

# Yandex. Algorithm 2011 < br > Round 2

# A. Reflection

time limit per test: 2 seconds memory limit per test: 256 megabytes input: standard input output: standard output

For each positive integer n consider the integer  $\psi(n)$  which is obtained from n by replacing every digit a in the decimal notation of n with the digit (9 - a). We say that  $\psi(n)$  is the *reflection* of n. For example, reflection of n should be omitted. So reflection of n equals n0, reflection of n1 equals n2.

Let us call the *weight* of the number the product of the number and its reflection. Thus, the weight of the number 10 = 890.

Your task is to find the maximum weight of the numbers in the given range [I, r] (boundaries are included).

#### Input

Input contains two space-separated integers l and r ( $1 \le l \le r \le 10^9$ ) — bounds of the range.

# **Output**

Output should contain single integer number: maximum value of the product  $n \cdot \psi(n)$ , where  $l \le n \le r$ .

Please, do not use %lld specificator to read or write 64-bit integers in C++. It is preferred to use cout (also you may use %I64d).

# **Examples**

input	
3 7	
output	
20	

input	
1 1	
output	
8	

input
8 10
output
890

#### Note

In the third sample weight of 8 equals  $8 \cdot 1 = 8$ , weight of 9 equals  $9 \cdot 0 = 0$ , weight of 10 equals 890.

Thus, maximum value of the product is equal to 890.

# B. Tetris revisited

time limit per test: 1 second memory limit per test: 256 megabytes input: standard input output: standard output

Physicist Woll likes to play one relaxing game in between his search of the theory of everything.

Game interface consists of a rectangular  $n \times m$  playing field and a dashboard. Initially some cells of the playing field are filled while others are empty. Dashboard contains images of all various connected (we mean connectivity by side) figures of 2, 3, 4 and 5 cells, with all their rotations and reflections. Player can copy any figure from the dashboard and place it anywhere at the still empty cells of the playing field. Of course any figure can be used as many times as needed.

Woll's aim is to fill the whole field in such a way that there are no empty cells left, and also... just have some fun.

Every initially empty cell should be filled with exactly one cell of some figure. Every figure should be entirely inside the board.

In the picture black cells stand for initially filled cells of the field, and one-colour regions represent the figures.

## Input

First line contains integers n and m ( $1 \le n, m \le 1000$ ) — the height and the width of the field correspondingly. Next n lines contain m symbols each. They represent the field in a natural way: j-th character of the i-th line is "#" if the corresponding cell is filled, and "." if it is empty.

#### **Output**

If there is no chance to win the game output the only number "-1" (without the quotes). Otherwise output any filling of the field by the figures in the following format: each figure should be represented by some digit and figures that touch each other by side should be represented by distinct digits. Every initially filled cell should be represented by "#".

# Examples input

2 3

#.#
output
000 #0#
input
3 3 .#.
 #
output
5#1 511 55#
input
3 3
 .## .#.
output
-1
input

#### **Note**

##

1 2 ## output

In the third sample, there is no way to fill a cell with no empty neighbours.

In the forth sample, Woll does not have to fill anything, so we should output the field from the input.

# C. Genetic engineering

time limit per test: 2 seconds memory limit per test: 256 megabytes input: standard input

output: standard output

"Multidimensional spaces are completely out of style these days, unlike genetics problems" — thought physicist Woll and changed his subject of study to bioinformatics. Analysing results of sequencing he faced the following problem concerning DNA sequences. We will further think of a DNA sequence as an arbitrary string of uppercase letters "A", "C", "G" and "T" (of course, this is a simplified interpretation).

Let W be a long DNA sequence and  $S_1, S_2, ..., S_m$  — collection of short DNA sequences. Let us say that the collection filters W iff W can be covered with the sequences from the collection. Certainly, substrings corresponding to the different positions of the string may intersect or even cover each other. More formally: denote by |W| the length of W, let symbols of W be numbered from W to W. Then for each position W in W there exist pair of indices W, W is such that the substring W in W is equals one of the elements W, W, W, W of the collection.

Woll wants to calculate the number of DNA sequences of a given length filtered by a given collection, but he doesn't know how to deal with it. Help him! Your task is to find the number of different DNA sequences of length n filtered by the collection  $\{S_i\}$ .

Answer may appear very large, so output it modulo 100000009.

#### Input

First line contains two integer numbers n and m ( $1 \le n \le 1000$ ,  $1 \le m \le 10$ ) — the length of the string and the number of sequences in the collection correspondently.

Next m lines contain the collection sequences  $S_i$ , one per line. Each  $S_i$  is a nonempty string of length not greater than 10. All the strings consist of uppercase letters "A", "C", "G", "T". The collection may contain identical strings.

