
 return 1;
```

#### FactorFast.h

Description: Pollard-rho randomized factorization algorithm. Returns prime factors of a number, in arbitrary order (e.g. 2299 -> {11, 19, 11}).

**Time:**  $\mathcal{O}\left(N^{1/4}\right)$ , less for numbers with small factors

```
ul pollard(ul n) { // return some nontrivial factor of n
  auto f = [n](ul x) \{ return modMul(x, x, n) + 1; \};
  ul x = 0, y = 0, t = 30, prd = 2, i = 1, q;
  while (t++ % 40 || gcd(prd, n) == 1) {
    if (x == y) x = ++i, y = f(x);
   if ((q = modMul(prd, max(x,y) - min(x,y), n))) prd = q;
   x = f(x), y = f(f(y));
  return gcd(prd, n);
void factor_rec(ul n, map<ul,int>& cnt) {
  if (n == 1) return;
  if (prime(n)) { ++cnt[n]; return; }
  ul u = pollard(n);
  factor_rec(u,cnt), factor_rec(n/u,cnt);
```

### **Euclidean Algorithm**

#### FracInterval.h

**Description:** Given fractions a < b with non-negative numerators and denominators, finds fraction f with lowest denominator such that a < f < b. Should work with all numbers less than  $2^{62}$ 

```
pl bet(pl a, pl b) {
  11 num = a.f/a.s; a.f = num*a.s, b.f = num*b.s;
  if (b.f > b.s) return {1+num,1};
  auto x = bet(\{b.s, b.f\}, \{a.s, a.f\});
  return {x.s+num*x.f,x.f};
```

#### Euclid.h

Description: Generalized Euclidean algorithm. euclid and invGeneral work for  $A, B < 2^{62}$ .

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

```
c7e528, 9 lines
// ceil(a/b)
// 11 cdiv(11 a, 11 b) { return a/b+((a^b)>0&&a%b); }
pl euclid(ll A, ll B) { // For A, B>=0, finds (x,y) s.t.
  // Ax+By=gcd(A,B), |Ax|, |By| \le AB/gcd(A,B)
  if (!B) return {1,0};
  pl p = euclid(B,A%B); return {p.s,p.f-A/B*p.s}; }
11 invGeneral(11 A, 11 B) { // find x in [0,B) such that Ax=1
  pl p = euclid(A,B); assert(p.f*A+p.s*B == 1);
```

```
return p.f+(p.f<0)*B; } // must have qcd(A,B)=1
```

### CRT.h

**Description:** Chinese Remainder Theorem.  $a.f \pmod{a.s}, b.f \pmod{b.s}$  $\implies$ ? (mod lcm(a.s, b.s)). Should work for  $ab < 2^{62}$ 

```
pl CRT(pl a, pl b) { assert(0 <= a.f && a.f < a.s && 0 <= b.f
  \hookrightarrow & & b.f < b.s);
 if (a.s < b.s) swap(a,b); // will overflow if b.s^2 > 2^{62}
 11 x,y; tie(x,y) = euclid(a.s,b.s);
 11 q = a.s*x+b.s*y, l = a.s/q*b.s;
 if ((b.f-a.f)%g) return {-1,-1}; // no solution
 // ?*a.s+a.f \equiv b.f \pmod{b.s}
 // ?= (b.f-a.f)/g*(a.s/g)^{-1} \pmod{b.s/g}
 x = (b.f-a.f) %b.s*x%b.s/q*a.s+a.f;
 return \{x+(x<0)*1,1\};
```

### ModArith.h

Description: Statistics on mod'ed arithmetic series. minBetween and minRemainder both assume that  $0 \le L \le R < B, \, AB < 2^{62}$ 

```
ll minBetween(ll A, ll B, ll L, ll R) {
 // min x s.t. exists y s.t. L \le A*x-B*y \le R
 A %= B;
 if (L == 0) return 0;
 if (A == 0) return -1;
 ll k = cdiv(L,A); if (A*k \le R) return k;
 11 x = minBetween (B, A, A-R%A, A-L%A); // min x s.t. exists y
 // s.t. -R <= Bx-Av <= -L
 return x == -1 ? x : cdiv(B*x+L,A); // solve for y
// find min((Ax+C)%B) for 0 <= x <= M
// aka find minimum non-negative value of A*x-B*y+C
// where 0 <= x <= M, 0 <= y
11 minRemainder(11 A, 11 B, 11 C, 11 M) {
 assert (A >= 0 && B > 0 && C >= 0 && M >= 0);
 A %= B, C %= B; ckmin(M,B-1);
 if (A == 0) return C;
 if (C >= A) { // make sure C < A
   11 \text{ ad} = \text{cdiv}(B-C,A);
   M \rightarrow ad; if (M < 0) return C;
    C += ad*A-B;
 11 q = B/A, new_B = B%A; // new_B < A
 if (new_B == 0) return C; // B-q*A
 // now minimize A*x-new_B*y+C
 // where \theta \le x, y and x+q*y \le M, \theta \le C < new_B < A
 // q*y -> C-new_B*y
 if (C/new_B > M/q) return C-M/q*new_B;
 M -= C/new_B*q; C %= new_B; // now C < new_B</pre>
 // given y, we can compute x = ceil[((B-q*A)*y-C)/A]
 // so x+q*y = ceil((B*y-C)/A) <= M
 11 \text{ max}_Y = (M*A+C)/B; // \text{ must have } y \leq \text{max}_Y
 ll\ max_X = cdiv(new_B*max_Y-C,A); // must\ have\ x <= max_X
 if (\max_X \times A - \text{new}_B \times \max_Y + C >= \text{new}_B) - -\max_X;
 // now we can remove upper bound on y
 return minRemainder(A, new_B, C, max_X);
```

### 4.4 Pythagorean Triples

The Pythagorean triples are uniquely generated by

$$a = k \cdot (m^2 - n^2), \ b = k \cdot (2mn), \ c = k \cdot (m^2 + n^2),$$

with m > n > 0, k > 0,  $m \perp n$ , and either m or n even.

### Lifting the Exponent

For n > 0, p prime, and ints x, y s.t.  $p \nmid x, y$  and p|x-y:

• 
$$p \neq 2$$
 or  $p = 2, 4|x - y$   
 $\implies v_p(x^n - y^n) = v_p(x - y) + v_p(n).$ 

• 
$$p = 2, 2|n \implies v_2(x^n - y^n) = v_2((x^2)^{n/2} - (y^2)^{n/2}).$$

# Combinatorial (5)

### Permutations

### **5.1.1** Cycles

Let  $q_S(n)$  be the number of n-permutations whose cycle lengths all belong to the set S. Then

$$\sum_{n=0}^{\infty} g_S(n) \frac{x^n}{n!} = \exp\left(\sum_{n \in S} \frac{x^n}{n}\right)$$

### 5.1.2 Burnside's lemma

Given a group G of symmetries and a set X, the number of elements of X up to symmetry equals

$$\frac{1}{|G|} \sum_{g \in G} |X^g|,$$

where  $X^g$  are the elements fixed by q (q.x = x).

If f(n) counts "configurations" (of some sort) of length n, we can ignore rotational symmetry using  $G = \mathbb{Z}_n$  to get

$$g(n) = \frac{1}{n} \sum_{k=0}^{n-1} f(\gcd(n,k)) = \frac{1}{n} \sum_{k|n} f(k)\phi(n/k).$$

### 5.2 Partitions and subsets

### 5.2.1 Partition function

Number of ways of writing n as a sum of positive integers, disregarding the order of the summands.

$$p(0) = 1, \ p(n) = \sum_{k \in \mathbb{Z} \setminus \{0\}} (-1)^{k+1} p(n - k(3k - 1)/2)$$

$$p(n) \sim 0.145/n \cdot \exp(2.56\sqrt{n})$$

### 5.2.2 Lucas' Theorem

Let n, m be non-negative integers and p a prime. Write  $n = n_k p^k + ... + n_1 p + n_0$  and  $m = m_k p^k + ... + m_1 p + m_0$ . Then  $\binom{n}{m} \equiv \prod_{i=0}^k \binom{n_i}{m_i} \pmod{p}$ .

### 5.3 General purpose numbers

### 5.3.1 Bernoulli numbers

EGF of Bernoulli numbers is  $B(t) = \frac{t}{e^t - 1}$  (FFT-able).  $B[0, \ldots] = [1, -\frac{1}{2}, \frac{1}{6}, 0, -\frac{1}{30}, 0, \frac{1}{42}, \ldots]$ 

Sums of powers:

$$\sum_{i=1}^{n} n^{m} = \frac{1}{m+1} \sum_{k=0}^{m} {m+1 \choose k} B_{k} (n+1)^{m+1-k}$$

Euler-Maclaurin formula for infinite sums:

$$\sum_{i=m}^{\infty} f(i) = \int_{m}^{\infty} f(x)dx - \sum_{k=1}^{\infty} \frac{B_k}{k!} f^{(k-1)}(m)$$

$$\approx \int_{-\infty}^{\infty} f(x)dx + \frac{f(m)}{2} - \frac{f'(m)}{12} + \frac{f'''(m)}{720} + O(f^{(5)}(m))$$

### 5.3.2 Stirling numbers of the first kind

Number of permutations on n items with k cycles.

$$c(n,k) = c(n-1,k-1) + (n-1)c(n-1,k), \ c(0,0) = 1$$
  
$$\sum_{k=0}^{n} c(n,k)x^{k} = x(x+1)\dots(x+n-1)$$

$$c(8, k) = 8, 0, 5040, 13068, 13132, 6769, 1960, 322, 28, 1$$
  
 $c(n, 2) = 0, 0, 1, 3, 11, 50, 274, 1764, 13068, 109584, \dots$ 

### 5.3.3 Eulerian numbers

Number of permutations  $\pi \in S_n$  in which exactly k elements are greater than the previous element. k j:s s.t.  $\pi(j) > \pi(j+1)$ , k+1 j:s s.t.  $\pi(j) \geq j$ , k j:s s.t.  $\pi(j) > j$ .

$$E(n,k) = (n-k)E(n-1,k-1) + (k+1)E(n-1,k)$$

$$E(n,0) = E(n,n-1) = 1$$

$$E(n,k) = \sum_{j=0}^{k} (-1)^{j} \binom{n+1}{j} (k+1-j)^{n}$$

### 5.3.4 Stirling numbers of the second kind

Partitions of n distinct elements into exactly k groups.

$$S(n,k) = S(n-1,k-1) + kS(n-1,k)$$

$$S(n,1) = S(n,n) = 1$$

$$S(n,k) = \frac{1}{k!} \sum_{j=0}^{k} (-1)^{k-j} \binom{k}{j} j^{n}$$

### 5.3.5 Bell numbers

Total number of partitions of n distinct elements. B(n) = 1, 1, 2, 5, 15, 52, 203, 877, 4140, 21147, .... For p prime,

$$B(p^m + n) \equiv mB(n) + B(n+1) \pmod{p}$$

### 5.3.6 Labeled unrooted trees

# on n vertices:  $n^{n-2}$ # on k existing trees of size  $n_i$ :  $n_1 n_2 \cdots n_k n^{k-2}$ # with degrees  $d_i$ :  $(n-2)!/((d_1-1)!\cdots(d_n-1)!)$ 

### 5.3.7 Catalan numbers

$$C_n = \frac{1}{n+1} {2n \choose n} = {2n \choose n} - {2n \choose n+1} = \frac{(2n)!}{(n+1)!n!}$$

$$C_0 = 1, \ C_{n+1} = \frac{2(2n+1)}{n+2} C_n, \ C_{n+1} = \sum_{i=1}^{n} C_i C_{n-i}$$

$$C_n = 1, 1, 2, 5, 14, 42, 132, 429, 1430, 4862, 16796, 58786, \dots$$

• sub-diagonal monotone paths in an  $n \times n$  grid.

- strings with n pairs of parenthesis, correctly nested.
- binary trees with with n+1 leaves (0 or 2 children).
- ordered trees with n+1 vertices.
- ways a convex polygon with n+2 sides can be cut into triangles by connecting vertices with straight lines.
- $\bullet$  permutations of [n] with no 3-term increasing subseq.

### 5.4 Young Tableaux

Let a **Young diagram** have shape  $\lambda = (\lambda_1 \ge \cdots \ge \lambda_k)$ , where  $\lambda_i$  equals the number of cells in the *i*-th (left-justified) row from the top. A **Young tableau** of shape  $\lambda$  is a filling of the  $n = \sum \lambda_i$  cells with a permutation of  $1 \dots n$  such that each row and column is increasing.

**Hook-Length Formula**: For the cell in position (i, j), let  $h_{\lambda}(i, j) = |\{(I, J)| i \leq I, j \leq J, (I = i \text{ or } J = j)\}|$ . The number of Young tableaux of shape  $\lambda$  is equal to  $f^{\lambda} = \frac{n!}{\prod h_{\lambda}(i, j)}$ .

Schensted's Algorithm: converts a permutation  $\sigma$  of length n into a pair of Young Tableaux  $(S(\sigma), T(\sigma))$  of the same shape. When inserting  $x = \sigma_i$ ,

- 1. Add x to the first row of S by inserting x in place of the largest y with x < y. If y doesn't exist, push x to the end of the row, set the value of T at that position to be i, and stop.
- 2. Add y to the second row using the same rule, keep repeating as necessary.

All pairs  $(S(\sigma), T(\sigma))$  of the same shape correspond to a unique  $\sigma$ , so  $n! = \sum (f^{\lambda})^2$ . Also,  $S(\sigma^R) = S(\sigma)^T$ .

Let  $d_k(\sigma), a_k(\sigma)$  be the lengths of the longest subseqs which are a union of k decreasing/ascending subseqs, respectively. Then  $a_k(\sigma) = \sum_{i=1}^k \lambda_i, d_k(\sigma) = \sum_{i=1}^k \lambda_i^*$ , where  $\lambda_i^*$  is size of the i-th column.

### 5.5 Other

### DeBruijnSeq.h

**Description:** Given alphabet [0, k) constructs a cyclic string of length  $k^n$  that contains every length n string as substr.

a6961b, 13 lines

```
vi deBruijnSeq(int k, int n) {
   if (k == 1) return {0};
   vi seq, aux(n+1);
   function<void(int,int)> gen = [&](int t, int p) {
      if (t > n) { // +lyndon word of len p
         if (n%p == 0) FOR(i,1,p+1) seq.pb(aux[i]);
   } else {
      aux[t] = aux[t-p]; gen(t+1,p);
```

b18e29, 21 lines

### NimProduct MatroidIsect Matrix MatrixInv

```
while (++aux[t] < k) gen(t+1,t);
    };
    gen(1,1); return seq;
}</pre>
```

### NimProduct.h

**Description:** Product of nimbers is associative, commutative, and distributive over addition (xor). Forms finite field of size  $2^{2^k}$ . Defined by  $ab = \max(\{a'b + ab' + a'b' : a' < a, b' < b\})$ . Application: Given 1D coin turning games  $G_1, G_2$   $G_1 \times G_2$  is the 2D coin turning game defined as follows. If turning coins at  $x_1, x_2, \ldots, x_m$  is legal in  $G_1$  and  $y_1, y_2, \ldots, y_n$  is legal in  $G_2$ , then turning coins at all positions  $(x_i, y_j)$  is legal assuming that the coin at  $(x_m, y_n)$  goes from heads to tails. Then the grundy function g(x, y) of  $G_1 \times G_2$  is  $g_1(x) \times g_2(y)$ .

**Time:** 64<sup>2</sup> xors per multiplication, memorize to speed up. 5afe17, 46 lines

```
using ul = uint64_t;
struct Precalc {
  ul tmp[64][64], y[8][8][256];
  unsigned char x[256][256];
  Precalc() { // small nim products, all < 256
   FOR(i, 256) FOR(j, 256) x[i][j] = mult < 8 > (i, j);
   FOR(i,8) FOR(j,i+1) FOR(k,256)
     y[i][j][k] = mult < 64 > (prod2(8*i, 8*j), k);
  ul prod2(int i, int j) { // nim prod of 2^i, 2^j
   ul& u = tmp[i][j]; if (u) return u;
   if (!(i&j)) return u = 1ULL<<(i|j);
    int a = (i\&j)\&-(i\&j); // a=2^k, consider 2^{2^k}
    return u=prod2(i^a, j)^prod2((i^a)|(a-1),(j^a)|(i&(a-1)));
    \frac{1}{2^{2^{i}}} *2^{2^{i}} = 2^{2^{i}} = 2^{2^{i}} if i < i
  template<int L> ul mult(ul a, ul b) {
    ul c = 0; FOR(i, L) if (a>>i&1)
     FOR(j,L) if (b>>j&1) c ^= prod2(i,j);
  // 2^{6*i}*(a>>(8*i)&255) * 2^{6*i}*(b>>(8*i)&255)
  // \rightarrow (2^{8*i}*2^{8*i})*((a>>(8*i)&255)*(b>>(8*i)&255))
  ul multFast(ul a, ul b) const { // faster nim product
    ul res = 0; auto f=[](ul c,int d) {return c>>(8*d)&255;};
     FOR(j,i) res ^= y[i][j][x[f(a,i)][f(b,j)]
             ^x[f(a,j)][f(b,i)]];
     res ^= y[i][i][x[f(a,i)][f(b,i)]];
   }
    return res;
};
const Precalc P;
struct nb { // nimber
 ul x; nb() { x = 0; }
  nb(ul _x): x(_x) {}
  explicit operator ul() { return x; }
  nb operator+(nb y) { return nb(x^y.x); }
  nb operator*(nb y) { return nb(P.multFast(x,y.x)); }
  friend nb pow(nb b, ul p) {
   nb res = 1; for (;p;p/=2,b=b*b) if (p&1) res = res*b;
   return res; } // b^{2^{2^{A}}-1}=1 where 2^{2^{A}} > b
  friend nb inv(nb b) { return pow(b,-2); }
};
```

#### MatroidIsect.h

**Description:** Computes a set of maximum size which is independent in both graphic and colorful matroids, aka a spanning forest where no two edges are of the same color. In general, construct the exchange graph and find a shortest path. Can apply similar concept to partition matroid.

Usage: MatroidIsect<Gmat, Cmat> M(sz (ed), Gmat (ed), Cmat (col)) Time:  $\mathcal{O}(GI^{1.5})$  calls to oracles, where G is size of ground set and I is size of G is size of G in G is size of G.

of independent set. "DSU.h" d0051c, 51 lines struct Gmat { // graphic matroid int V = 0; vpi ed; DSU D; Gmat(vpi ed):ed( ed) { map < int, int > m; each(t, ed) m[t.f] = m[t.s] = 0;each(t,m) t.s = V++;each(t,ed) t.f = m[t.f], t.s = m[t.s]; void clear() { D.init(V); } void ins(int i) { assert(D.unite(ed[i].f,ed[i].s)); } bool indep(int i) { return !D.sameSet(ed[i].f,ed[i].s); } struct Cmat { // colorful matroid int C = 0; vi col; V<bool> used; Cmat(vi col):col(col) {each(t,col) ckmax(C,t+1); } void clear() { used.assign(C,0); } void ins(int i) { used[col[i]] = 1; } bool indep(int i) { return !used[col[i]]; } template<class M1, class M2> struct MatroidIsect { int n; V<bool> iset; M1 m1; M2 m2; bool augment() { vi pre(n+1,-1); queue<int> q({n}); while (sz(q)) { int x = q.ft; q.pop();if (iset[x]) { m1.clear(); F0R(i,n) if (iset[i] && i != x) m1.ins(i); FOR(i,n) if (!iset[i] && pre[i] == -1 && ml.indep(i)) pre[i] = x, q.push(i);} else { auto backE = [&]() { // back edge m2.clear();  $FOR(c, 2) FOR(i, n) if((x==i | | iset[i]) &&(pre[i]==-1) == c) {$ if (!m2.indep(i))return c?pre[i]=x,q.push(i),i:-1; m2.ins(i); } return n; for (int y; (y = backE()) != -1;) if  $(y == n) {$ for(; x != n; x = pre[x]) iset[x] = !iset[x]; return 1: } } return 0: MatroidIsect(int n, M1 m1, M2 m2):n(n), m1(m1), m2(m2) { iset.assign(n+1,0); iset[n] = 1;ml.clear(); m2.clear(); // greedily add to basis R0F(i,n) if (m1.indep(i) && m2.indep(i)) iset[i] = 1, m1.ins(i), m2.ins(i);while (augment()); };

# Numerical (6)

### 6.1 Matrix

#### Matrix.h

**Description:** 2D matrix operations. "ModInt.h"

```
using T = mi;
using Mat = V<V<T>>; // use array instead if tight TL
Mat makeMat(int r, int c) { return Mat(r, V<T>(c)); }
Mat makeId(int n) {
 Mat m = makeMat(n,n); FOR(i,n) m[i][i] = 1;
 return m:
Mat operator*(const Mat& a, const Mat& b) {
 int x = sz(a), y = sz(a[0]), z = sz(b[0]);
 assert (y == sz(b)); Mat c = makeMat(x,z);
 FOR(i,x) FOR(j,y) FOR(k,z) c[i][k] += a[i][j]*b[j][k];
 return c:
Mat& operator*=(Mat& a, const Mat& b) { return a = a*b; }
Mat pow(Mat m, 11 p) {
 int n = sz(m); assert (n == sz(m[0]) \&\& p >= 0);
 Mat res = makeId(n);
 for (; p; p /= 2, m \star= m) if (p&1) res \star= m;
 return res;
```

#### MatrixInv.h

**Description:** Uses gaussian elimination to convert into reduced row echelon form and calculates determinant. For determinant via arbitrary modulos, use a modified form of the Euclidean algorithm because modular inverse may not exist. If you have computed  $A^{-1} \pmod{p^k}$ , then the inverse  $\pmod{p^{2k}}$  is  $A^{-1}(2I-AA^{-1})$ .

**Time:**  $\mathcal{O}(N^3)$ , determinant of  $1000 \times 1000$  matrix of modints in 1 second if you reduce # of operations by half

```
const db EPS = 1e-9; // adjust?
int getRow(V<V<db>>& m, int R, int i, int nex) {
    pair<db,int> bes{0,-1}; // find row with max abs value
    FOR(j,nex,R) ckmax(bes,{abs(m[j][i]),j});
    return bes.f < EPS ? -1 : bes.s; }
int getRow(V<vmi>& m, int R, int i, int nex) {
    FOR(j, nex, R) if (m[j][i] != 0) return j;
    return -1; }
pair<T,int> gauss(Mat& m) { // convert to reduced row echelon
   \hookrightarrow form
    if (!sz(m)) return {1,0};
    int R = sz(m), C = sz(m[0]), rank = 0, nex = 0;
    T prod = 1; // determinant
    F0R(i,C) {
        int row = getRow(m,R,i,nex);
        if (row == -1) { prod = 0; continue; }
        if (row != nex) prod \star = -1, swap(m[row], m[nex]);
        prod *= m[nex][i]; rank++;
        T x = 1/m[nex][i]; FOR(k,i,C) m[nex][k] *= x;
        FOR(j,R) if (j != nex) {
            T v = m[j][i]; if (v == 0) continue;
            FOR(k,i,C) m[j][k] \rightarrow v*m[nex][k];
        nex++;
    return {prod,rank};
Mat inv(Mat m) {
```

int R = sz(m); assert(R == sz(m[0]));

```
Mat x = makeMat(R,2*R);
FOR(i,R) {
    x[i][i+R] = 1;
    FOR(j,R) x[i][j] = m[i][j];
}
if (gauss(x).s != R) return Mat();
Mat res = makeMat(R,R);
FOR(i,R) FOR(j,R) res[i][j] = x[i][j+R];
return res;
```

#### MatrixTree.h

**Description:** Kirchhoff's Matrix Tree Theorem. Given adjacency matrix, calculates # of spanning trees.

```
"MatrixInv.h" 066e59, 11 lines
T numSpan(Mat m) {
   int n = sz(m); Mat res(n-1,n-1);
   FOR(i,n) FOR(j,i+1,n) {
      mi ed = m[i][j]; res[i][i] += ed;
      if (j != n-1) {
       res[j][j] += ed;
      res[i][j] -= ed, res[j][i] -= ed;
    }
   }
   return gauss(res).f;
}
```

#### ShermanMorrison.h

**Description:** Calculates  $(A + uv^T)^{-1}$  given  $B = A^{-1}$ . Not invertible if sum=0.

## 6.2 Polynomials

#### Poly.h

**Description:** Basic poly ops including division. Can replace T with double, complex.

```
"ModInt.h"
                                                     cd218a, 73 lines
using T = mi; using poly = V<T>;
void remz(poly& p) { while (sz(p)\&\&p.bk==T(0)) p.pop\_back(); }
poly REMZ(poly p) { remz(p); return p; }
poly rev(poly p) { reverse(all(p)); return p; }
poly shift (poly p, int x) {
  if (x \ge 0) p.insert(begin(p), x, 0);
  else assert(sz(p)+x \ge 0), p.erase(begin(p),begin(p)-x);
  return p;
poly RSZ(const poly& p, int x) {
  if (x \le sz(p)) return poly(begin(p), begin(p)+x);
  poly q = p; q.rsz(x); return q; }
T eval(const poly& p, T x) { // evaluate at point x
 T res = 0; ROF(i,sz(p)) res = x*res+p[i];
  return res; }
poly dif(const poly& p) { // differentiate
  poly res; FOR(i,1,sz(p)) res.pb(T(i)*p[i]);
  return res; }
poly integ(const poly& p) { // integrate
  static poly invs{0,1};
  for (int i = sz(invs); i \le sz(p); ++i)
```

```
invs.pb(-MOD/i*invs[MOD%i]);
  poly res(sz(p)+1); FOR(i,sz(p)) res[i+1] = p[i]*invs[i+1];
 return res;
poly& operator+=(poly& 1, const poly& r) {
 1.rsz(max(sz(1),sz(r))); FOR(i,sz(r)) 1[i] += r[i];
 return 1; }
poly& operator = (poly& 1, const poly& r) {
 1.rsz(max(sz(1),sz(r))); F0R(i,sz(r)) 1[i] -= r[i];
poly& operator *= (poly& 1, const T& r) { each(t,1) t *= r;
 return 1; }
poly& operator/=(poly& 1, const T& r) { each(t,1) t /= r;
 return 1; }
poly operator+(poly 1, const poly& r) { return 1 += r; }
poly operator-(poly 1, const poly& r) { return 1 -= r; }
poly operator-(poly 1) { each(t,1) t *= -1; return 1; }
poly operator*(poly 1, const T& r) { return 1 *= r; }
poly operator* (const T& r, const poly& 1) { return 1*r; }
poly operator/(poly 1, const T& r) { return 1 /= r; }
poly operator* (const poly& 1, const poly& r) {
 if (!min(sz(l),sz(r))) return {};
 polv x(sz(1)+sz(r)-1);
 FOR(i, sz(1)) FOR(j, sz(r)) x[i+j] += l[i]*r[j];
 return x:
poly& operator*=(poly& 1, const poly& r) { return 1 = 1*r; }
pair<poly, poly> quoRemSlow(poly a, poly b) {
  remz(a); remz(b); assert(sz(b));
 T lst = b.bk, B = T(1)/lst; each(t,a) t *= B;
  each (t,b) t *= B;
  poly q(max(sz(a)-sz(b)+1,0));
  for (int dif; (dif=sz(a)-sz(b)) >= 0; remz(a)) {
   q[dif] = a.bk; FOR(i,sz(b)) a[i+dif] -= q[dif]*b[i]; }
  each(t,a) t *= 1st;
 return {q,a}; // quotient, remainder
poly operator% (const poly& a, const poly& b) {
 return quoRemSlow(a,b).s; }
T resultant(poly a, poly b) { // R(A,B)
 // =b_m^n*prod_{j=1}^mA(mu_j)
  // =b_m^na_n^m*prod_{i=1}^nprod_{j=1}^m (mu_j-lambda_i)
  // = (-1) ^{mn}a_n^m*prod_{i=1}^nB(lambda_i)
  // = (-1) ^{nm}R(B, A)
  // Also, R(A,B)=b_m^{deg(A)-deg(A-CB)}R(A-CB,B)
  int ad = sz(a)-1, bd = sz(b)-1;
  if (bd <= 0) return bd < 0 ? 0 : pow(b.bk,ad);
  int pw = ad; a = a\%b; pw -= (ad = sz(a)-1);
  return resultant(b,a) *pow(b.bk,pw) *T((bd&ad&1)?-1:1);
```

### PolyInterpolate.h

**Description:** n points determine unique polynomial of degree  $\leq n-1$ . For numerical precision pick  $v[k] \cdot f = c * \cos(k/(n-1) * \pi), k = 0 \dots n-1$ . **Time:**  $\mathcal{O}(n^2)$ 

#### FFT.h

**Description:** Multiply polynomials of ints for any modulus  $< 2^{31}$ . For XOR convolution ignore m within fft.

**Time:**  $\mathcal{O}(N \log N)$ . For  $N = 10^6$ , conv is  $\sim 0.13$ ms, conv\_general is  $\sim 320$ ms. "ModInt.h" 19ab20, 39 lines // const int MOD = 998244353; tcT> void fft(V<T>& A, bool invert = 0) { // NTT int n = sz(A); assert((T::mod-1)%n == 0); V<T> B(n); for (int b = n/2; b; b /= 2, swap (A,B)) { // w = n/b'th root T w = pow(T::rt(), (T::mod-1)/n\*b), m = 1;for (int i = 0; i < n; i += b\*2, m \*= w) FOR (i,b) { T u = A[i+j], v = A[i+j+b]\*m;B[i/2+j] = u+v; B[i/2+j+n/2] = u-v;if (invert) { reverse(1+all(A));  $Tz = inv(T(n)); each(t,A) t *= z; }$ } // for NTT-able moduli tcT> V<T> conv(V<T> A, V<T> B) { if (!min(sz(A),sz(B))) return {}; int s = sz(A) + sz(B) - 1, n = 1; for (; n < s; n \*= 2); A.rsz(n), fft(A); B.rsz(n), fft(B);  $FOR(i,n) A[i] \star= B[i];$ fft(A,1); A.rsz(s); return A; template<class M, class T> V<M> mulMod(const V<T>& x, const V<T auto con = [](const V<T>& v) { V<M> w(sz(v)); FOR(i,sz(v)) w[i] = (int)v[i];return w; };

tcT> V<T> conv general (const V<T>& A, const V<T>& B) {

m2 r02 = inv(m2(m0::mod)), r12 = inv(m2(m1::mod));

int a = c0[i].v, b = ((c1[i]-a)\*r01).v,

c = (((c2[i]-a)\*r02-b)\*r12).v;

using m0 = mint<(119<<23)+1,62>; auto c0 = mulMod<m0>(A,B);

using m1 = mint < (5 << 25) +1, 62>; auto c1 = mulMod < m1 > (A, B);

using m2 = mint < (7 << 26) +1, 62>; auto c2 = mulMod < m2> (A, B);

FOR(i,n) { // a=remainder mod m0::mod, b fixes it mod m1::mod

res[i] = (T(c)\*m1::mod+b)\*m0::mod+a; // c fixes m2::mod

int  $n = sz(c\theta)$ ; V < T > res(n);  $m1 r\theta1 = inv(m1(m\theta::mod))$ ;

### PolyInvSimpler.h

return res;

return conv(con(x), con(y));

} // arbitrary moduli

**Description:** computes  $A^{-1}$  such that  $AA^{-1} \equiv 1 \pmod{x^n}$ . Newton's method: If you want F(x) = 0 and  $F(Q_k) \equiv 0 \pmod{x^a}$  then  $Q_{k+1} = Q_k - \frac{F(Q_k)}{F(Q_k)} \pmod{x^{2a}}$  satisfies  $F(Q_{k+1}) \equiv 0 \pmod{x^{2a}}$ . Application: if f(n), g(n) are the #s of forests and trees on n nodes then  $\sum_{n=0}^{\infty} f(n)x^n = \exp\left(\sum_{n=1}^{\infty} \frac{g(n)}{n!}\right)$ .

**Usage:** vmi v{1,5,2,3,4}; ps(exp(2\*log(v,9),9)); // squares v **Time:**  $\mathcal{O}(N \log N)$ . For  $N = 5 \cdot 10^5$ , inv~270ms, log ~350ms, exp~550ms "FFT.h", "Poly.h" 6e5362, 30 line

```
poly inv(poly A, int n) { // Q-(1/Q-A)/(-Q^{-2})}
poly B{inv(A[0])};
for (int x = 2; x/2 < n; x *= 2)
    B = 2*B-RSZ(conv(RSZ(A,x),conv(B,B)),x);
return RSZ(B,n);
}
poly sqrt(const poly& A, int n) { // Q-(Q^2-A)/(2Q)}
assert(A[0].v == 1); poly B{1};
for (int x = 2; x/2 < n; x *= 2)
    B = inv(T(2))*RSZ(B+conv(RSZ(A,x),inv(B,x)),x);
return RSZ(B,n);</pre>
```

```
// return {quotient, remainder}
pair<poly, poly> quoRem(const poly& f, const poly& g) {
 if (sz(f) < sz(g)) return {{},f};
  poly q = conv(inv(rev(g), sz(f) - sz(g) + 1), rev(f));
  q = rev(RSZ(q, sz(f) - sz(q) + 1));
  poly r = RSZ(f-conv(q,q),sz(q)-1); return \{q,r\};
poly log(poly A, int n) { assert(A[0].v == 1); // (ln A)' = A'/
  \hookrightarrow A
  A.rsz(n); return integ(RSZ(conv(dif(A),inv(A,n-1)),n-1)); }
poly exp(poly A, int n) { assert(A[0].v == 0);
  poly B{1}, IB{1}; // inverse of B
  for (int x = 1; x < n; x *= 2) {
    IB = 2 * IB - RSZ (conv(B, conv(IB, IB)), x);
    poly Q = dif(RSZ(A,x)); Q += RSZ(conv(IB, dif(B) - conv(B,Q))
       \hookrightarrow, 2*x-1);
    B = B+RSZ (conv(B,RSZ(A,2*x)-integ(Q)),2*x);
  return RSZ(B,n);
```

### 6.3 Misc

#### LinearRecurrence.h

```
Description: Berlekamp-Massey. Computes linear recurrence C of order N for sequence s of 2N terms. C[0] = 1 and for all i \geq sz(C) - 1, \sum_{j=0}^{sz(C)-1} C[j]s[i-j] = 0. Usage: LinRec L; L.init(\{0,1,1,2,3\}); L.eval(5); L.eval(6); // 5, 8

Time: init \Rightarrow \mathcal{O}(N|C|), eval \Rightarrow \mathcal{O}(|C|^2 \log p) or faster with FFT
```

39ea71, 29 lines "Poly.h" struct LinRec { poly s, C, rC; void BM() { int x = 0; T b = 1; poly B;  $B = C = \{1\}$ ; // B is fail vector FOR(i, sz(s)) { // update C after adding a term of s ++x; int L = sz(C), M = i+3-L; T d = 0; FOR(j,L) d += C[j]\*s[i-j];  $// [D^i]C*s$ if (d.v == 0) continue;  $// [D^i]C*s=0$ poly  $_{C} = C; T coef = d*inv(b);$ C.rsz(max(L,M)); F0R(j,sz(B)) C[j+x] -= coef\*B[j];if  $(L < M) B = _C, b = d, x = 0;$ void init(const poly& \_s) {  $s = _s; BM();$ rC = C; reverse(all(rC)); C.erase(begin(C)); each(t,C) t \*=-1; poly getPow(ll p) { // get x^p mod rC if (p == 0) return {1}; poly r = getPow(p/2); r = (r\*r) %rC; return p&1?(r\*poly{0,1})%rC:r; T dot(poly v) { // dot product with s T ans = 0; FOR(i,sz(v)) ans +=v[i]\*s[i];return ans; } // get p-th term of rec T eval(ll p) { assert(p >= 0); return dot(getPow(p)); }

### Integrate.h

**Description:** Integration of a function over an interval using Simpson's rule, exact for polynomials of degree up to 3. The error should be proportional to  $dif^4$ , although in practice you will want to verify that the result is stable to desired precision when epsilon changes.

```
Usage: quad([](db x) { return x*x+3*x+1; }, 2, 3) // 34.833.6 hies

template<class F> db quad(F f, db a, db b) {
  const int n = 1000;
  db dif = (b-a)/2/n, tot = f(a)+f(b);
  FOR(i,1,2*n) tot += f(a+i*dif)*(i&1?4:2);
  return tot*dif/3;
}
```

### IntegrateAdaptive.h

**Description:** Unused. Fast integration using adaptive Simpson's rule, exact for polynomials of degree up to 5. **Usage:** db z, y;

```
db h(db x) { return x*x + y*y + z*z <= 1; }
db g(db y) { ::y = y; return quad(h, -1, 1); }
db f(db z) { ::z = z; return quad(g, -1, 1); }
db sphereVol = quad(f, -1, 1), pi = sphereVol*3/4;

template<class F> db simpson(F f, db a, db b) {
    db c = (a+b)/2; return (f(a)+4*f(c)+f(b))*(b-a)/6; }

template<class F> db rec(F& f, db a, db b, db eps, db S) {
    db c = (a+b)/2;
    db S1 = simpson(f, a, c), S2 = simpson(f, c, b), T = S1+S2;
    if (abs(T-S)<=15*eps || b-a<1e-10) return T+(T-S)/15;
    return rec(f, a, c, eps/2, S1)+rec(f, c, b, eps/2, S2);
}

template<class F> db quad(F f, db a, db b, db eps = 1e-8) {
    return rec(f, a, b, eps, simpson(f, a, b)); }
```

### Simplex.h

**Description:** Solves a general linear maximization problem: maximize  $c^Tx$  subject to  $Ax \leq b, x \geq 0$ . Returns -inf if there is no solution, inf if there are arbitrarily good solutions, or the maximum value of  $c^Tx$  otherwise. The input vector is set to an optimal x (or in the unbounded case, an arbitrary solution fulfilling the constraints). Numerical stability is not guaranteed. For better performance, define variables such that x=0 is viable.

```
Usage: vvd A\{\{1,-1\}, \{-1,1\}, \{-1,-2\}\}; vd b\{1,1,-4\}, c\{-1,-1\}, x; T val = LPSolver(A, b, c).solve(x);
```

**Time:**  $\mathcal{O}(NM \cdot \#pivots)$ , where a pivot may be e.g. an edge relaxation.  $\mathcal{O}(2^N)$  in the general case.