#### **Output**

Output should contain a single integer — the number of strings filtered by the collection modulo  $100000009 (10^9 + 9)$ .

#### **Examples**

input	
2 1	
output	
output	

input	
6 2 CAT TACT	
output	
2	

#### Note

In the first sample, a string has to be filtered by "A". Clearly, there is only one such string: "AA".

In the second sample, there exist exactly two different strings satisfying the condition (see the pictures below).



# D. Powerful array

time limit per test: 5 seconds memory limit per test: 256 megabytes input: standard input

output: standard output

An array of positive integers  $a_1, a_2, ..., a_n$  is given. Let us consider its arbitrary subarray  $a_l, a_{l+1}..., a_r$ , where  $1 \le l \le r \le n$ . For every positive integer S denote by  $K_S$  the number of occurrences of S into the subarray. We call the *power* of the subarray the sum of products  $K_S \cdot K_S \cdot S$  for every positive integer S. The sum contains only finite number of nonzero summands as the number of different values in the array is indeed finite.

You should calculate the power of t given subarrays.

#### Input

First line contains two integers n and t ( $1 \le n$ ,  $t \le 200000$ ) — the array length and the number of queries correspondingly.

Second line contains n positive integers  $a_i$  ( $1 \le a_i \le 10^6$ ) — the elements of the array.

Next t lines contain two positive integers l, r ( $1 \le l \le r \le n$ ) each — the indices of the left and the right ends of the corresponding subarray.

#### **Output**

Output t lines, the i-th line of the output should contain single positive integer — the power of the i-th query subarray.

Please, do not use %lld specificator to read or write 64-bit integers in C++. It is preferred to use cout stream (also you may use %I64d).

#### **Examples**

```
input

3 2
1 2 1
1 2 1
1 3

output

3 6
```

```
input

8 3
1 1 2 2 1 3 1 1
2 7
1 6
2 7

output

20
20
20
```

#### Note

Consider the following array (see the second sample) and its [2, 7] subarray (elements of the subarray are colored):

Then  $K_1=3$ ,  $K_2=2$ ,  $K_3=1$ , so the power is equal to  $3^2 \cdot 1 + 2^2 \cdot 2 + 1^2 \cdot 3 = 20$ .

# E. Long sequence

time limit per test: 2 seconds memory limit per test: 256 megabytes

input: standard input output: standard output

A sequence  $a_0, a_1, \ldots$  is called a *recurrent binary sequence*, if each term  $a_i$  ( $i = 0, 1, \ldots$ ) is equal to 0 or 1 and there exist coefficients  $c_1, c_2, \ldots, c_k \in \{0,1\}$  such that

$$a_n = c_1 \cdot a_{n-1} + c_2 \cdot a_{n-2} + \dots + c_k \cdot a_{n-k} \pmod{2},$$

for all  $n \ge k$ . Assume that not all of  $C_i$  are zeros.

Note that such a sequence can be uniquely recovered from any k-tuple  $\{a_s, a_{s+1}, ..., a_{s+k-1}\}$  and so it is periodic. Moreover, if a k-tuple contains only zeros, then the sequence contains only zeros, so this case is not very interesting. Otherwise the minimal period of the sequence is not greater than  $2^k - 1$ , as k-tuple determines next element, and there are  $2^k - 1$  non-zero k-tuples. Let us call a sequence *long* if its minimal period is exactly  $2^k - 1$ . Your task is to find a long sequence for a given k, if there is any.

## Input

Input contains a single integer k ( $2 \le k \le 50$ ).

#### Output

If there is no long sequence for a given K, output "-1" (without quotes). Otherwise the first line of the output should contain K integer numbers:  $C_1, C_2, ..., C_K$  (coefficients). The second line should contain first K elements of the sequence:  $A_0, A_1, ..., A_{K-1}$ . All of them (elements and coefficients) should be equal to 0 or 1, and at least one  $C_i$  has to be equal to 1.

If there are several solutions, output any.

# **Examples**

input	
2	
output	
1 1 1 0	

## input

3

# output

0 1 1 1 1 1

# Note

1. In the first sample:  $c_1 = 1$ ,  $c_2 = 1$ , so  $a_n = a_{n-1} + a_{n-2} \pmod{2}$ . Thus the sequence will be:

so its period equals  $3 = 2^2 - 1$ .

2. In the second sample:  $c_1 = 0$ ,  $c_2 = 1$ ,  $c_3 = 1$ , so  $a_n = a_{n-2} + a_{n-3} \pmod{2}$ . Thus our sequence is:

and its period equals  $7 = 2^3 - 1$ .

1 1 1 0 0 1 0 1 1 1 0 0 1 0 ...

101101101...

Periods are colored.