```
c99f9c, 67 lines
using T = db; // double probably suffices
using vd = V<T>; using vvd = V<vd>;
const T eps = 1e-8, inf = 1/.0;
#define ltj(X) if (s=-1 \mid | mp(X[j],N[j]) < mp(X[s],N[s])) s=j
struct LPSolver {
 int m, n; // # m = contraints, # n = variables
 vi N, B; // N[j] = non-basic variable (j-th column), = 0
  vvd D; // B[j] = basic variable (j-th row)
 LPSolver(const vvd& A, const vd& b, const vd& c) :
    m(sz(b)), n(sz(c)), N(n+1), B(m), D(m+2), vd(n+2)) {
    FOR(i,m) FOR(j,n) D[i][j] = A[i][j];
    FOR(i,m) B[i] = n+i, D[i][n] = -1, D[i][n+1] = b[i];
    // B[i]: basic variable for each constraint
    // D[i][n]: artificial variable for testing feasibility
    FOR(j,n) N[j] = j, D[m][j] = -c[j];
    // D[m] stores negation of objective,
    // which we want to minimize
    N[n] = -1; D[m+1][n] = 1; // to find initial feasible
  } // solution, minimize artificial variable
  void pivot(int r, int s) { // swap B[r] (row)
   T inv = 1/D[r][s]; // with N[r] (column)
```

```
FOR(i, m+2) if (i != r && abs(D[i][s]) > eps) {
    T binv = D[i][s]*inv;
    FOR(j, n+2) if (j != s) D[i][j] -= D[r][j]*binv;
    D[i][s] = -binv;
  D[r][s] = 1; FOR(j, n+2) D[r][j] *= inv; // scale r-th row
  swap(B[r],N[s]);
bool simplex(int phase) {
  int x = m + phase - 1;
  while (1) { // if phase=1, ignore artificial variable
    int s = -1; FOR(j, n+1) if (N[j] != -phase) ltj(D[x]);
    // find most negative col for nonbasic (NB) variable
    if (D[x][s] >= -eps) return 1;
    // can't get better sol by increasing NB variable
    int r = -1;
    F0R(i,m) {
      if (D[i][s] <= eps) continue;</pre>
      if (r == -1 \mid | mp(D[i][n+1] / D[i][s], B[i])
             < mp(D[r][n+1] / D[r][s], B[r])) r = i;
      // find smallest positive ratio
    } // -> max increase in NB variable
    if (r == -1) return 0; // objective is unbounded
    pivot(r,s);
T solve (vd& x) { // 1. check if x=0 feasible
  int r = 0; FOR(i,1,m) if (D[i][n+1] < D[r][n+1]) r = i;
  if (D[r][n+1] < -eps) { // if not, find feasible start
    pivot(r,n); // make artificial variable basic
    assert(simplex(2)); // I think this will always be true??
    if (D[m+1][n+1] < -eps) return -inf;</pre>
    // D[m+1][n+1] is max possible value of the negation of
    // artificial variable, optimal value should be zero
    // if exists feasible solution
    FOR(i,m) if (B[i] == -1) { // artificial var basic
      int s = 0; FOR(j,1,n+1) Itj(D[i]); // -> nonbasic
      pivot(i,s);
  bool ok = simplex(1); x = vd(n);
  FOR(i,m) if (B[i] < n) x[B[i]] = D[i][n+1];
  return ok ? D[m][n+1] : inf;
```

# Graphs (7)

**Erdos-Gallai:**  $d_1 \ge \cdots \ge d_n$  can be degree sequence of simple graph on n vertices iff their sum is even and  $\sum_{i=1}^k d_i \le k(k-1) + \sum_{i=k+1}^n \min(d_i, k), \forall 1 \le k \le n.$ 

### 7.1 Basics

#### DSU.h

**Description:** Disjoint Set Union with path compression and union by size. Add edges and test connectivity. Use for Kruskal's or Boruvka's minimum spanning tree. Time:  $\mathcal{O}(\alpha(N))$ 

e42a83, 11 lines

```
struct DSU {
  vi e; void init(int N) { e = vi(N,-1); }
  int get(int x) { return e[x] < 0 ? x : e[x] = get(e[x]); }
  bool sameSet(int a, int b) { return get(a) == get(b); }
  int size(int x) { return -e[get(x)]; }
  bool unite(int x, int y) { // union by size</pre>
```

### NegativeCycle LCAjump LCArmq HLD Centroid

vi depth, pos, par, rev; // rev is for compress

void init(int \_N) { N = \_N; adj.rsz(N);

depth = pos = par = rev = vi(N); }

vpi tmp; RMQ<pi> r;

int lca(int x, int y) {

template <class BinaryOp>

for (; root[x] != root[y]; y = par[root[y]])

return depth[x] < depth[y] ? x : y;</pre>

LazySeg<11,SZ> tree; // segtree for sum

if (depth[x] > depth[y]) swap(x,y);

op(pos[root[y]],pos[y]); }

void processPath(int x, int y, BinaryOp op) {

for (; root[x] != root[y]; y = par[root[y]]) {

if (depth[root[x]] > depth[root[y]]) swap(x,y);

if (depth[root[x]] > depth[root[y]]) swap(x,y);

```
x = get(x), y = get(y); if (x == y) return 0;
   if (e[x] > e[y]) swap(x,y);
   e[x] += e[y]; e[y] = x; return 1;
};
```

### NegativeCvcle.h

**Description:** use Bellman-Ford (make sure no underflow)

688ec8, 11 lines

```
vi negCyc(int N, V<pair<pi,int>> ed) {
  vl d(N); vi p(N); int x = -1;
  rep(N) {
   x = -1; each(t,ed) if (ckmin(d[t.f.s],d[t.f.f]+t.s))
     p[t.f.s] = t.f.f, x = t.f.s;
    if (x == -1) return \{\};
  rep(N) x = p[x]; // enter cycle
  vi cyc{x}; while (p[cyc.bk] != x) cyc.pb(p[cyc.bk]);
  reverse(all(cvc)); return cvc;
```

### Trees

### LCAjump.h

**Description:** Calculates least common ancestor in tree with verts  $0 \dots N-1$ and root R using binary jumping.

**Memory:**  $\mathcal{O}(N \log N)$ 

**Time:**  $\mathcal{O}(N \log N)$  build,  $\mathcal{O}(\log N)$  query

```
6b0ee9, 28 lines
struct LCA {
  int N; V<vi> par, adj; vi depth;
  void init(int _N) { N = _N;
    int d = 1; while ((1 << d) < N) ++d;
   par.assign(d, vi(N)); adj.rsz(N); depth.rsz(N);
  void ae(int x, int y) { adj[x].pb(y), adj[y].pb(x); }
  void gen(int R = 0) \{ par[0][R] = R; dfs(R); \}
  void dfs(int x = 0) {
   FOR(i, 1, sz(par)) par[i][x] = par[i-1][par[i-1][x]];
    each(y,adj[x]) if (y != par[0][x])
      depth[y] = depth[par[0][y]=x]+1, dfs(y);
  int jmp(int x, int d) {
   FOR(i, sz(par)) if ((d>>i)&1) x = par[i][x];
    return x; }
  int lca(int x, int v) {
    if (depth[x] < depth[y]) swap(x,y);</pre>
    x = jmp(x, depth[x]-depth[y]); if (x == y) return x;
    R0F(i,sz(par)) {
     int X = par[i][x], Y = par[i][y];
     if (X != Y) x = X, y = Y;
    return par[0][x];
  int dist(int x, int v) { // # edges on path
    return depth[x]+depth[y]-2*depth[lca(x,y)]; }
};
```

### LCArmq.h

**Description:** Euler Tour LCA. Compress takes a subset S of nodes and computes the minimal subtree that contains all the nodes pairwise LCAs and compressing edges. Returns a list of (par, origindex) representing a tree rooted at 0. The root points to itself.

**Time:**  $\mathcal{O}(N \log N)$  build,  $\mathcal{O}(1)$  LCA,  $\mathcal{O}(|S| \log |S|)$  compress "RMQ.h" e5a035, 28 lines

```
struct LCA {
 int N; V<vi> adj;
```

void ae(int x, int y) { adj[x].pb(y), adj[y].pb(x); } void dfs(int x) { pos[x] = sz(tmp); tmp.eb(depth[x],x);each(y,adj[x]) if (y != par[x]) { depth[y] = depth[par[y]=x]+1, dfs(y);tmp.eb(depth[x],x); } void gen(int R = 0) { par[R] = R; dfs(R); r.init(tmp); } int lca(int u, int v){ u = pos[u], v = pos[v]; if (u > v) swap(u, v); return r.querv(u,v).s; } int dist(int u, int v) { return depth[u]+depth[v]-2\*depth[lca(u,v)]; } vpi compress(vi S) { auto cmp = [&](int a, int b) { return pos[a] < pos[b]; };</pre> sort(all(S), cmp); R0F(i, sz(S)-1) S.pb(lca(S[i], S[i+1]));sort(all(S),cmp); S.erase(unique(all(S)),end(S));  $vpi ret{\{0,S[0]\}\}; F0R(i,sz(S)) rev[S[i]] = i;}$ FOR(i,1,sz(S)) ret.eb(rev[lca(S[i-1],S[i])],S[i]); return ret; }; HLD.h Description: Heavy-Light Decomposition, add val to verts and query sum in path/subtree. **Time:** any tree path is split into  $\mathcal{O}(\log N)$  parts "LazySeg.h" 1802e2, 48 lines template<int SZ, bool VALS IN EDGES> struct HLD { int N: vi adi[SZ]: int par[SZ], root[SZ], depth[SZ], sz[SZ], ti; int pos[SZ]; vi rpos; // rpos not used but could be useful void ae(int x, int y) { adj[x].pb(y), adj[y].pb(x); } void dfsSz(int x) { sz[x] = 1;each(y,adj[x]) { par[y] = x; depth[y] = depth[x]+1;adj[y].erase(find(all(adj[y]),x)); dfsSz(y); sz[x] += sz[y];if (sz[y] > sz[adj[x][0]]) swap(y,adj[x][0]); void dfsHld(int x) { pos[x] = ti++; rpos.pb(x);each(y,adj[x]) { root[y] = (y == adj[x][0] ? root[x] : y);dfsHld(y); } void init(int N, int R = 0) { N = N; par[R] = depth[R] = ti = 0; dfsSz(R);root[R] = R; dfsHld(R);

```
op(pos[x]+VALS_IN_EDGES,pos[y]);
void modifyPath(int x, int y, int v) {
  processPath(x,y,[this,&v](int 1, int r) {
    tree.upd(1,r,v); }); }
11 queryPath(int x, int y) {
  11 res = 0; processPath(x,y,[this,&res](int 1, int r) {
    res += tree.query(1,r); });
  return res; }
void modifvSubtree(int x, int v) {
  tree.upd(pos[x]+VALS_IN_EDGES, pos[x]+sz[x]-1,v); }
```

#### Centroid.h

**Description:** The centroid of a tree of size N is a vertex such that after removing it, all resulting subtrees have size at most  $\frac{N}{2}$ . Supports updates in the form "add 1 to all verts v such that  $dist(x, v) \leq \overline{y}$ ."

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

```
Time: \mathcal{O}(N \log N) build, \mathcal{O}(\log N) update and query
void ad(vi& a, int b) { ckmin(b,sz(a)-1); if (b>=0) a[b]++; }
void prop(vi& a) { R0F(i,sz(a)-1) a[i] += a[i+1]; }
template<int SZ> struct Centroid {
 vi adj[SZ]; void ae(int a, int b) {adj[a].pb(b), adj[b].pb(a);}
 bool done[SZ]; // processed as centroid yet
  int N, sub[SZ], cen[SZ], lev[SZ]; // subtree size, centroid and
  int dist[32-__builtin_clz(SZ)][SZ]; // dists to all ancs
 vi stor[SZ], STOR[SZ];
  void dfs(int x, int p) { sub[x] = 1;
    each(y,adj[x]) if (!done[y] && y != p)
      dfs(y,x), sub[x] += sub[y];
 int centroid(int x) {
    dfs(x,-1);
    for (int sz = sub[x];;) {
      pi mx = \{0, 0\};
      each(y,adj[x]) if (!done[y] \&\& sub[y] < sub[x])
        ckmax(mx, {sub[y],y});
      if (mx.f*2 \le sz) return x;
      x = mx.s;
 void genDist(int x, int p, int lev) {
    dist[lev][x] = dist[lev][p]+1;
    each(y,adj[x]) if (!done[y] \&\& y != p) genDist(y,x,lev); }
  void gen(int CEN, int _x) { // CEN = centroid above x
    int x = centroid(_x); done[x] = 1; cen[x] = CEN;
    sub[x] = sub[x]; lev[x] = (CEN == -1 ? 0 : lev[CEN]+1);
    dist[lev[x]][x] = 0;
    stor[x].rsz(sub[x]), STOR[x].rsz(sub[x]+1);
    each(y,adj[x]) if (!done[y]) genDist(y,x,lev[x]);
    each(y,adj[x]) if (!done[y]) gen(x,y);
 void init(int N) { N = N; FOR(i, 1, N+1) done[i] = 0;
    gen(-1,1); } // start at vert 1
  void upd(int x, int y) {
    int cur = x, pre = -1;
    ROF(i, lev[x]+1) {
      ad(stor[cur],y-dist[i][x]);
      if (pre != -1) ad(STOR[pre], y-dist[i][x]);
      if (i > 0) pre = cur, cur = cen[cur];
 } // call propAll() after all updates
 void propAll() { FOR(i,1,N+1) prop(stor[i]), prop(STOR[i]); }
 int query(int x) { // get value at vertex x
    int cur = x, pre = -1, ans = 0;
    ROF(i, lev[x]+1) { // if pre != -1, subtract those from
      ans += stor[cur][dist[i][x]]; // same subtree
      if (pre != -1) ans -= STOR[pre][dist[i][x]];
```

### 7.2.1 SqrtDecompton

HLD generally suffices. If not, here are some common strategies:

- Rebuild the tree after every  $\sqrt{N}$  queries.
- Consider vertices with > or  $<\sqrt{N}$  degree separately.
- For subtree updates, note that there are  $O(\sqrt{N})$  distinct sizes among child subtrees of any node.

Block Tree: Use a DFS to split edges into contiguous groups of size  $\sqrt{N}$  to  $2\sqrt{N}$ .

Mo's Algorithm for Tree Paths: Maintain an array of vertices where each one appears twice, once when a DFS enters the vertex (st) and one when the DFS exists (en). For a tree path  $u \leftrightarrow v$  such that st[u] < st[v],

- If u is an ancestor of v, query [st[u], st[v]].
- Otherwise, query [en[u], st[v]] and consider LCA(u, v) separately.

Solutions with worse complexities can be faster if you optimize the operations that are performed most frequently. Use arrays instead of vectors whenever possible. Iterating over an array in order is faster than iterating through the same array in some other order (ex. one given by a random permutation) or DFSing on a tree of the same size. Also, the difference between  $\sqrt{N}$  and the optimal block (or buffer) size can be quite large. Try up to 5x smaller or larger (at least).

## 7.3 DFS Algorithms

#### EulerPath.h

**Description:** Eulerian path starting at src if it exists, visits all edges exactly once. Works for both directed and undirected. Returns vector of {vertex,label of edge to vertex}. Second element of first pair is always -1. **Time:**  $\mathcal{O}\left(N+M\right)$ 

```
template<bool directed> struct Euler {
  int N; V<vpi> adj; V<vpi::iterator> its; vb used;
  void init(int _N) { N = _N; adj.rsz(N); }
  void ae(int a, int b) {
    int M = sz(used); used.pb(0);
    adj[a].eb(b,M); if (!directed) adj[b].eb(a,M); }
  vpi solve(int src = 0) {
    its.rsz(N); F0R(i,N) its[i] = begin(adj[i]);
```

```
vpi ans, s{{src,-1}}; // {{vert,prev vert},edge label}
int lst = -1; // ans generated in reverse order
while (sz(s)) {
   int x = s.bk.f; auto& it=its[x], en=end(adj[x]);
   while (it != en && used[it->s]) ++it;
   if (it == en) { // no more edges out of vertex
        if (lst != -1 && lst != x) return {};
        // not a path, no tour exists
        ans.pb(s.bk); s.pop_back(); if (sz(s)) lst=s.bk.f;
   } else s.pb(*it), used[it->s] = 1;
   } // must use all edges
   if (sz(ans) != sz(used)+1) return {};
   reverse(all(ans)); return ans;
}
};
```

### SCCT.h

**Description:** Tarjan's, DFS once to generate strongly connected components in topological order. a, b in same component if both  $a \to b$  and  $b \to a$  exist. Uses less memory than Kosaraju b/c doesn't store reverse edges.

Time:  $\mathcal{O}(N+M)$  a36e0c, 22 lin-

```
struct SCC {
 int N, ti = 0; V<vi> adj;
 vi disc, comp, stk, comps;
 void init(int _N) { N = _N, adj.rsz(N);
   disc.rsz(N), comp.rsz(N,-1);}
 void ae(int x, int y) { adj[x].pb(y); }
 int dfs(int x) {
   int low = disc[x] = ++ti; stk.pb(x);
   each(y,adj[x]) if (comp[y] == -1) // comp[y] == -1,
     ckmin(low, disc[y]?:dfs(y)); // disc[y] != 0 -> in stack
   if (low == disc[x]) { // make new SCC
     // pop off stack until you find x
     comps.pb(x); for (int y = -1; y != x;)
       comp[y = stk.bk] = x, stk.pop_back();
   return low;
 void gen() {
   FOR(i, N) if (!disc[i]) dfs(i);
   reverse(all(comps));
};
```

### TwoSAT.h

**Description:** Calculates a valid assignment to boolean variables a, b, c,... to a 2-SAT problem, so that an expression of the type (a|||b)&&(!a|||c)&&(d|||!b)&&... becomes true, or reports that it is unsatisfiable. Negated variables are represented by bit-inversions ( $\sim$ x).

```
Usage: TwoSat ts;

ts.either(0, ~3); // Var 0 is true or var 3 is false
ts.setVal(2); // Var 2 is true
ts.atMostOne({0,~1,2}); // <= 1 of vars 0, ~1 and 2 are true
ts.solve(N); // Returns true iff it is solvable
ts.ans[0..N-1] holds the assigned values to the vars

"SCC.h"

805elc, 32 lines
```

```
struct TwoSAT {
  int N = 0; vpi edges;
  void init(int _N) { N = _N; }
  int addVar() { return N++; } // for atMostOne
  void either(int x, int y) {
    x = max(2*x,-1-2*x), y = max(2*y,-1-2*y);
    edges.eb(x,y); }
  void implies(int x, int y) { either(~x,y); }
  void must(int x) { either(x,x); }
  void atMostOne(const vi& li) {
    if (sz(li) <= 1) return;
}</pre>
```

```
int cur = \simli[0];
    FOR(i, 2, sz(li)) {
      int next = addVar();
      either(cur,~li[i]); either(cur,next);
      either(~li[i],next); cur = ~next;
    either(cur,~li[1]);
  vb solve(int _N = -1) {
    if (N != -1) N = N;
    SCC S; S.init(2*N);
    each (e,edges) S.ae (e.f^1,e.s), S.ae (e.s^1,e.f);
    S.gen(); reverse(all(S.comps)); // reverse topo order
    for (int i = 0; i < 2*N; i += 2)
     if (S.comp[i] == S.comp[i^1]) return {};
    vi tmp(2*N); each(i,S.comps) if (!tmp[i])
      tmp[i] = 1, tmp[S.comp[i^1]] = -1;
    vb ans(N); FOR(i,N) ans[i] = tmp[S.comp[2*i]] == 1;
};
```

### BCC.h

};

**Description:** Biconnected components of edges. Removing any vertex in BCC doesn't disconnect it. To get block-cut tree, create a bipartite graph with the original vertices on the left and a vertex for each BCC on the right. Draw edge  $u \leftrightarrow v$  if u is contained within the BCC for v. Self-loops are not included in any BCC while BCCS of size 1 represent bridges.

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

```
0625a6, 35 lines
struct BCC {
 V<vpi> adj; vpi ed;
 V<vi> edgeSets, vertSets; // edges for each bcc
 int N, ti = 0; vi disc, stk;
 void init(int _N) { N = _N; disc.rsz(N), adj.rsz(N); }
 void ae(int x, int y) {
   adj[x].eb(y,sz(ed)), adj[y].eb(x,sz(ed)), ed.eb(x,y); }
 int dfs(int x, int p = -1) { // return lowest disc
   int low = disc[x] = ++ti;
    each(e,adj[x]) if (e.s != p) {
     if (!disc[e.f]) {
       stk.pb(e.s); // disc[x] < LOW -> bridge
       int LOW = dfs(e.f,e.s); ckmin(low,LOW);
       if (disc[x] <= LOW) { // get edges in bcc
         edgeSets.eb(); vi& tmp = edgeSets.bk; // new bcc
          for (int y = -1; y != e.s; )
            tmp.pb(y = stk.bk), stk.pop_back();
     } else if (disc[e.f] < disc[x]) // back-edge</pre>
        ckmin(low, disc[e.f]), stk.pb(e.s);
   return low;
 void gen() {
   FOR(i, N) if (!disc[i]) dfs(i);
    each(c,edgeSets) { // edges contained within each BCC
     vertSets.eb(); // so you can easily create block cut tree
     auto ad = [\&] (int x) {
       if (!in[x]) in[x] = 1, vertSets.bk.pb(x); };
      each(e,c) ad(ed[e].f), ad(ed[e].s);
      each(e,c) in[ed[e].f] = in[ed[e].s] = 0;
```

### MaximalCliques.h

Description: Used only once. Finds all maximal cliques.

Time:  $\mathcal{O}\left(3^{N/3}\right)$ f5cd93, 16 lines using B = bitset<128>; B adj[128]; // possibly in clique, not in clique, in clique void cliques (B P =  $\sim$ B(), B X={}, B R={}) { if (!P.anv()) { if (!X.anv()) // do smth with R return; int q = (P|X).\_Find\_first(); // clique must contain q or non-neighbor of q B cands =  $P\&\sim adj[q];$ F0R(i,N) if (cands[i]) { R[i] = 1; cliques(P&adj[i], X&adj[i], R); R[i] = P[i] = 0; X[i] = 1;

### 7.4 Flows

**Konig's Theorem:** In a bipartite graph, max matching = min vertex cover.

**Dilworth's Theorem:** For any partially ordered set, the sizes of the max antichain and of the min chain decomposition are equal. Equivalent to Konig's theorem on the bipartite graph (U, V, E) where U = V = S and (u, v) is an edge when u < v. Those vertices outside the min vertex cover in both U and V form a max antichain.

### Dinic.h

**Description:** Fast flow. After computing flow, edges  $\{u,v\}$  such that  $lev[u] \neq 0$ , lev[v] = 0 are part of min cut.

Time:  $\mathcal{O}\left(N^2M\right)$  flow

```
c76643, 43 lines
template<class F> struct Dinic {
  struct Edge { int to, rev; F cap; };
  int N; V<V<Edge>> adj;
  void init(int _N) { N = _N; adj.rsz(N); }
  pi ae(int a, int b, F cap, F rcap = 0) {
    assert (min(cap,rcap) >= 0); // saved me > once
    adj[a].pb({b,sz(adj[b]),cap});
   adj[b].pb({a,sz(adj[a])-1,rcap});
    return {a,sz(adj[a])-1};
  F edgeFlow(pi loc) { // get flow along original edge
   const Edge& e = adj.at(loc.f).at(loc.s);
    return adj.at(e.to).at(e.rev).cap;
 vi lev, ptr;
  bool bfs(int s, int t) { // level=shortest dist from source
   lev = ptr = vi(N);
   lev[s] = 1; queue < int > q({s});
   while (sz(q)) { int u = q.ft; q.pop();
     each(e,adj[u]) if (e.cap && !lev[e.to]) {
       q.push(e.to), lev[e.to] = lev[u]+1;
        if (e.to == t) return 1;
    return 0;
```

### MaximalCliques Dinic GomoryHu MCMF Hungarian

```
F dfs(int v, int t, F flo) {
   if (v == t) return flo;
    for (int& i = ptr[v]; i < sz(adj[v]); i++) {</pre>
     Edge& e = adj[v][i];
     if (lev[e.to]!=lev[v]+1||!e.cap) continue;
     if (F df = dfs(e.to,t,min(flo,e.cap))) {
       e.cap -= df; adj[e.to][e.rev].cap += df;
       return df; } // saturated >=1 one edge
   return 0;
 F maxFlow(int s, int t) {
   F tot = 0; while (bfs(s,t)) while (F df =
     dfs(s,t,numeric_limits<F>::max())) tot += df;
};
```

### GomorvHu.h

**Description:** Returns edges of Gomory-Hu tree (second element is weight). Max flow between pair of vertices of undirected graph is given by min edge weight along tree path. Uses the fact that for any  $i, j, k, \lambda_{ik} \geq \min(\lambda_{ij}, \lambda_{jk})$ , where  $\lambda_{ij}$  denotes the flow between i and j.

**Time:** N - 1 calls to Dinic

```
"Dinic.h"
                                                     0d712e, 16 lines
template<class F> V<pair<pi,F>> gomoryHu(int N,
   const V<pair<pi,F>>& ed) {
 vi par(N); Dinic<F> D; D.init(N);
 vpi ed_locs; each(t,ed)ed_locs.pb(D.ae(t.f.f,t.f.s,t.s,t.s));
 V<pair<pi,F>> ans;
 FOR(i,1,N) {
   each(p,ed_locs) { // reset capacities
     auto& e = D.adj.at(p.f).at(p.s);
     auto& e_rev = D.adj.at(e.to).at(e.rev);
     e.cap = e_rev.cap = (e.cap+e_rev.cap)/2;
   ans.pb({{i,par[i]},D.maxFlow(i,par[i])});
   FOR(j, i+1, N) if (par[j] == par[i] \&\& D.lev[j]) par[j] = i;
 return ans;
```

#### MCMF.h

Description: Minimum-cost maximum flow, assumes no negative cycles. It is possible to choose negative edge costs such that the first run of Dijkstra is slow, but this hasn't been an issue in the past. Edge weights  $\geq 0$  for every subsequent run. To get flow through original edges, assign ID's during ae. **Time:** Ignoring first run of Dijkstra,  $\mathcal{O}(FM \log M)$  if caps are integers and F is max flow.

```
77bfb0, 46 lines
using F = 11; using C = 11; // flow type, cost type
struct Edge { int to, rev; F flo, cap; C cost; };
int N; V<C> p, dist; vpi pre; V<V<Edge>> adj;
void init(int _N) { N = _N;
  p.rsz(N), adj.rsz(N), dist.rsz(N), pre.rsz(N); }
void ae(int u, int v, F cap, C cost) { assert(cap >= 0);
  adj[u].pb({v,sz(adj[v]),0,cap,cost});
  adj[v].pb({u,sz(adj[u])-1,0,0,-cost});
} // use asserts, don't try smth dumb
bool path(int s, int t) { // send flow through lowest cost
  const C inf = numeric_limits<C>::max(); dist.assign(N,inf);
  using T = pair<C, int>;
  priority_queue<T, V<T>, greater<T>> todo;
  todo.push(\{dist[s] = 0, s\});
  while (sz(todo)) { // Dijkstra
    T x = todo.top(); todo.pop();
```

```
if (x.f > dist[x.s]) continue;
      each(e,adj[x.s]) { // all weights should be non-negative
        if (e.flo < e.cap && ckmin(dist[e.to],
            x.f+e.cost+p[x.s]-p[e.to]))
          pre[e.to]={x.s,e.rev}, todo.push({dist[e.to],e.to});
    } // if costs are doubles, add some EPS so you
    // don't traverse ~0-weight cycle repeatedly
    return dist[t] != inf; // true if augmenting path
  pair<F,C> calc(int s, int t) { assert(s != t);
    FOR(_,N) FOR(i,N) each(e,adj[i]) // Bellman-Ford
      if (e.cap) ckmin(p[e.to],p[i]+e.cost);
    F \text{ totFlow} = 0; C \text{ totCost} = 0;
    while (path(s,t)) { // p -> potentials for Dijkstra
      FOR(i,N) p[i] += dist[i]; // don't matter for unreachable
      F df = numeric_limits<F>::max();
      for (int x = t; x != s; x = pre[x].f) {
        Edge& e = adj[pre[x].f][adj[x][pre[x].s].rev];
        ckmin(df,e.cap-e.flo); }
      totFlow += df; totCost += (p[t]-p[s])*df;
      for (int x = t; x != s; x = pre[x].f) {
        Edge& e = adj[x][pre[x].s]; e.flo -= df;
        adj[pre[x].f][e.rev].flo += df;
    } // get max flow you can send along path
    return {totFlow,totCost};
};
```

### Matching

### Hungarian.h

**Description:** Given array of (possibly negative) costs to complete each of N(1-indexed) jobs w/ each of M workers  $(N \leq M)$ , finds min cost to complete all jobs such that each worker is assigned to at most one job. Dijkstra with potentials works in almost the same way as MCMF.

```
Time: \mathcal{O}\left(N^2M\right)
                                                                                                     09d0ec, 28 lines
```

```
using C = 11;
C hungarian(const V<V<C>>& a) {
 int N = sz(a)-1, M = sz(a[0])-1; assert (N \le M);
 V<C> u(N+1), v(M+1); // potentials to make edge weights >= 0
  vi job(M+1);
  FOR(i,1,N+1) { // find alternating path with job i
    const C inf = numeric limits<C>::max();
    int w = 0; job[w] = i; // add "dummy" worker 0
    V<C> dist(M+1,inf); vi pre(M+1,-1); vb done(M+1);
    while (job[w]) { // dijkstra
      done[w] = 1; int j = job[w], nexW; C delta = inf;
      // fix dist[j], update dists from j
      FOR(W,M+1) if (!done[W]) { // try all workers
        if (ckmin(dist[W],a[j][W]-u[j]-v[W])) pre[W] = w;
        if (ckmin(delta,dist[W])) nexW = W;
      FOR(W,M+1) { // subtract constant from all edges going
        // from done -> not done vertices, lowers all
        // remaining dists by constant
        if (done[W]) u[job[W]] += delta, v[W] -= delta;
        else dist[W] -= delta;
    } // potentials adjusted so all edge weights \geq= 0
    for (int W; w; w = W) job[w] = job[W = pre[w]];
 } // job[w] = 0, found alternating path
 return -v[0]; // min cost
```

### $General Match Blossom \ \ General Weighted Match$

### GeneralMatchBlossom.h

Description: Variant on Gabow's Impl of Edmond's Blossom Algorithm. General unweighted max matching with 1-based indexing. If white [v] = 0 after solve() returns, v is part of every max matching.

**Time:**  $\mathcal{O}(NM)$ , faster in practice

```
fd5cc7, 50 lines
struct MaxMatching {
  int N; V<vi> adj;
  V<int> mate, first; vb white; vpi label;
  void init(int _N) { N = _N; adj = V<vi>(N+1);
    mate = first = vi(N+1); label = vpi(N+1); white = vb(N+1);
  void ae(int u, int v) { adj.at(u).pb(v), adj.at(v).pb(u); }
  int group(int x) { if (white[first[x]]) first[x] = group(
    \hookrightarrowfirst[x]);
    return first[x]; }
  void match (int p, int b) {
    swap(b,mate[p]); if (mate[b] != p) return;
    if (!label[p].s) mate[b] = label[p].f, match(label[p].f,b);

→ // vertex label

    else match(label[p].f,label[p].s), match(label[p].s,label[p
       \hookrightarrow].f); // edge label
  bool augment(int st) { assert(st);
    white[st] = 1; first[st] = 0; label[st] = \{0,0\};
    queue<int> q; q.push(st);
    while (!q.empty()) {
      int a = q.ft; q.pop(); // outer vertex
      each(b,adj[a]) { assert(b);
        if (white[b]) { // two outer vertices, form blossom
          int x = group(a), y = group(b), lca = 0;
          while (x||y) {
            if (y) swap(x,y);
            if (label[x] == pi{a,b}) { lca = x; break; }
            label[x] = {a,b}; x = group(label[mate[x]].first);
          for (int v: {group(a), group(b)}) while (v != lca) {
            assert(!white[v]); // make everything along path
            g.push(v); white[v] = true; first[v] = lca;
            v = group(label[mate[v]].first);
        } else if (!mate[b]) { // found augmenting path
          mate[b] = a; match(a,b); white = vb(N+1); // reset
          return true;
        } else if (!white[mate[b]]) {
          white[mate[b]] = true; first[mate[b]] = b;
          label[b] = \{0,0\}; label[mate[b]] = pi\{a,0\};
          q.push(mate[b]);
    return false;
  int solve() {
    int ans = 0;
   FOR(st,1,N+1) if (!mate[st]) ans += augment(st);
   FOR(st,1,N+1) if (!mate[st] && !white[st]) assert(!augment(
       \hookrightarrowst));
    return ans:
};
```

### GeneralWeightedMatch.h

**Description:** General max weight max matching with 1-based indexing. Edge weights must be positive, combo of UnweightedMatch and Hungarian. Time:  $\mathcal{O}(N^3)$ ?

120873, 145 lines

```
struct edge { int u,v,w; }; edge g[SZ*2][SZ*2];
void ae(int u, int v, int w) { g[u][v].w = g[v][u].w = w; }
int N, NX, lab[SZ*2], match[SZ*2], slack[SZ*2], st[SZ*2];
int par[SZ*2],floFrom[SZ*2][SZ],S[SZ*2],aux[SZ*2];
vi flo[SZ*2]; queue<int> q;
void init(int _N) { N = _N; // init all edges
  FOR(u, 1, N+1) FOR(v, 1, N+1) g[u][v] = \{u, v, 0\};
int eDelta(edge e) { // >= 0 at all times
  return lab[e.u]+lab[e.v]-q[e.u][e.v].w*2; }
void updSlack(int u, int x) { // smallest edge -> blossom x
  if (!slack[x] || eDelta(q[u][x]) < eDelta(q[slack[x]][x]))</pre>
    slack[x] = u; }
void setSlack(int x) {
  slack[x] = 0; FOR(u, 1, N+1) if (q[u][x].w > 0
   && st[u] != x && S[st[u]] == 0) updSlack(u,x); }
void gPush(int x) {
  if (x \le N) q.push(x);
  else each(t,flo[x]) qPush(t); }
void setSt(int x, int b) {
  st[x] = b; if (x > N) each(t,flo[x]) setSt(t,b); }
int getPr(int b, int xr) { // get even position of xr
  int pr = find(all(flo[b]), xr)-begin(flo[b]);
  if (pr&1) { reverse(1+all(flo[b])); return sz(flo[b])-pr; }
  return pr; }
void setMatch(int u, int v) { // rearrange flo[u], matches
  edge e = g[u][v]; match[u] = e.v; if (u <= N) return;</pre>
  int xr = floFrom[u][e.u], pr = getPr(u,xr);
  FOR(i,pr) setMatch(flo[u][i],flo[u][i^1]);
  setMatch(xr,v); rotate(begin(flo[u]),pr+all(flo[u])); }
void augment (int u, int v) { // set matches including u->v
  while (1) { // and previous ones
    int xnv = st[match[u]]; setMatch(u,v);
    if (!xnv) return;
    setMatch(xnv,st[par[xnv]]);
    u = st[par[xnv]], v = xnv;
int lca(int u, int v) { // same as in unweighted
  static int t = 0; // except maybe return 0
  for (++t;u||v;swap(u,v)) {
    if (!u) continue;
    if (aux[u] == t) return u;
    aux[u] = t; u = st[match[u]];
    if (u) u = st[par[u]];
  return 0:
void addBlossom(int u, int anc, int v) {
  int b = N+1; while (b \le NX \&\& st[b]) ++b;
  if (b > NX) ++NX; // new blossom
  lab[b] = S[b] = 0; match[b] = match[anc]; flo[b] = {anc};
  auto blossom = [&](int x) {
    for (int y; x != anc; x = st[par[y]])
      flo[b].pb(x), flo[b].pb(y = st[match[x]]), qPush(y);
  blossom(u); reverse(1+all(flo[b])); blossom(v); setSt(b,b);
  // identify all nodes in current blossom
  FOR(x, 1, NX+1) q[b][x].w = q[x][b].w = 0;
  FOR(x, 1, N+1) floFrom[b][x] = 0;
  each(xs,flo[b]) { // find tightest constraints
    FOR(x,1,NX+1) if (g[b][x].w == 0 \mid \mid eDelta(g[xs][x]) <
      eDelta(g[b][x])) g[b][x]=g[xs][x], g[x][b]=g[x][xs];
    FOR(x, 1, N+1) if (floFrom[xs][x]) floFrom[b][x] = xs;
  } // floFrom to deconstruct blossom
  setSlack(b); // since didn't qPush everything
void expandBlossom(int b) {
  each(t,flo[b]) setSt(t,t); // undo setSt(b,b)
  int xr = floFrom[b][g[b][par[b]].u], pr = getPr(b,xr);
```

```
for(int i = 0; i < pr; i += 2) {
    int xs = flo[b][i], xns = flo[b][i+1];
    par[xs] = q[xns][xs].u; S[xs] = 1; // no setSlack(xns)?
    S[xns] = slack[xs] = slack[xns] = 0; qPush(xns);
  S[xr] = 1, par[xr] = par[b];
  FOR(i,pr+1,sz(flo[b])) { // matches don't change
    int xs = flo[b][i]; S[xs] = -1, setSlack(xs); }
  st[b] = 0; // blossom killed
bool onFoundEdge(edge e) {
  int u = st[e.u], v = st[e.v];
  if (S[v] == -1) { // v unvisited, matched with smth else
    par[v] = e.u, S[v] = 1; slack[v] = 0;
    int nu = st[match[v]]; S[nu] = slack[nu] = 0; qPush(nu);
  } else if (S[v] == 0) {
    int anc = lca(u,v); // if 0 then match found!
    if (!anc) return augment(u,v), augment(v,u),1;
    addBlossom(u,anc,v);
  return 0;
bool matching() {
  q = queue<int>();
  FOR(x,1,NX+1) {
    S[x] = -1, slack[x] = 0; // all initially unvisited
    if (st[x] == x \&\& !match[x]) par[x] = S[x] = 0, qPush(x);
  if (!sz(q)) return 0;
  while (1) {
    while (sz(q)) { // unweighted matching with tight edges
      int u = q.ft; q.pop(); if (S[st[u]] == 1) continue;
      FOR(v, 1, N+1) if (q[u][v].w > 0 && st[u] != st[v]) {
        if (eDelta(g[u][v]) == 0) \{ // condition is strict \}
          if (onFoundEdge(g[u][v])) return 1;
        } else updSlack(u,st[v]);
    int d = INT_MAX;
    FOR(b, N+1, NX+1) if (st[b] == b \&\& S[b] == 1)
      ckmin(d,lab[b]/2); // decrease lab[b]
    FOR(x, 1, NX+1) if (st[x] == x \&\& slack[x]) {
      if (S[x] == -1) ckmin(d,eDelta(g[slack[x]][x]));
      else if (S[x] == 0) ckmin(d, eDelta(g[slack[x]][x])/2);
    } // edge weights shouldn't go below 0
    FOR(u, 1, N+1) {
      if (S[st[u]] == 0) {
        if (lab[u] <= d) return 0; // why?
        lab[u] -= d;
      } else if (S[st[u]] == 1) lab[u] += d;
    } // lab has opposite meaning for verts and blossoms
    FOR(b, N+1, NX+1) if (st[b] == b \&\& S[b] != -1)
      lab[b] += (S[b] == 0 ? 1 : -1) *d*2;
    q = queue<int>();
    FOR(x, 1, NX+1) if (st[x] == x \&\& slack[x] // new tight edge
      && st[slack[x]] != x && eDelta(q[slack[x]][x]) == 0)
        if (onFoundEdge(g[slack[x]][x])) return 1;
    FOR (b, N+1, NX+1) if (st[b] == b \&\& S[b] == 1 \&\& lab[b] == 0)
      expandBlossom(b); // odd dist blossom taken apart
  return 0;
pair<ll, int> calc() {
  NX = N; st[0] = 0; FOR(i, 1, 2*N+1) aux[i] = 0;
  FOR(i, 1, N+1) match[i] = 0, st[i] = i, flo[i].clear();
  int wMax = 0;
  FOR(u, 1, N+1) FOR(v, 1, N+1)
    floFrom[u][v] = (u == v ? u : 0), ckmax(wMax,q[u][v].w);
  FOR(u, 1, N+1) lab[u] = wMax; // start high and decrease
```

```
int num = 0; 11 wei = 0; while (matching()) ++num;
    FOR(u, 1, N+1) if (match[u] \&\& match[u] < u)
      wei += q[u][match[u]].w; // edges in matching
    return {wei, num};
};
```

#### MaxMatchFast.h

Description: Fast bipartite matching.

Time:  $\mathcal{O}\left(M\sqrt{N}\right)$ 

ec6c96, 31 lines

```
vpi maxMatch(int L, int R, const vpi& edges) {
 V < vi > adj = V < vi > (L);
  vi nxt(L,-1), prv(R,-1), lev, ptr;
  FOR(i,sz(edges)) adj.at(edges[i].f).pb(edges[i].s);
   lev = ptr = vi(L); int max lev = 0;
    queue<int> q; F0R(i,L) if (nxt[i]==-1) lev[i]=1, q.push(i);
    while (sz(q)) {
     int x = q.ft; q.pop();
      for (int y: adj[x]) {
       int z = prv[y];
       if (z == -1) \max_{lev} = lev[x];
       else if (!lev[z]) lev[z] = lev[x]+1, q.push(z);
      if (max_lev) break;
    if (!max_lev) break;
   FOR(i,L) if (lev[i] > max_lev) lev[i] = 0;
    auto dfs = [&](auto self, int x) -> bool {
     for (;ptr[x] < sz(adj[x]);++ptr[x]) {</pre>
       int y = adj[x][ptr[x]], z = prv[y];
       if (z == -1 \mid | (lev[z] == lev[x]+1 \&\& self(self,z)))
          return nxt[x]=y, prv[y]=x, ptr[x]=sz(adj[x]), 1;
     return 0;
    };
   FOR(i,L) if (nxt[i] == -1) dfs(dfs,i);
  vpi ans; FOR(i,L) if (nxt[i] != -1) ans.pb(\{i,nxt[i]\});
 return ans;
```

### Advanced

### ChordalGraphRecognition.h

**Description:** Recognizes graph where every induced cycle has length exactly 3 using maximum adjacency search. 6cc97d, 58 lines

```
int N,M;
set<int> adj[MX];
int cnt[MX];
vi ord, rord;
vi find_path(int x, int y, int z) {
  vi pre(N,-1);
  queue<int> q; q.push(x);
  while (sz(q)) {
    int t = q.ft; q.pop();
    if (adj[t].count(y)) {
      pre[y] = t; vi path = {y};
     while (path.bk != x) path.pb(pre[path.bk]);
     path.pb(z);
      return path;
    each(u,adj[t]) if (u != z \&\& !adj[u].count(z) \&\& pre[u] ==
       →-1) {
```

```
pre[u] = t;
      q.push(u);
 assert(0);
int main() {
 setIO(); re(N,M);
 F0R(i,M) {
    int a,b; re(a,b);
    adj[a].insert(b), adj[b].insert(a);
 rord = vi(N, -1);
 priority_queue<pi> pq;
 FOR(i,N) pq.push(\{0,i\});
 while (sz(pq)) {
   pi p = pq.top(); pq.pop();
    if (rord[p.s] != -1) continue;
    rord[p.s] = sz(ord); ord.pb(p.s);
    each(t,adj[p.s]) pq.push({++cnt[t],t});
 assert(sz(ord) == N);
 each(z,ord) {
   pi biq = \{-1, -1\};
    each(y,adj[z]) if (rord[y] < rord[z])
     ckmax(big, mp(rord[y],y));
    if (big.f == -1) continue;
    int y = big.s;
    each(x,adj[z]) if (rord[x] < rord[y]) if (!adj[y].count(x))
       \hookrightarrow {
      ps("NO");
     vi v = find_path(x,y,z);
     ps(sz(v));
     each(t,v) pr(t,' ');
      exit(0);
 ps("YES");
 reverse(all(ord));
 each(z,ord) pr(z,' ');
```

#### DominatorTree.h

**Description:** Used only a few times. Assuming that all nodes are reachable from root, a dominates b iff every path from root to b passes through a. Time:  $\mathcal{O}\left(M\log N\right)$ 

```
template<int SZ> struct Dominator {
 vi adj[SZ], ans[SZ]; // input edges, edges of dominator tree
 vi radj[SZ], child[SZ], sdomChild[SZ];
 int label[SZ], rlabel[SZ], sdom[SZ], dom[SZ], co = 0;
 int par[SZ], bes[SZ];
 void ae(int a, int b) { adj[a].pb(b); }
 int get(int x) { // DSU with path compression
    // get vertex with smallest sdom on path to root
   if (par[x] != x) {
     int t = get(par[x]); par[x] = par[par[x]];
     if (sdom[t] < sdom[bes[x]]) bes[x] = t;</pre>
   return bes[x];
 void dfs(int x) { // create DFS tree
   label[x] = ++co; rlabel[co] = x;
   sdom[co] = par[co] = bes[co] = co;
   each(y,adj[x]) {
     if (!label[y]) {
        dfs(y); child[label[x]].pb(label[y]); }
      radj[label[y]].pb(label[x]);
```

```
void init(int root) {
    dfs(root);
    ROF(i, 1, co+1) {
      each(j,radj[i]) ckmin(sdom[i],sdom[get(j)]);
      if (i > 1) sdomChild[sdom[i]].pb(i);
      each(j,sdomChild[i]) {
       int k = qet(j);
        if (sdom[i] == sdom[k]) dom[i] = sdom[i];
        else dom[j] = k;
      each(j,child[i]) par[j] = i;
    FOR(i, 2, co+1) {
      if (dom[i] != sdom[i]) dom[i] = dom[dom[i]];
      ans[rlabel[dom[i]]].pb(rlabel[i]);
};
```

### EdgeColor.h

**Description:** Used only once. Naive implementation of Misra & Gries edge coloring. By Vizing's Theorem, a simple graph with max degree d can be edge colored with at most d+1 colors

**Time:**  $\mathcal{O}(N^2M)$ , faster in practice

cc2b29, 40 lines

```
template<int SZ> struct EdgeColor {
 int N = 0, maxDeg = 0, adj[SZ][SZ], deg[SZ];
 void init(int N) { N = N;
    FOR(i,N) \{ deg[i] = 0; FOR(j,N) adj[i][j] = 0; \} \}
  void ae(int a, int b, int c) {
   adj[a][b] = adj[b][a] = c; }
  int delEdge(int a, int b) {
   int c = adj[a][b]; adj[a][b] = adj[b][a] = 0;
    return c; }
 V<bool> genCol(int x) {
    V < bool > col(N+1); FOR(i,N) col[adj[x][i]] = 1;
    return col; }
  int freeCol(int u) {
    auto col = genCol(u); int x = 1;
    while (col[x]) ++x; return x; }
  void invert(int x, int d, int c) {
    FOR(i,N) if (adj[x][i] == d)
      delEdge(x,i), invert(i,c,d), ae(x,i,c); }
  void ae(int u, int v) {
    // check if you can add edge w/o doing any work
    assert(N); ckmax(maxDeg,max(++deg[u],++deg[v]));
    auto a = genCol(u), b = genCol(v);
    FOR(i,1,maxDeg+2) if (!a[i] && !b[i])
      return ae(u, v, i);
    V < bool > use(N); vi fan = \{v\}; use[v] = 1;
    while (1) {
      auto col = genCol(fan.bk);
      if (sz(fan) > 1) col[adj[fan.bk][u]] = 0;
      int i=0; while (i<N && (use[i] || col[adj[u][i]])) i++;</pre>
      if (i < N) fan.pb(i), use[i] = 1;</pre>
      else break:
    int c = freeCol(u), d = freeCol(fan.bk); invert(u,d,c);
    int i = 0; while (i < sz(fan) && genCol(fan[i])[d]
      && adj[u][fan[i]] != d) i ++;
    assert (i != sz(fan));
    FOR(j,i) ae(u,fan[j],delEdge(u,fan[j+1]));
    ae(u,fan[i],d);
};
```

### DirectedMST LCT ComplexComp

#### DirectedMST.h

**Description:** Chu-Liu-Edmonds algorithm. Computes minimum weight directed spanning tree rooted at r, edge from  $par[i] \rightarrow i$  for all  $i \neq r$ . Use DSU with rollback if need to return edges.

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

"DSUrb.h" 5d5c10, 61 lines struct Edge { int a, b; ll w; }; struct Node { // lazv skew heap node Edge key; Node \*1, \*r; ll delta; void prop() { key.w += delta; if (1) 1->delta += delta; if (r) r->delta += delta; delta = 0;Edge top() { prop(); return key; } Node \*merge(Node \*a, Node \*b) { if (!a || !b) return a ?: b; a->prop(), b->prop(); if (a->key.w > b->key.w) swap(a, b); swap(a->1, a->r = merge(b, a->r));return a; void pop(Node\*& a) { a->prop(); a = merge(a->1, a->r); } pair<ll, vi> dmst(int n, int r, const vector<Edge>& g) { DSUrb dsu; dsu.init(n); vector<Node\*> heap(n); // store edges entering each vertex // in increasing order of weight each(e,q) heap[e.b] = merge(heap[e.b], new Node{e}); ll res = 0; vi seen(n,-1); seen[r] = r; vpi in(n, {-1,-1}); // edge entering each vertex in MST vector<pair<int, vector<Edge>>> cvcs; F0R(s,n) { int u = s, w;vector<pair<int, Edge>> path; while (seen[u] < 0) {</pre> if (!heap[u]) return {-1,{}}; seen[u] = s; Edge e = heap[u]->top(); path.pb({u,e}); heap[u]->delta -= e.w, pop(heap[u]); res += e.w, u = dsu.get(e.a);if (seen[u] == s) { // found cycle, contract Node\* cyc = 0; cycs.eb(); cyc = merge(cyc, heap[w = path.bk.f]); cycs.bk.s.pb(path.bk.s); path.pop\_back(); } while (dsu.unite(u,w)); u = dsu.get(u); heap[u] = cyc, seen[u] = -1;cycs.bk.f = u;each (t,path) in  $[dsu.get(t.s.b)] = \{t.s.a,t.s.b\};$ } // found path from root to s, done while (sz(cycs)) { // expand cycs to restore sol auto c = cycs.bk; cycs.pop\_back(); pi inEdge = in[c.f]; each(t,c.s) dsu.rollback();  $each(t,c.s) in[dsu.get(t.b)] = \{t.a,t.b\};$ in[dsu.get(inEdge.s)] = inEdge; vi par(n); FOR(i,n) par[i] = in[i].f;// i == r ? in[i].s == -1 : in[i].s == ireturn {res,par};

### LCT.h

**Description:** Link-Cut Tree. Given a function  $f(1\dots N)\to 1\dots N$ , evaluates  $f^b(a)$  for any a,b. sz is for path queries; sub, vsub are for subtree queries. x->access() brings x to the top and propagates it; its left subtree will be the path from x to the root and its right subtree will be empty. Then sub will be the number of nodes in the connected component of x and vsub will be the number of nodes under x. Use makeRoot for arbitrary path queries.

Usage: FOR(i,1,N+1)LCT[i]=new snode(i); link(LCT[1],LCT[2],1); Time:  $\mathcal{O}(\log N)$ 

```
typedef struct snode* sn;
struct snode { ////// VARIABLES
 sn p, c[2]; // parent, children
 sn extra; // extra cycle node for "The Applicant"
 bool flip = 0; // subtree flipped or not
 int val, sz; // value in node, # nodes in current splay tree
 int sub, vsub = 0; // vsub stores sum of virtual children
 snode(int val) : val( val) {
   p = c[0] = c[1] = extra = NULL; calc(); }
 friend int getSz(sn x) { return x?x->sz:0; }
 friend int getSub(sn x) { return x?x->sub:0; }
 void prop() { // lazy prop
   if (!flip) return;
   swap(c[0],c[1]); flip = 0;
   FOR(i,2) if (c[i]) c[i]->flip ^= 1;
 void calc() { // recalc vals
   FOR(i,2) if (c[i]) c[i]->prop();
   sz = 1+getSz(c[0])+getSz(c[1]);
   sub = 1+getSub(c[0])+getSub(c[1])+vsub;
 ////// SPLAY TREE OPERATIONS
 int dir() {
   if (!p) return -2;
   FOR(i,2) if (p->c[i] == this) return i;
   return -1; // p is path-parent pointer
 } // -> not in current splay tree
 // test if root of current splay tree
 bool isRoot() { return dir() < 0; }</pre>
 friend void setLink(sn x, sn y, int d) {
   if (y) y -> p = x;
   if (d >= 0) x -> c[d] = y; }
 void rot() { // assume p and p->p propagated
   assert(!isRoot()); int x = dir(); sn pa = p;
   setLink(pa->p, this, pa->dir());
   setLink(pa, c[x^1], x); setLink(this, pa, x^1);
   pa->calc();
 void splay() {
   while (!isRoot() && !p->isRoot()) {
     p->p->prop(), p->prop(), prop();
     dir() == p->dir() ? p->rot() : rot();
     rot();
    if (!isRoot()) p->prop(), prop(), rot();
   prop(); calc();
 sn fbo(int b) { // find by order
   prop(); int z = getSz(c[0]); // of splay tree
   if (b == z) { splay(); return this; }
    return b < z ? c[0] -> fbo(b) : c[1] -> fbo(b-z-1);
 ////// BASE OPERATIONS
 void access() { // bring this to top of tree, propagate
   for (sn v = this, pre = NULL; v; v = v->p) {
     v->splay(); // now switch virtual children
     if (pre) v->vsub -= pre->sub;
      if (v->c[1]) v->vsub += v->c[1]->sub;
```

```
v->c[1] = pre; v->calc(); pre = v;
    splay(); assert(!c[1]); // right subtree is empty
  void makeRoot() {
    access(); flip ^= 1; access(); assert(!c[0] && !c[1]); }
  ////// OUERIES
  friend sn lca(sn x, sn y) {
    if (x == y) return x;
    x->access(), y->access(); if (!x->p) return NULL;
    x->splay(); return x->p?:x; // y was below x in latter case
  } // access at y did not affect x -> not connected
  friend bool connected(sn x, sn y) { return lca(x,y); }
  // # nodes above
  int distRoot() { access(); return getSz(c[0]); }
  sn getRoot() { // get root of LCT component
    access(); sn a = this;
    while (a\rightarrow c[0]) a = a\rightarrow c[0], a\rightarrow prop();
    a->access(); return a;
  sn getPar(int b) { // get b-th parent on path to root
    access(); b = getSz(c[0]) - b; assert(b >= 0);
    return fbo(b);
  } // can also get min, max on path to root, etc
  ////// MODIFICATIONS
  void set(int v) { access(); val = v; calc(); }
  friend void link(sn x, sn y, bool force = 0) {
    assert(!connected(x,y));
    if (force) v->makeRoot(); // make x par of v
    else { y->access(); assert(!y->c[0]); }
    x->access(); setLink(v,x,0); v->calc();
  friend void cut(sn y) { // cut y from its parent
    v->access(); assert(v->c[0]);
    y->c[0]->p = NULL; y->c[0] = NULL; y->calc(); }
  friend void cut(sn x, sn y) { // if x, y adj in tree
    x->makeRoot(); y->access();
    assert (y->c[0] == x && !x->c[0] && !x->c[1]); cut (y); }
};
sn LCT[MX];
////// THE APPLICANT SOLUTION
void setNex(sn a, sn b) { // set f[a] = b
 if (connected(a,b)) a->extra = b;
 else link(b,a); }
void delNex(sn a) { // set f[a] = NULL
  auto t = a->getRoot();
  if (t == a) { t->extra = NULL; return; }
  cut(a); assert(t->extra);
  if (!connected(t,t->extra))
    link(t->extra,t), t->extra = NULL;
sn getPar(sn a, int b) { // get f^b[a]
  int d = a->distRoot(); if (b <= d) return a->getPar(b);
  b -= d+1; auto r = a->getRoot()->extra; assert(r);
  d = r->distRoot()+1; return r->getPar(b%d);
```

# Geometry (8)

### 8.1 Primitives

### ${\bf ComplexComp.h}$

**Description:** Allows you to sort complex numbers.

6f828b, 5 lines

```
#define x real()
#define y imag()
```

```
using P = complex<db>;
namespace std {
  bool operator<(P 1,P r) { return mp(1.x,1.y)<mp(r.x,r.y); } }</pre>
```

### PointShort.h

 $\begin{tabular}{ll} \textbf{Description:} \end{tabular} Use in place of $$\operatorname{\texttt{complex}}$<&$T>$. \end{tabular}$ 

cefef8, 36 lines

```
using T = db; // or 11
const T EPS = 1e-9; // adjust as needed
using P = pair<T,T>; using vP = V<P>; using Line = pair<P,P>;
int sqn(T a) { return (a>EPS)-(a<-EPS); }</pre>
T sq(T a) { return a*a; }
T norm(P p) { return sq(p.f)+sq(p.s); }
T abs(P p) { return sqrt(norm(p)); }
T arg(P p) { return atan2(p.s,p.f); }
P conj(P p) { return P(p.f,-p.s); }
P perp(P p) { return P(-p.s,p.f); }
P dir(T ang) { return P(cos(ang), sin(ang)); }
P operator+(P 1, P r) { return P(1.f+r.f,1.s+r.s); }
P operator-(P 1, P r) { return P(1.f-r.f,1.s-r.s); }
P operator*(P 1, T r) { return P(1.f*r,1.s*r); }
P operator/(P 1, T r) { return P(1.f/r,1.s/r); }
P operator*(P 1, P r) { // complex # multiplication
  return P(1.f*r.f-l.s*r.s,l.s*r.f+l.f*r.s); }
P operator/(P 1, P r) { return 1*conj(r)/norm(r); }
P unit (const P& p) { return p/abs(p); }
T dot(const P& a, const P& b) { return a.f*b.f+a.s*b.s; }
T dot(const P& p, const P& a, const P& b) { return dot(a-p,b-p)
T cross(const P& a, const P& b) { return a.f*b.s-a.s*b.f; }
T cross(const P& p, const P& a, const P& b) {
  return cross(a-p,b-p); }
P reflect (const P& p, const Line& 1) {
  P = 1.f, d = 1.s-1.f;
  return a+conj((p-a)/d)*d; }
P foot (const P& p, const Line& 1) {
  return (p+reflect(p,1))/(T)2; }
bool onSeg(const P& p, const Line& 1) {
  return sgn(cross(1.f,l.s,p)) == 0 \&\& sgn(dot(p,l.f,l.s)) <= 0
ostream& operator << (ostream& os, P p) {
  return os << "(" << p.f << "," << p.s << ")"; }
```

### AngleCmp.h

**Description:** Sorts points in ccw order about origin in the same way as atan2, which returns real in  $(-\pi, \pi]$  so points on negative x-axis come last.

"Point.h"

2df5fc, 12 lines

#### SegDist.h

**Description:** computes distance between P and line (segment) AB

"Point.h" Ocb69a, 6 lines

```
T lineDist(const P& p, const Line& 1) {
  return abs(cross(p,1.f,1.s))/abs(1.f-1.s); }
T segDist(const P& p, const Line& 1) {
  if (dot(1.f,p,1.s) <= 0) return abs(p-1.f);
  if (dot(1.s,p,1.f) <= 0) return abs(p-1.s);
  return lineDist(p,1); }</pre>
```

### SegIsect.h

**Description:** computes the intersection point(s) of line (segments) a and b

```
// {unique intersection point} if it exists
// {b.f,b.s} if input lines are the same
// empty if lines do not intersect
vP lineIsect(const Line& a, const Line& b) {
 T = a\theta = cross(a.f.a.s.b.f), al = cross(a.f.a.s.b.s);
 if (a0 == a1) return a0 == 0 ? vP{b.f,b.s} : vP{};
 return { (b.s*a0-b.f*a1) / (a0-a1) };
// point in interior of both segments a and b, if it exists
vP strictIsect(const Line& a, const Line& b) {
 T = a0 = cross(a.f,a.s,b.f), a1 = cross(a.f,a.s,b.s);
 T b0 = cross(b.f,b.s,a.f), b1 = cross(b.f,b.s,a.s);
 if (sgn(a0) * sgn(a1) < 0 && sgn(b0) * sgn(b1) < 0)
   return { (b.s*a0-b.f*a1) / (a0-a1) };
 return {}:
// intersection of segments, a and b may be degenerate
vP segIsect(const Line& a, const Line& b) {
 vP v = strictIsect(a,b); if (sz(v)) return v;
 set<P> s;
 \#define i(x,y) if (onSeg(x,y)) s.ins(x)
 i(a.f,b); i(a.s,b); i(b.f,a); i(b.s,a);
 return {all(s)};
```

# 8.2 Polygons

#### PolygonCenArea.h

**Description:** centroid (center of mass) of a polygon with constant mass per unit area and SIGNED area

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

### InPolygon.h

**Description:** Tests whether point is inside, on, or outside of a polygon (returns -1, 0, or 1). Both CW and CCW polygons are ok.

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

#### ConvexHull.h

 $\textbf{Description:} \ \operatorname{top-bottom} \ \operatorname{convex} \ \operatorname{hull}$ 

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

```
"Point.h"
                                                      868655, 18 lines
pair<vi, vi> ulHull(const vP& v) {
 vi p(sz(v)), u, 1; iota(all(p), 0);
  sort(all(p), [&v](int a, int b) { return v[a] < v[b]; });</pre>
  each(i,p) {
    #define ADDP(C, cmp) while (sz(C) > 1 && cross(\
     v[C[sz(C)-2]], v[C.bk], v[i]) cmp 0) C.pop_back(); C.pb(i);
    ADDP(u, >=); ADDP(1, <=);
 return {u,1};
vi hullInd(const vP& v) { // returns indices in CCW order
 vi u, l; tie(u, l) = ulHull(v); if (sz(l) <= 1) return l;
 if (v[1[0]] == v[1[1]]) return {0};
 1.insert (end(1),1+rall(u)-1); return 1;
vP hull(const vP& v) {
 vi w = hullInd(v); vP res; each(t,w) res.pb(v[t]);
 return res; }
```

### HullDiameter.h

**Description:** rotating caliphers, gives greatest distance between two points in  ${\cal P}$ 

**Time:**  $\mathcal{O}(N)$  given convex hull

#### LineHull.h

Description: lineHull accepts line and ccw convex polygon. If all vertices in poly lie to one side of the line, returns a vector of closest vertices to line as well as orientation of poly with respect to line (±1 for above/below). Otherwise, returns the range of vertices that lie on or below the line. extrVertex returns the point of a hull with the max projection onto a line.

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

```
"Point.h"
                                                      40e5a6, 41 lines
using Line = AR<P.2>:
#define cmp(i,j) sgn(-dot(dir,poly[(i)%n]-poly[(j)%n]))
#define extr(i) cmp(i+1,i) >= 0 && cmp(i,i-1+n) < 0
int extrVertex(const vP& poly, P dir) {
  int n = sz(polv), lo = 0, hi = n;
  if (extr(0)) return 0;
  while (lo+1 < hi) {
    int m = (lo+hi)/2;
    if (extr(m)) return m;
    int ls = cmp(lo+1, lo), ms = cmp(m+1, m);
    (ls < ms \mid | (ls == ms \&\& ls == cmp(lo, m)) ? hi : lo) = m;
  return lo;
vi same(Line line, const vP& poly, int a) {
  // points on same parallel as a
  int n = sz(poly); P dir = perp(line[0]-line[1]);
  if (cmp(a+n-1,a) == 0) return \{(a+n-1) n,a\};
  if (cmp(a,a+1) == 0) return \{a,(a+1)\%n\};
  return {a};
```

```
#define cmpL(i) sqn(cross(line[0],line[1],poly[i]))
pair<int, vi> lineHull(Line line, const vP& poly) {
  int n = sz(poly); assert(n>1);
  int endA = extrVertex(poly,perp(line[0]-line[1])); // lowest
  if (cmpL(endA) >= 0) return {1, same(line, poly, endA)};
  int endB = extrVertex(poly,perp(line[1]-line[0])); // highest
  if (cmpL(endB) <= 0) return {-1, same(line, poly, endB)};</pre>
  AR<int,2> res;
  F0R(i,2) {
    int lo = endA, hi = endB; if (hi < lo) hi += n;
    while (lo < hi) {
     int m = (lo+hi+1)/2;
     if (cmpL(m%n) == cmpL(endA)) lo = m;
     else hi = m-1;
   res[i] = lo%n; swap(endA,endB);
  if (cmpL((res[0]+1)%n) == 0) res[0] = (res[0]+1)%n;
  return {0, {(res[1]+1)%n, res[0]}};
```

#### HalfPlaneIsect.h

MIT

**Description:** Returns vertices of half-plane intersection. A half-plane is the area to the left of a ray, which is defined by a point p and a direction dp. Area of intersection should be sufficiently precise when all inputs are integers with magnitude  $\leq 10^5$ . Intersection must be bounded. Probably works with floating point too (but EPS might need to be adjusted?).

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

```
"AngleCmp.h"
                                                        18f712, 52 lines
struct Ray {
 P p, dp; // origin, direction
  P isect(const Ray& L) const {
    return p+dp*(cross(L.dp,L.p-p)/cross(L.dp,dp)); }
  bool operator<(const Ray& L) const {
    return angleCmp(dp,L.dp); }
vP halfPlaneIsect(V<Ray> rays, bool add bounds = false) {
  if (add bounds) {
    int DX = 1e9, DY = 1e9; // bound input by rectangle [\theta, DX]
       \hookrightarrow x [0, DY]
    rays.pb(\{P\{0,0\},P\{1,0\}\});
    rays.pb(\{P\{DX, 0\}, P\{0, 1\}\});
    rays.pb(\{P\{DX, DY\}, P\{-1, 0\}\});
    rays.pb(\{P\{0,DY\},P\{0,-1\}\});
  sor(rays); // sort rays by angle
  { // remove parallel rays
    V<Ray> nrays;
    each(t,rays) {
      if (!sz(nrays) || cross(nrays.bk.dp,t.dp) > EPS) { nrays.
         \hookrightarrow pb(t); continue; }
      // last two rays are parallel, keep only one
      if (cross(t.dp,t.p-nrays.bk.p) > 0) nrays.bk = t;
    swap(rays, nrays);
  auto bad = [&] (const Ray& a, const Ray& b, const Ray& c) {
    P p1 = a.isect(b), p2 = b.isect(c);
    if (dot(p2-p1,b.dp) \le EPS) {
      if (cross(a.dp,c.dp) \le 0) return 2; // isect(a,b,c) =
      return 1; // isect(a,c) == isect(a,b,c)
    return 0; // all three rays matter
  #define reduce(t) \
    while (sz(poly) > 1) \{ \
```

```
int b = bad(poly.at(sz(poly)-2),poly.bk,t); \
   if (b == 2) return {}; \
    if (b == 1) poly.pop_back(); \
   else break; \
deque<Ray> poly;
each(t,rays) { reduce(t); poly.pb(t); }
for(;;poly.pop_front()) {
 reduce(poly[0]);
 if (!bad(poly.bk,poly[0],poly[1])) break;
assert(sz(poly) >= 3); // expect nonzero area
vP poly points; F0R(i,sz(poly))
 poly_points.pb(poly[i].isect(poly[(i+1)%sz(poly)]));
return poly_points;
```

### 8.3 Circles

#### Circle.h

Description: represent circle as {center,radius}

```
"Point.h"
                                                       91f3fc, 6 lines
using Circ = pair<P,T>;
int in(const Circ& x, const P& y) { // -1 if inside, 0, 1
 return sqn(abs(y-x.f)-x.s); }
T arcLength(const Circ& x, P a, P b) {
    // precondition: a and b on x
 P d = (a-x.f)/(b-x.f); return x.s*acos(d.f); }
```

#### CircleIsect.h

Description: Circle intersection points and intersection area. Tangents will be returned twice.

```
"Circle.h"
                                                     a0b0f8, 22 lines
vP isect(const Circ& x, const Circ& y) { // precondition: x!=y
 T d = abs(x.f-y.f), a = x.s, b = y.s;
 if (sqn(d) == 0) { assert(a != b); return {}; }
 T C = (a*a+d*d-b*b) / (2*a*d);
  if (abs(C) > 1+EPS) return {};
 T S = sqrt(max(1-C*C, (T)0)); P tmp = (y.f-x.f)/d*x.s;
 return \{x.f+tmp*P(C,S),x.f+tmp*P(C,-S)\};
vP isect(const Circ& x, const Line& y) {
 Pc = foot(x.f,y); Tsq_dist = sq(x.s)-norm(x.f-c);
 if (sqn(sq_dist) < 0) return {};</pre>
 P offset = unit(y.s-y.f) *sqrt(max(sq_dist,T(0)));
 return {c+offset,c-offset};
T isect_area(Circ x, Circ y) { // not thoroughly tested
 T d = abs(x.f-y.f), a = x.s, b = y.s; if (a < b) swap(a,b);
 if (d \ge a+b) return 0:
 if (d <= a-b) return PI*b*b;
 T ca = (a*a+d*d-b*b)/(2*a*d), cb = (b*b+d*d-a*a)/(2*b*d);
 T s = (a+b+d)/2, h = 2*sqrt(s*(s-a)*(s-b)*(s-d))/d;
 return a*a*acos(ca)+b*b*acos(cb)-d*h;
```

### CircleTangents.h

Description: internal and external tangents between two circles

```
P tangent (P x, Circ y, int t = 0) {
 y.s = abs(y.s); // abs needed because internal calls y.s < 0
 if (y.s == 0) return y.f;
 T d = abs(x-y.f);
 P = pow(y.s/d, 2) * (x-y.f) + y.f;
 P b = sqrt(d*d-y.s*y.s)/d*y.s*unit(x-y.f)*dir(PI/2);
 return t == 0 ? a+b : a-b;
```

```
V<pair<P,P>> external(Circ x, Circ y) {
 V<pair<P,P>> v;
 if (x.s == y.s) {
   P \text{ tmp} = \text{unit}(x.f-y.f)*x.s*dir(PI/2);
    v.eb(x.f+tmp,y.f+tmp);
    v.eb(x.f-tmp,y.f-tmp);
 } else {
    P p = (y.s*x.f-x.s*y.f)/(y.s-x.s);
    FOR(i,2) v.eb(tangent(p,x,i),tangent(p,y,i));
 return v;
V<pair<P,P>> internal(Circ x, Circ y) {
 return external({x.f,-x.s},v); }
```

#### Circumcenter.h

**Description:** returns {circumcenter,circumradius}

```
"Circle.h"
                                                        a2c6a6, 5 lines
Circ ccCenter(P a, P b, P c) {
 b -= a; c -= a;
 P res = b*c*(conj(c)-conj(b))/(b*conj(c)-conj(b)*c);
 return {a+res, abs(res)};
```

### MinEnclosingCirc.h

Description: minimum enclosing circle

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

```
"Circumcenter.h"
                                                      53963d, 13 lines
circ mec(vP ps) {
  shuffle(all(ps), rng);
 P \circ = ps[0]; T r = 0, EPS = 1+1e-8;
  FOR(i,sz(ps)) if (abs(o-ps[i]) > r*EPS) {
    o = ps[i], r = 0; // point is on MEC
    FOR(j,i) if (abs(o-ps[j]) > r*EPS) {
      o = (ps[i]+ps[j])/2, r = abs(o-ps[i]);
      FOR(k,j) if (abs(o-ps[k]) > r*EPS)
        tie(o,r) = ccCenter(ps[i],ps[j],ps[k]);
  return {o,r};
```

### 8.4 Misc

#### ClosestPair.h

Description: Line sweep to find two closest points .

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

```
"Point.h"
                                                      2b60fa, 17 lines
pair<P,P> solve(vP v) {
 pair<db, pair<P, P>> bes; bes.f = INF;
 set < P > S; int ind = 0;
  sort(all(v));
 FOR(i,sz(v)) {
    if (i && v[i] == v[i-1]) return {v[i],v[i]};
    for (; v[i].f-v[ind].f >= bes.f; ++ind)
      S.erase({v[ind].s,v[ind].f});
    for (auto it = S.ub({v[i].s-bes.f,INF});
      it != end(S) && it->f < v[i].s+bes.f; ++it) {
      P t = \{it->s, it->f\};
      ckmin(bes, {abs(t-v[i]), {t,v[i]}});
    S.insert({v[i].s,v[i].f});
  return bes.s;
```

return q;

### DelaunayIncremental.h

Description: Bowyer-Watson where not all points collinear. Works for  $|x|, |y| \le 10^4$ , assuming that all circumradii in final triangulation are  $\ll 10^9$ . Time:  $\mathcal{O}\left(N^2 \log N\right)$ 

```
"DelaunayFast.h"
                                                      57c54d, 23 lines
// include inCircle from DelaunayFast
const T BIG = 1e9; // >> (10^4)^2
V<AR<int,3>> triIncrement(vP v) {
  v.pb({-BIG,-BIG}); v.pb({BIG,0}); v.pb({0,BIG});
  V<AR<int,3>> ret, tmp;
  ret.pb(\{sz(v)-3, sz(v)-2, sz(v)-1\});
  FOR(i, sz(v) - 3) {
   map<pi,int> m;
    each(a,ret) {
      if (inCircle(v[i], v[a[0]], v[a[1]], v[a[2]]))
       m[{a[0],a[1]}]++, m[{a[1],a[2]}]++, m[{a[0],a[2]}]++;
      else tmp.pb(a);
    each(a, m) if (a.s == 1) {
     AR<int,3> x{a.f.f,a.f.s,i};
      sor(x); tmp.pb(x);
   swap(ret,tmp); tmp.clear();
  each (a, ret) if (a[2] < sz(v)-3) tmp.pb(a);
  return tmp;
```

### DelaunavFast.h

Description: Fast Delaunay triangulation assuming no duplicates and not all points collinear (in latter case, result will be empty). Should work for doubles as well, though there may be precision issues in 'circ'. Returns triangles in ccw order. Each circumcircle will contain none of the input points. If coordinates are into at most B then T should be large enough to support ints on the order of  $B^4$ .

```
Time: \mathcal{O}(N \log N)
"Point.h"
                                                       0e7085, 82 lines
// using T = 11; (if coords are < 2e4)
using 111 = __int128;
// return true if p strictly within circumcircle(a,b,c)
bool inCircle(P p, P a, P b, P c) {
  a -= p, b -= p, c -= p; // assert(cross(a,b,c)>0);
  lll x = (lll) norm(a) *cross(b,c) + (lll) norm(b) *cross(c,a)
      +(111) norm(c) *cross(a,b);
  return x*(cross(a,b,c)>0?1:-1) > 0;
P arb(LLONG_MAX, LLONG_MAX); // not equal to any other point
using Q = struct Quad*;
struct Quad {
  bool mark; Q o, rot; P p;
  P F() { return r()->p; }
  Q r() { return rot->rot; }
  Q prev() { return rot->o->rot; }
  Q next() { return r()->prev(); }
Q makeEdge(P orig, P dest) {
  Q q[]{new Quad{0,0,0,oriq}, new Quad{0,0,0,arb},
      new Quad{0,0,0,dest}, new Quad{0,0,0,arb}};
  FOR(i, 4) q[i] -> o = q[-i \& 3], q[i] -> rot = q[(i+1) \& 3];
  return *q;
void splice(Q a, Q b) { swap(a->o->rot->o, b->o->rot->o); swap(
   \hookrightarrowa->o, b->o); }
Q connect(Q a, Q b) {
  Q = makeEdge(a->F(), b->p);
  splice(q, a->next()); splice(q->r(), b);
```

```
pair<Q,Q> rec(const vP& s) {
  if (sz(s) \le 3) {
    Q = makeEdge(s[0], s[1]), b = makeEdge(s[1], s.bk);
    if (sz(s) == 2) return { a, a->r() };
    splice(a->r(), b);
    auto side = cross(s[0], s[1], s[2]);
    Q c = side ? connect(b, a) : 0;
    return {side < 0 ? c -> r() : a, side < 0 ? c : b -> r() };
#define H(e) e \rightarrow F(), e \rightarrow p
#define valid(e) (cross(e->F(),H(base)) > 0)
  O A, B, ra, rb;
  int half = sz(s) / 2;
  tie(ra, A) = rec({all(s)-half});
  tie(B, rb) = rec({sz(s)-half+all(s)});
  while ((cross(B->p,H(A)) < 0 \&\& (A = A->next()))
       (cross(A->p,H(B)) > 0 && (B = B->r()->o)));
  O base = connect(B->r(), A);
  if (A->p == ra->p) ra = base->r();
  if (B->p == rb->p) rb = base;
#define DEL(e, init, dir) O e = init->dir; if (valid(e)) \
    while (inCircle(e->dir->F(), H(base), e->F())) { \
      0 t = e \rightarrow dir; \
      splice(e, e->prev()); \
      splice(e->r(), e->r()->prev()); \
      e = t; \
  while (1) {
    DEL(LC, base->r(), o); DEL(RC, base, prev());
    if (!valid(LC) && !valid(RC)) break;
    if (!valid(LC) || (valid(RC) && inCircle(H(RC), H(LC))))
      base = connect(RC, base->r());
    else base = connect(base->r(), LC->r());
  return {ra, rb};
V<AR<P,3>> triangulate(vP pts) {
  sor(pts); assert(unique(all(pts)) == end(pts)); // no
     \hookrightarrowduplicates
  if (sz(pts) < 2) return {};
  Q = rec(pts).f; V<Q> q = {e};
  while (cross(e->o->F(), e->F(), e->p) < 0) e = e->o;
#define ADD { Q c = e; do { c->mark = 1; pts.pb(c->p); \
  q.pb(c->r()); c = c->next(); } while (c != e); }
  ADD; pts.clear();
  int qi = 0; while (qi < sz(q)) if (!(e = q[qi++]) \rightarrow mark) ADD;
  V<AR<P,3>> ret(sz(pts)/3);
  FOR(i,sz(pts)) ret[i/3][i%3] = pts[i];
  return ret;
```

### ManhattanMST.h

**Description:** Given N points, returns up to 4N edges which are guaranteed to contain a MST for graph with edge weights w(p,q) = |p.x-q.x| + |p.y-q.y|. Edges are in the form {dist, {src, dst}}.

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

```
"DSU.h"
// use standard MST algorithm on result to find final MST
V<pair<int,pi>> manhattanMst(vpi v) {
 vi id(sz(v)); iota(all(id), 0);
 V<pair<int,pi>> ed;
 F0R(k, 4) {
    sort(all(id),[&](int i, int j) {
     return v[i].f+v[i].s < v[j].f+v[j].s; });</pre>
```

```
map<int,int> sweep; // find first octant neighbors
  each(i,id) { // those in sweep haven't found neighbor yet
    for (auto it = sweep.lb(-v[i].s);
     it != end(sweep); sweep.erase(it++)) {
      int j = it -> s;
      pi d{v[i].f-v[j].f,v[i].s-v[j].s};if (d.s>d.f)break;
      ed.pb({d.f+d.s,{i,j}});
    sweep[-v[i].s] = i;
  each(p,v) {
    if (k\&1) p.f *=-1;
    else swap(p.f,p.s);
return ed;
```

#### 8.5 3D

### Point3D.h

**Description:** Basic 3D geometry.

```
using P3 = AR<T,3>; using Tri = AR<P3,3>; using vP3 = V<P3>;
T norm(const P3& x) {
 T sum = 0; FOR(i,3) sum += sq(x[i]);
 return sum; }
T abs(const P3& x) { return sqrt(norm(x)); }
P3& operator+= (P3& 1, const P3& r) { F0R(i,3) 1[i] += r[i];
 return 1: }
P3& operator = (P3& 1, const P3& r) { F0R(i,3) 1[i] -= r[i];
P3& operator*=(P3& 1, const T& r) { F0R(i,3) 1[i] *= r;
 return 1; }
P3& operator/=(P3& 1, const T& r) { F0R(i,3) 1[i] /= r;
 return 1; }
P3 operator-(P3 1) { 1 *= -1; return 1; }
P3 operator+(P3 1, const P3& r) { return 1 += r; }
P3 operator-(P3 1, const P3& r) { return 1 -= r; }
P3 operator* (P3 1, const T& r) { return 1 *= r; }
P3 operator*(const T& r, const P3& 1) { return 1*r; }
P3 operator/(P3 1, const T& r) { return 1 /= r; }
P3 unit(const P3& x) { return x/abs(x); }
T dot(const P3& a, const P3& b) {
 T sum = 0; FOR(i,3) sum += a[i]*b[i];
 return sum; }
P3 cross(const P3& a, const P3& b) {
 return {a[1]*b[2]-a[2]*b[1],a[2]*b[0]-a[0]*b[2],
      a[0]*b[1]-a[1]*b[0]; }
P3 cross(const P3& a, const P3& b, const P3& c) {
 return cross(b-a,c-a); }
P3 perp(const P3& a, const P3& b, const P3& c) {
 return unit(cross(a,b,c)); }
bool isMult(const P3& a, const P3& b) { // for long longs
 P3 c = cross(a,b); FOR(i,sz(c)) if (c[i] != 0) return 0;
 return 1; }
bool collinear(const P3& a, const P3& b, const P3& c) {
 return isMult(b-a,c-a); }
T DC(const P3&a,const P3&b,const P3&c,const P3&p) {
 return dot(cross(a,b,c),p-a); }
bool coplanar(const P3&a,const P3&b,const P3&c,const P3&p) {
 return DC(a,b,c,p) == 0; }
bool op(const P3& a, const P3& b) {
```

### Hull3D PolySaVol Delaunay3 KMP Z

```
int ind = 0; // going in opposite directions?
  FOR(i,1,3) if (std::abs(a[i]*b[i])>std::abs(a[ind]*b[ind]))
  return a[ind] *b[ind] < 0;
// coplanar points, b0 and b1 on opposite sides of a0-a1?
bool opSide(const P3&a,const P3&b,const P3&c,const P3&d) {
  return op(cross(a,b,c),cross(a,b,d)); }
// coplanar points, is a in Triangle b
bool inTri(const P3& a, const Tri& b) {
  FOR(i,3) if (opSide(b[i],b[(i+1)%3],b[(i+2)%3],a)) return 0;
// point-seg dist
T psDist(const P3&p,const P3&a,const P3&b) {
  if (dot(a-p,a-b) \le 0) return abs(a-p);
  if (dot(b-p,b-a) \le 0) return abs(b-p);
  return abs(cross(p,a,b))/abs(a-b);
// projection onto line
P3 foot(const P3& p, const P3& a, const P3& b) {
 P3 d = unit(b-a); return a+dot(p-a,d)*d; }
// rotate p about axis
P3 rotAxis(const P3& p, const P3& a, const P3& b, T theta) {
  P3 dz = unit(b-a), f = foot(p,a,b);
  P3 dx = p-f, dy = cross(dz, dx);
  return f+cos(theta)*dx+sin(theta)*dy;
// projection onto plane
P3 foot(const P3& a, const Tri& b) {
  P3 c = perp(b[0],b[1],b[2]);
  return a-c*(dot(a,c)-dot(b[0],c)); }
// line-plane intersection
P3 lpIntersect(const P3&a0,const P3&a1,const Tri&b) {
  P3 c = unit(cross(b[2]-b[0],b[1]-b[0]));
  T \times = dot(a0,c)-dot(b[0],c), y = dot(a1,c)-dot(b[0],c);
  return (y*a0-x*a1)/(y-x);
```

### Hull3D.h

**Description:** Incremental 3D convex hull where not all points are coplanar. Normals to returned faces point outwards. If coordinates are into at most B then T should be large enough to support ints on the order of  $B^3$ . Changes order of points. The number of returned faces may depend on the random seed, because points that are on the boundary of the convex hull may or may not be included in the output.

Time:  $\mathcal{O}(N^2)$ ,  $\mathcal{O}(N \log N)$ 

5f4f0e, 91 lines

```
// using T = 11;
bool above (const P3&a, const P3&b, const P3&c, const P3&p) {
  return DC(a,b,c,p) > 0; } // is p strictly above plane
void prep(vP3& p) { // rearrange points such that
  shuffle(all(p),rng); // first four are not coplanar
  int dim = 1;
  FOR(i, 1, sz(p))
    if (dim == 1) {
      if (p[0] != p[i]) swap(p[1],p[i]), ++dim;
    } else if (dim == 2) {
      if (!collinear(p[0],p[1],p[i]))
        swap(p[2],p[i]), ++dim;
    } else if (dim == 3) {
      if (!coplanar(p[0],p[1],p[2],p[i]))
        swap(p[3], p[i]), ++dim;
  assert(dim == 4);
using F = AR<int,3>; // face
V<F> hull3d(vP3& p) {
```

```
// s.t. first four points form tetra
 prep(p); int N = sz(p); V<F> hull; // triangle for each face
 auto ad = [\&] (int a, int b, int c) { hull.pb(\{a,b,c\}); };
 // +new face to hull
 ad(0,1,2), ad(0,2,1); // initialize hull as first 3 points
 V<vb> in(N, vb(N)); // is zero before each iteration
 FOR(i,3,N) { // incremental construction
   V<F> def, HULL; swap(hull, HULL);
    // HULL now contains old hull
    auto ins = [&](int a, int b, int c) {
      if (in[b][a]) in[b][a] = 0; // kill reverse face
     else in[a][b] = 1, ad(a,b,c);
   };
    each(f, HULL) {
     if (above(p[f[0]],p[f[1]],p[f[2]],p[i]))
       FOR(j,3) ins(f[j],f[(j+1)%3],i);
        // recalc all faces s.t. point is above face
     else def.pb(f);
    each(t,hull) if (in[t[0]][t[1]]) // edge exposed,
     in[t[0]][t[1]] = 0, def.pb(t); // add a new face
    swap(hull,def);
 return hull;
V<F> hull3dFast(vP3& p) {
 prep(p); int N = sz(p); V<F> hull;
 vb active; // whether face is active
 V<vi> rvis; // points visible from each face
 V<AR<pi, 3>> other; // other face adjacent to each edge of
    \hookrightarrow face
 V<vi> vis(N); // faces visible from each point
 auto ad = [&](int a, int b, int c) {
   hull.pb({a,b,c}); active.pb(1); rvis.eb(); other.eb(); };
 auto ae = [&](int a, int b) { vis[b].pb(a), rvis[a].pb(b); };
 auto abv = [&](int a, int b) {
   F f=hull[a]; return above(p[f[0]],p[f[1]],p[f[2]],p[b]);};
 auto edge = [&](pi e) -> pi {
   return {hull[e.f][e.s],hull[e.f][(e.s+1)%3]}; };
 auto glue = [&] (pi a, pi b) { // link two faces by an edge
   pi x = edge(a); assert(edge(b) == mp(x.s, x.f));
   other[a.f][a.s] = b, other[b.f][b.s] = a;
 \}; // ensure face 0 is removed when i=3
 ad(0,1,2), ad(0,2,1); if (abv(1,3)) swap(p[1],p[2]);
 FOR(i,3) glue(\{0,i\},\{1,2-i\});
 FOR(i,3,N) ae(abv(1,i),i); // coplanar points go in rvis[\theta]
 vi label (N, -1);
 FOR(i,3,N) { // incremental construction
   vi rem; each(t,vis[i]) if (active[t]) active[t]=0, rem.pb(t
    if (!sz(rem)) continue; // hull unchanged
    int st = -1;
    each(r, rem) FOR(j, 3) {
     int o = other[r][j].f;
     if (active[o]) { // create new face!
        int a,b; tie(a,b) = edge(\{r,j\}); ad(a,b,i); st = a;
       int cur = sz(rvis)-1; label[a] = cur;
       vi tmp; set_union(all(rvis[r]),all(rvis[o]),
                 back_inserter(tmp));
        // merge sorted vectors ignoring duplicates
       each(x,tmp) if (abv(cur,x)) ae(cur,x);
       glue({cur,0},other[r][j]); // glue old w/ new face
    for (int x = st, y; x = y) { // glue new faces together
     int X = label[x]; qlue({X,1},{label[y=hull[X][1]],2});
     if (y == st) break;
```

```
V<F> ans; F0R(i,sz(hull)) if (active[i]) ans.pb(hull[i]);
return ans;
```

### PolySaVol.h

Description: surface area and volume of polyhedron, normals to faces must point outwards

```
"Hull3D.h"
                                                       52fc2b, 8 lines
pair<T,T> SaVol(vP3 p, V<F> faces) {
 T s = 0, v = 0;
 each(i,faces) {
    P3 a = p[i[0]], b = p[i[1]], c = p[i[2]];
    s += abs(cross(a,b,c)); v += dot(cross(a,b),c);
 return {s/2, v/6};
```

### Delaunav3.h

**Description:** Delaunay triangulation with 3D hull. Assumes no duplicate points and that not all points are collinear. If coordinates are ints at most B, T should be large enough to support ints on the order of  $B^4$ .

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

```
"Point.h", "Hull3D.h"
                                                       a4497e, 15 lines
V<AR<P,3>> triHull(vP p) {
 V < P3 > p3; V < AR < P, 3 > res; each(x,p) p3.pb({x.f,x.s,norm(x)});
 bool ok = 0; each(t,p3) ok |= !coplanar(p3[0],p3[1],p3[2],t);
 if (!ok) { // all points concyclic
    sort(1+all(p),[&p](P a, P b) {
      return cross(a-p.ft,b-p.ft)>0; });
    FOR(i, 1, sz(p)-1) res.pb({p.ft,p[i],p[i+1]});
 } else {
    #define nor(x) P(p3[x][0],p3[x][1])
    each(t,hull3dFast(p3))
      if (dot(cross(p3[t[0]],p3[t[1]],p3[t[2]]), \{0,0,1\}) < 0)
        res.pb(\{nor(t[0]), nor(t[2]), nor(t[1])\});
 return res;
```

# Strings (9)

## 9.1 Light

### KMP.h

**Description:** f[i] is length of the longest proper suffix of the *i*-th prefix of s that is a prefix of s

Time:  $\mathcal{O}(N)$ 4538e4, 13 lines

```
vi kmp(str s) {
 int N = sz(s); vi f(N+1); f[0] = -1;
  FOR(i, 1, N+1) {
    for (f[i]=f[i-1];f[i]!=-1&&s[f[i]]!=s[i-1];)f[i]=f[f[i]];
    ++f[i]; }
 return f;
vi getOc(str a, str b) { // find occurrences of a in b
 vi f = kmp(a+"@"+b), ret;
 FOR(i, sz(a), sz(b)+1) if (f[i+sz(a)+1] == sz(a))
   ret.pb(i-sz(a));
 return ret;
```

**Description:** f[i] is the max len such that s.substr(0,len) == s.substr(i,len)

27a566, 30 lines

```
Time: \mathcal{O}(N)
                                                       566170, 15 lines
vi z(str s) {
 int N = sz(s), L = 1, R = 0; s += '#';
  vi ans(N); ans[\theta] = N;
  FOR(i,1,N) {
    if (i \le R) ans [i] = min(R-i+1, ans[i-L]);
    while (s[i+ans[i]] == s[ans[i]]) ++ans[i];
    if (i+ans[i]-1 > R) L = i, R = i+ans[i]-1;
  return ans:
vi getPrefix(str a, str b) { // find prefixes of a in b
 vi t = z(a+b); t = vi(sz(a)+all(t));
 each(u,t) ckmin(u,sz(a));
  return t;
```

#### Manacher.h

Description: length of largest palindrome centered at each character of string and between every consecutive pair

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

fcc3f7, 13 lines

```
vi manacher(str _S) {
 str S = "@"; each(c,_S) S += c, S += "#";
  S.bk = '&';
  vi ans(sz(S)-1); int lo = 0, hi = 0;
  FOR(i,1,sz(S)-1) {
   if (i != 1) ans[i] = min(hi-i, ans[hi-i+lo]);
   while (S[i-ans[i]-1] == S[i+ans[i]+1]) ++ans[i];
   if (i+ans[i] > hi) lo = i-ans[i], hi = i+ans[i];
  ans.erase(begin(ans));
 FOR(i, sz(ans)) if (i%2 == ans[i]%2) ++ans[i];
  return ans:
```

### LyndonFactor.h

**Description:** A string is "simple" if it is strictly smaller than any of its own nontrivial suffixes. The Lyndon factorization of the string s is a factorization  $s = w_1 w_2 \dots w_k$  where all strings  $w_i$  are simple and  $w_1 \geq w_2 \geq \dots \geq w_k$ . Min rotation gets min index i such that cyclic shift of s starting at i is minimum.

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

af38ba, 19 lines

```
vs duval(str s) {
  int N = sz(s); vs factors;
  for (int i = 0; i < N; ) {
    int j = i+1, k = i;
    for (; j < N \&\& s[k] \le s[j]; ++j) {
     if (s[k] < s[j]) k = i;
    for (; i \le k; i += j-k) factors.pb(s.substr(i, j-k));
  return factors;
int minRotation(str s) {
  int N = sz(s); s += s;
  vs d = duval(s); int ind = 0, ans = 0;
  while (ans+sz(d[ind]) < N) ans += sz(d[ind++]);
  while (ind && d[ind] == d[ind-1]) ans -= sz(d[ind--]);
  return ans;
```

#### HashRange.h

Description: Polynomial hash for substrings with two bases. fc0b90, 24 lines

using H = AR<int,2>; // bases not too close to ends

```
H makeH(char c) { return {c,c}; }
uniform int distribution<int> BDIST(0.1*MOD, 0.9*MOD);
const H base{BDIST(rng),BDIST(rng)};
H operator+(H l, H r) {
 FOR(i,2) if ((l[i] += r[i]) >= MOD) l[i] -= MOD;
  return 1: }
H operator-(H l, H r) {
  FOR(i,2) if ((1[i] -= r[i]) < 0) 1[i] += MOD;
 return 1: }
H operator*(H l, H r) {
 FOR(i,2) 1[i] = (11)1[i] *r[i] %MOD;
  return 1; }
V<H> pows{{1,1}};
struct HashRange {
 str S; V<H> cum{{}};
  void add(char c) { S += c; cum.pb(base*cum.bk+makeH(c)); }
  void add(str s) { each(c,s) add(c); }
  void extend(int len) { while (sz(pows) <= len)</pre>
    pows.pb(base*pows.bk); }
  H hash(int 1, int r) { int len = r+1-1; extend(len);
    return cum[r+1]-pows[len]*cum[l]; }
```

#### ReverseBW.h

**Description:** Used only once. Burrows-Wheeler Transform appends # to a string, sorts the rotations of the string in increasing order, and constructs a new string that contains the last character of each rotation. This function reverses the transform.

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

e400d8, 7 lines

```
str reverseBW(str t) {
 vi nex(sz(t)); iota(all(nex),0);
 stable_sort(all(nex),[&t](int a,int b){return t[a]<t[b];});</pre>
 str ret; for (int i = nex[0]; i;)
   ret += t[i = nex[i]];
 return ret:
```

### AhoCorasickFixed.h

Description: Aho-Corasick for fixed alphabet. For each prefix, stores link to max length suffix which is also a prefix.

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

96dfcc, 27 lines

```
template<size_t ASZ> struct ACfixed {
 struct Node { AR<int, ASZ> to; int link; };
 V<Node> d{{}};
 int add(str s) { // add word
   int v = 0:
   each(C,s) {
     int c = C-'a';
     if (!d[v].to[c]) d[v].to[c] = sz(d), d.eb();
     v = d[v].to[c];
   return v;
 void init() { // generate links
   d[0].link = -1;
   queue<int> q; q.push(0);
   while (sz(q)) {
     int v = q.ft; q.pop();
     FOR(c, ASZ) {
       int u = d[v].to[c]; if (!u) continue;
       d[u].link = d[v].link == -1 ? 0 : d[d[v].link].to[c];
       q.push(u);
     if (v) FOR(c,ASZ) if (!d[v].to[c])
       d[v].to[c] = d[d[v].link].to[c];
```

```
};
```

### SuffixArrav.h

**Description:** Sort suffixes. First element of sa is sz(S), isa is the inverse of sa, and 1cp stores the longest common prefix between every two consecutive elements of sa.

Time:  $\mathcal{O}(N \log N)$ "RMQ.h"

```
struct SuffixArray {
 str S; int N; vi sa, isa, lcp;
 void init(str \_S) { N = sz(S = \_S)+1; genSa(); genLcp(); }
 void genSa() { // sa has size sz(S)+1, starts with sz(S)
    sa = isa = vi(N); sa[0] = N-1; iota(1+all(sa),0);
    sort(1+all(sa),[&](int a, int b) { return S[a] < S[b]; });</pre>
    FOR(i,1,N) { int a = sa[i-1], b = sa[i];
      isa[b] = i > 1 \&\& S[a] == S[b] ? isa[a] : i; }
    for (int len = 1; len < N; len *= 2) { // currently sorted
      // by first len chars
      vi s(sa), is(isa), pos(N); iota(all(pos),\theta);
      each(t,s) {int T=t-len; if (T>=0) sa[pos[isa[T]]++] = T;}
      FOR(i, 1, N)  { int a = sa[i-1], b = sa[i];
        isa[b] = is[a] == is[b] &&is[a+len] == is[b+len]?isa[a]:i;
  void genLcp() { // Kasai's Algo
    lcp = vi(N-1); int h = 0;
    FOR(b, N-1)  { int a = sa[isa[b]-1];
      while (a+h < sz(S) \&\& S[a+h] == S[b+h]) ++h;
      lcp[isa[b]-1] = h; if (h) h--; }
    R.init(lcp);
  RMO<int> R:
  int getLCP(int a, int b) { // lcp of suffixes starting at a,b
    if (a == b) return sz(S)-a;
    int l = isa[a], r = isa[b]; if (l > r) swap(l,r);
    return R.query(1,r-1);
};
```

### SuffixArrayLinear.h

**Description:** Linear-time suffix array.

Usage: sa\_is(s, 26) // all entries must be in [0, 26) **Time:** O(N), ~100ms for  $N = 5 \cdot 10^5$ 

```
ed0bb4, 46 lines
vi sa_is(const vi& s, int upper) {
 int n = sz(s); if (!n) return {};
 vi sa(n); vb ls(n);
 ROF(i, n-1) ls[i] = s[i] == s[i+1] ? ls[i+1] : s[i] < s[i+1];
 vi sum_l(upper), sum_s(upper);
 FOR(i,n) (ls[i] ? sum_l[s[i]+1] : sum_s[s[i]])++;
 F0R(i,upper) {
    if (i) sum_l[i] += sum_s[i-1];
    sum_s[i] += sum_l[i];
 auto induce = [&](const vi& lms) {
    fill(all(sa),-1);
    vi buf = sum s;
    for (int d: lms) if (d != n) sa[buf[s[d]] ++] = d;
    buf = sum_1; sa[buf[s[n-1]]++] = n-1;
    F0R(i,n) {
      int v = sa[i]-1;
      if (v >= 0 \&\& !ls[v]) sa[buf[s[v]]++] = v;
    buf = sum_1;
    R0F(i,n) {
     int v = sa[i]-1;
     if (v >= 0 \&\& ls[v]) sa[--buf[s[v]+1]] = v;
```

### TandemRepeats PalTree SuffixAutomaton SuffixTree

```
};
  vi lms_map(n+1,-1), lms; int m = 0;
  FOR(i,1,n) if (!ls[i-1] \&\& ls[i]) lms_map[i]=m++, lms.pb(i);
  induce(lms); // sorts LMS prefixes
  vi sorted_lms;each(v,sa)if (lms_map[v]!=-1)sorted_lms.pb(v);
  vi rec_s(m); int rec_upper = 0; // smaller subproblem
  FOR(i,1,m) { // compare two lms substrings in sorted order
   int l = sorted_lms[i-1], r = sorted_lms[i];
    int end 1 = 1 \text{ms map}[1] + 1 < m ? 1 \text{ms}[1 \text{ms map}[1] + 1] : n;
    int end_r = lms_map[r]+1 < m ? <math>lms[lms_map[r]+1] : n;
   bool same = 0; // whether lms substrings are same
    if (end 1-1 == end r-r) {
     for (; 1 < end_1 && s[1] == s[r]; ++1, ++r);
     if (1 != n \&\& s[1] == s[r]) same = 1;
   rec_s[lms_map[sorted_lms[i]]] = (rec_upper += !same);
  vi rec_sa = sa_is(rec_s,rec_upper+1);
  FOR(i,m) sorted lms[i] = lms[rec sa[i]];
  induce(sorted_lms); // sorts LMS suffixes
  return sa:
TandemRepeats.h
Description: Find all (i, p) such that s.substr(i,p) ==
Usage: solve("aaabababa") // {{0, 1, 1}, {2, 5, 2}}
Time: \mathcal{O}(N \log N)
                                                      661326, 13 lines
```

s.substr(i+p,p). No two intervals with the same period intersect

```
"SuffixArray.h"
V<AR<int,3>> solve(str s) {
 int N = sz(s); SuffixArray A,B;
  A.init(s); reverse(all(s)); B.init(s);
  V<AR<int,3>> runs;
  for (int p = 1; 2*p \le N; ++p) { // do in O(N/p) for period p
    for (int i = 0, lst = -1; i+p \le N; i += p) {
     int 1 = i-B.getLCP(N-i-p,N-i), r = i-p+A.getLCP(i,i+p);
      if (1 > r \mid \mid 1 == 1st) continue;
      runs.pb(\{lst = l,r,p\}); // for each i in [l,r],
    } // s.substr(i,p) == s.substr(i+p,p)
  return runs;
```

### 9.2 Heavy

#### PalTree.h

Description: Used infrequently. Palindromic tree computes number of occurrences of each palindrome within string. ans[i][0] stores min even xsuch that the prefix s[1..i] can be split into exactly x palindromes, ans [i] [1] does the same for odd x.

```
Time: \mathcal{O}\left(N\sum\right) for addChar, \mathcal{O}\left(N\log N\right) for updAns
```

```
8a7d31, 41 lines
```

```
struct PalTree {
  static const int ASZ = 26;
  struct node {
    AR < int, ASZ > to = AR < int, ASZ > ();
    int len, link, oc = 0; // # occurrences of pal
    int slink = 0, diff = 0;
   AR<int,2> seriesAns;
   node(int _len, int _link) : len(_len), link(_link) {}
  str s = "@"; V<AR<int, 2>> ans = {{0, MOD}};
 V < node > d = \{\{0,1\}, \{-1,0\}\}; // dummy pals of len 0,-1
  int last = 1;
  int getLink(int v) {
```

```
while (s[sz(s)-d[v].len-2] != s.bk) v = d[v].link;
  return v;
void updAns() { // serial path has O(log n) vertices
  ans.pb({MOD,MOD});
  for (int v = last; d[v].len > 0; v = d[v].slink) {
    d[v].seriesAns=ans[sz(s)-1-d[d[v].slink].len-d[v].diff];
    if (d[v].diff == d[d[v].link].diff)
      FOR(i,2) ckmin(d[v].seriesAns[i],
            d[d[v].link].seriesAns[i]);
    // start of previous oc of link[v]=start of last oc of v
    FOR(i,2) ckmin(ans.bk[i],d[v].seriesAns[i^1]+1);
void addChar(char C) {
  s += C; int c = C-'a'; last = getLink(last);
  if (!d[last].to[c]) {
    d.eb(d[last].len+2,d[getLink(d[last].link)].to[c]);
    d[last].to[c] = sz(d)-1;
    auto& z = d.bk; z.diff = z.len-d[z.link].len;
    z.slink = z.diff == d[z.link].diff
     ? d[z.link].slink : z.link;
  } // max suf with different dif
  last = d[last].to[c]; ++d[last].oc;
  updAns();
void numOc() { ROF(i,2,sz(d)) d[d[i].link].oc += d[i].oc; }
```

#### SuffixAutomaton.h

Description: Used infrequently. Constructs minimal deterministic finite automaton (DFA) that recognizes all suffixes of a string. len corresponds to the maximum length of a string in the equivalence class, pos corresponds to the first ending position of such a string, 1nk corresponds to the longest suffix that is in a different class. Suffix links correspond to suffix tree of the reversed string!

```
Time: \mathcal{O}(N \log \Sigma)
```

```
struct SuffixAutomaton {
 int N = 1; vi lnk\{-1\}, len\{0\}, pos\{-1\}; // suffix link,
 // max length of state, last pos of first occurrence of state
 V<map<char,int>> nex{1}; V<bool> isClone{0};
 // transitions, cloned -> not terminal state
 V<vi> iLnk; // inverse links
 int add(int p, char c) { // \sim p \text{ nonzero if } p != -1
   auto getNex = [&]() {
     if (p == -1) return 0;
     int q = nex[p][c]; if (len[p]+1 == len[q]) return q;
     int clone = N++; lnk.pb(lnk[q]); lnk[q] = clone;
     len.pb(len[p]+1), nex.pb(nex[q]),
     pos.pb(pos[q]), isClone.pb(1);
     for (; \sim p \&\& nex[p][c] == q; p = lnk[p]) nex[p][c]=clone;
     return clone;
    // if (nex[p].count(c)) return getNex();
    // ^ need if adding > 1 string
   int cur = N++; // make new state
   lnk.eb(), len.pb(len[p]+1), nex.eb(),
   pos.pb(pos[p]+1), isClone.pb(0);
   for (; ~p && !nex[p].count(c); p = lnk[p]) nex[p][c] = cur;
   int x = getNex(); lnk[cur] = x; return cur;
 void init(str s) { int p = 0; each(x,s) p = add(p,x); }
 // inverse links
 void genIlnk() {iLnk.rsz(N); FOR(v,1,N)iLnk[lnk[v]].pb(v);}
 // APPLICATIONS
 void getAllOccur(vi& oc, int v) {
   if (!isClone[v]) oc.pb(pos[v]); // terminal position
   each(u,iLnk[v]) getAllOccur(oc,u); }
```

```
vi allOccur(str s) { // get all occurrences of s in automaton
    int cur = 0;
    each(x,s) {
     if (!nex[cur].count(x)) return {};
      cur = nex[cur][x]; }
    // convert end pos -> start pos
    vi oc; getAllOccur(oc,cur); each(t,oc) t += 1-sz(s);
    sort(all(oc)); return oc;
 vl distinct;
 11 getDistinct(int x) {
    // # distinct strings starting at state x
    if (distinct[x]) return distinct[x];
    distinct[x]=1;each(y,nex[x]) distinct[x]+=getDistinct(y.s);
    return distinct[x]; }
 11 numDistinct() { // # distinct substrings including empty
    distinct.rsz(N); return getDistinct(0); }
 11 numDistinct2() { // assert(numDistinct() == numDistinct2());
    ll ans = 1; FOR(i,1,N) ans += len[i]-len[lnk[i]];
    return ans: }
};
SuffixAutomaton S;
vi sa; str s;
void dfs(int x) {
 if (!S.isClone[x]) sa.pb(sz(s)-1-S.pos[x]);
 V<pair<char,int>> chr;
  each(t, S.iLnk[x]) chr.pb({s[S.pos[t]-S.len[x]],t});
 sort(all(chr)); each(t,chr) dfs(t.s);
int main() {
 re(s); reverse(all(s));
 S.init(s); S.genIlnk();
  dfs(0); ps(sa); // generating suffix array for s
```

### SuffixTree.h

a99c6d, 67 lines

Description: Used infrequently. Ukkonen's algorithm for suffix tree. Longest non-unique suffix of s has length len[p]+lef after each call to add terminates. Each iteration of loop within add decreases this quantity by one. Time:  $\mathcal{O}(N \log \Sigma)$ 

```
39751c, 51 lines
struct SuffixTree {
 str s; int N = 0;
 vi pos, len, lnk; V<map<char,int>> to;
 int make (int POS, int LEN) { // lnk[x] is meaningful when
    // x!=0 and len[x] != MOD
    pos.pb(POS);len.pb(LEN);lnk.pb(-1);to.eb();return N++; }
 void add(int& p, int& lef, char c) { // longest
    // non-unique suffix is at node p with lef extra chars
    s += c; ++lef; int lst = 0;
    for (; lef; p?p=lnk[p]: lef--) { // if p != root then <math>lnk[p]
      // must be defined
     while (lef>1 && lef>len[to[p][s[sz(s)-lef]]])
       p = to[p][s[sz(s)-lef]], lef -= len[p];
      // traverse edges of suffix tree while you can
     char e = s[sz(s)-lef]; int& q = to[p][e];
      // next edge of suffix tree
     if (!q) q = make(sz(s)-lef, MOD), lnk[lst] = p, lst = 0;
      // make new edge
     else {
       char t = s[pos[q]+lef-1];
       if (t == c) { lnk[lst] = p; return; } // suffix not
          ∽unique
       int u = make(pos[q],lef-1);
        // new node for current suffix-1, define its link
       to[u][c] = make(sz(s)-1,MOD); to[u][t] = q;
        // new, old nodes
```

### CircularLCS SMAWK FastIO

```
pos[q] += lef-1; if (len[q] != MOD) len[q] -= lef-1;
       q = u, lnk[lst] = u, lst = u;
  void init(str _s) {
   make (-1, 0); int p = 0, lef = 0;
   each(c,_s) add(p,lef,c);
   add(p,lef,'$'); s.pop_back(); // terminal char
  int maxPre(str x) { // max prefix of x which is substring
    for (int p = 0, ind = 0;;) {
     if (ind == sz(x) || !to[p].count(x[ind])) return ind;
     p = to[p][x[ind]];
     FOR(i, len[p]) {
       if (ind == sz(x) || x[ind] != s[pos[p]+i]) return ind;
  vi sa; // generate suffix array
  void genSa(int x = 0, int Len = 0) {
   if (!sz(to[x])) sa.pb(pos[x]-Len); // found terminal node
   else each(t,to[x]) genSa(t.s,Len+len[x]);
};
```

# Various (10)

### 10.1 Dynamic programming

When doing DP on intervals:  $a[i][j] = \min_{i < k < j} (a[i][k] + a[k][j]) + f(i, j),$  where the (minimal) optimal k increases with both i and j,

- one can solve intervals in increasing order of length, and search k = p[i][j] for a[i][j] only between p[i][j-1]and p[i+1][j].
- This is known as Knuth DP. Sufficient criteria for this are if  $f(b,c) \leq f(a,d)$  and  $f(a,c) + f(b,d) \le f(a,d) + f(b,c)$  for all  $a \le b \le c \le d$ .
- Consider also: LineContainer (ch. Data structures), monotone queues, ternary search.

### CircularLCS.h

**Description:** Used only twice. For strs A, B calculates longest common subsequence of A with all rotations of B

```
Time: \mathcal{O}(|A| \cdot |B|)
                                                         db21cf, 26 lines
int circular_lcs(str A, str B) {
 B += B;
  int max_lcs = 0;
  V < vb > dif_left(sz(A)+1, vb(sz(B)+1)), dif_up(sz(A)+1, vb(sz(B)))
     \hookrightarrow+1));
  auto recalc = [&](int x, int y) { assert(x && y);
    int res = (A.at(x-1) == B.at(y-1))
      dif_up[x][y-1] \mid dif_left[x-1][y];
    dif_left[x][y] = res-dif_up[x][y-1];
   dif_up[x][y] = res-dif_left[x-1][y];
  FOR(i,1,sz(A)+1) FOR(j,1,sz(B)+1) recalc(i,j);
```

```
FOR(j, sz(B)/2) {
  // 1. zero out dp[.][j], update dif_left and dif_right
  if (j) for (int x = 1, y = j; x \le sz(A) && y \le sz(B); ) {
    int pre_up = dif_up[x][y];
    if (y == j) dif_up[x][y] = 0;
    else recalc(x,y);
    (pre\_up == dif\_up[x][y]) ? ++x : ++y;
  // 2. calculate LCS(A[0:sz(A)),B[j:j+sz(B)/2))
  int cur lcs = 0;
  FOR(x,1,sz(A)+1) cur_lcs += dif_up[x][j+sz(B)/2];
  ckmax(max_lcs,cur_lcs);
return max_lcs;
```

#### SMAWK.h

Description: Given negation of totally monotone matrix with entries of type D, find indices of row maxima (their indices increase for every submatrix). If tie, take lesser index. f returns matrix entry at (r,c) in O(1). Use in place of divide & conquer to remove a log factor.

**Time:**  $\mathcal{O}(R+C)$ , can be reduced to  $\mathcal{O}(C(1+\log R/C))$  evaluations of f

```
template<class F, class D=11> vi smawk (F f, vi x, vi y) {
 vi ans(sz(x),-1); // x = rows, y = cols
 \#define upd() if (ans[i] == -1 || w > mx) ans[i] = c, mx = w
 if (\min(sz(x), sz(y)) \le 8) {
   FOR(i,sz(x)) { int r = x[i]; D mx;
     each(c, y) \{ D w = f(r, c); upd(); \} \}
   return ans:
 if (sz(x) < sz(y)) { // reduce subset of cols to consider
   vi Y; each(c,y) {
     for (; sz(Y); Y.pop\_back()) \{ int X = x[sz(Y)-1]; \}
       if (f(X,Y.bk) >= f(X,c)) break; }
     if (sz(Y) < sz(x)) Y.pb(c);
 } // recurse on half the rows
 vi X; for (int i = 1; i < sz(x); i += 2) X.pb(x[i]);
 vi ANS = smawk(f,X,y); FOR(i,sz(ANS)) ans[2*i+1] = ANS[i];
 for (int i = 0, k = 0; i < sz(x); i += 2) {
   int to = i+1 < sz(ans) ? ans[i+1] : y.bk; D mx;
   for(int r = x[i];;++k) {
     int c = y[k]; D w = f(r,c); upd();
     if (c == to) break; }
 return ans;
```

#### Debugging tricks 10.2

- signal(SIGSEGV, [](int) { \_Exit(0); }); converts segfaults into Wrong Answers. Similarly one can catch SIGABRT (assertion failures) and SIGFPE (zero divisions). \_GLIBCXX\_DEBUG violations generate SIGABRT (or SIGSEGV on gcc 5.4.0 apparently).
- feenableexcept (29); kills the program on NaNs (1), 0-divs (4), infinities (8) and denormals (16).

### 10.3 Optimization tricks

### 10.3.1 Bit hacks

- x & -x is the least bit in x.
- for (int x = m; x;) { --x &= m; ... } loops over all subset masks of m (except m itself).
- c = x&-x, r = x+c;  $(((r^x) >> 2)/c) | r$ is the next number after x with the same number of bits set.
- FOR(b,k) FOR(i,1<<K) if (i&1<<b) D[i] += D[i^(1<<b)]; computes all sums of subsets.

### 10.3.2 Pragmas

- #pragma GCC optimize ("Ofast") will make GCC auto-vectorize for loops and optimizes floating points better (assumes associativity and turns off denormals).
- #pragma GCC target ("avx,avx2") can double performance of vectorized code, but causes crashes on old machines. Also consider older #pragma GCC target ("sse4").
- #pragma GCC optimize ("trapv") kills the program on integer overflows (but is really slow).

```
FastIO.h
```

```
Description: Fast input and output.
Usage: initO(); int a,b; ri(a,b); wi(b,'
n'); wi(a,'
Time: input is \sim 300 \text{ms} faster for 10^6 long longs on CF
                                                       cbf279, 39 lines
inline namespace FastIO {
const int BSZ = 1<<15; ///// INPUT</pre>
char ibuf[BSZ]; int ipos, ilen;
char nc() { // next char
 if (ipos == ilen) {
    ipos = 0; ilen = fread(ibuf,1,BSZ,stdin);
    if (!ilen) return EOF;
 return ibuf[ipos++];
void rs(str& x) { // read str
 char ch; while (isspace(ch = nc()));
 do { x += ch; } while (!isspace(ch = nc()) && ch != EOF);
tcT> void ri(T& x) { // read int or 11
 char ch; int sgn = 1;
 while (!isdigit(ch = nc())) if (ch == '-') sgn *= -1;
```

x = ch'' 0'; while (isdigit (ch = nc())) x = x\*10+(ch'' 0');

tcT, class... Ts> void ri(T& t, Ts&... ts) {

ri(t); ri(ts...); } // read ints ///// OUTPUT (call initO() at start)

char obuf[BSZ], numBuf[100]; int opos;

```
void flushOut() { fwrite(obuf,1,opos,stdout); opos = 0; }
void wc(char c) { // write char
   if (opos == BSZ) flushOut();
   obuf[opos++] = c; }
void ws(str s) { each(c,s) wc(c); } // write str
tcT> void wi(T x, char after = '\0') {
   if (x < 0) wc('-'), x *= -1;
   int len = 0; for (;x>=10;x/=10) numBuf[len++] = '0'+(x%10);
   wc('0'+x); R0F(i,len) wc(numBuf[i]);
   if (after) wc(after);
}
void initO() { assert(atexit(flushOut) == 0); }
}
```

### 10.4 Other languages

### Python3.py

**Description:** Solution to CF Factorisation Collaboration. Demonstrates usage of Decimal.

47 lines

```
from math import *
import sys, random
def nextInt():
 return int(input())
def nextStrs():
 return input().split()
def nextInts():
 return list(map(int,nextStrs()))
n = nextInt()
v = [n]
def process(x):
  global v
  x = abs(x)
  V = []
  for t in v:
    g = gcd(t, x)
    if g != 1:
      V.append(g)
    if g != t:
      V.append(t//g)
for i in range (50):
  x = random.randint(0, n-1)
  if gcd(x,n) != 1:
    process(x)
  else:
    sx = x * x % n \# assert(gcd(sx,n) == 1)
    print(f"sqrt {sx}")
    sys.stdout.flush()
    X = nextInt()
    process(x+X)
    process(x-X)
print(f'! {len(v)}',end='')
for i in v:
  print(f' {i}',end='')
sys.stdout.flush() # sys.exit(0) -> exit
# sys.setrecursionlimit(int(1e9)) -> stack size
# print(f'{ans:=.6f}') -> print ans to 6 decimal places
from decimal import * # arbitrary precision decimals
ctx = getcontext()
ctx.prec = 28
print(Decimal(1) / Decimal(7)) # 0.1428571428571428571428571429
print(ctx.power(Decimal(10),-30)) # 1E-30
```