P17

DNA (Deoxyribonucleic Acid) is the molecule which contains the genetic instructions. It consists of four different nucleotides, namely Adenine, Thymine, Guanine, and Cytosine as shown in Figure 1. If we represent a nucleotide by its initial character, a DNA strand can be regarded as a long string (sequence of characters) consisting of the four characters A, T, G, and C. For example, assume we are given some part of a DNA strand which is composed of the following sequence of nucleotides:

"Thymine-Adenine-Cytosine-Thymine-Guanine-Cytosine-Cytosine-Guanine-Adenine-Thymine"

Then we can represent the above DNA strand with the string "TAACTGCCGAT."

The biologist Prof. Ahn found that a gene X commonly exists in the DNA strands of five different kinds of an imals, namely dogs, cats, horses, cows, and monkeys. He also discovered that the DNA sequences of the gene X from each animal were very alike. See Figure 2.

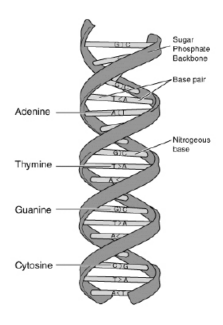


Figure 1.

	DNA sequence of gene X
Cat:	GCATATGGCTGTGCA
Dog:	GCAAATGGCTGTGCA
Horse:	GCTAATGGGTGTCCA
Cow:	GCAAATGGCTGTGCA
Monkey:	GCAAATCGGTGAGCA

Figure 2. DNA sequences of gene X in five animals.

Prof. Ahn thought that humans might also have the gene X and decided to search for the DNA sequence of X in human DNA. However, before searching, he should define a representative DNA sequence of gene X because its sequences are not exactly the same in the DNA of the five animals. He decided to use the Hamming distance to define the representative sequence.

The Hamming distance is the number of different characters at each position from two strings of equal length. For example, assume we are given the two strings "AGCAT" and "GGAAT." The Hamming distance of these two strings is 2 because the 1st and the 3rd characters of the two strings are different. Using the Hamming distance, we can define a representative string for a set of multiple strings of equal length. Given a set of strings $S = \{s_1, \ldots, s_m\}$ of length n, the consensus error between a string y of length n and the set S is the sum of the Hamming distances between y and each s_i in S. If the consensus error between y and S is the minimum among all possible strings S of length S is called a consensus string of S. For example, given the three strings "AGCAT" "AGACT" and "GGAAT" the consensus string of the given strings is "AGAAT" because the sum of the Hamming distances between "AGAAT" and the three strings is 3 which is minimal. (In this case, the consensus string is unique, but in general, there can be more than one consensus string.) We use the consensus string as a representative of the DNA sequence. For the example of Figure 2 above, a consensus string of gene S is "GCAAATGGCTGTGCA" and the consensus error is 7.

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Input

Your program is to read from standard input. The input consists of T test cases. The number of test cases T is given in the first line of the input. Each test case starts with a line containing two integers m and n which are separated by a single space. The integer m ($4 \le m \le 50$) represents the number of DNA sequences and n ($4 \le n \le 1000$) represents the length of the DNA sequences, respectively. In each of the next m lines, each DNA sequence is given.

Output

Your program is to write to standard output. Print the consensus string in the first line of each case and the consensus error in the second line of each case. If there exists more than one consensus string, print the lexicographically smallest consensus string.

Sample Input

3

5 8

TATGATAC

TAAGCTAC

AAAGATCC

TGAGATAC

TAAGATGT

4 10

ACGTACGTAC

CCGTACGTAG

GCGTACGTAT

TCGTACGTAA

6 10

ATGTTACCAT

AAGTTACGAT

AACAAAGCAA

AAGTTACCTT

AAGTTACCAA

TACTTACCAA

Sample Output

TAAGATAC

7

ACGTACGTAA

6

AAGTTACCAA

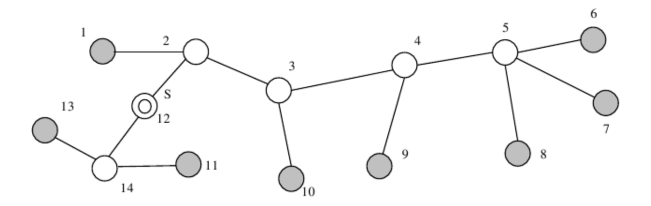
12

P18

Consider a tree network with n nodes where the internal nodes correspond to servers and the terminal nodes correspond to clients. The nodes are numbered from 1 to n. Among the servers, there is an original server S which provides VOD (Video On Demand) service. To ensure the quality of service for the clients, the distance from each client to the VOD server S should not exceed a certain value k. The distance from a node u to a node v in the tree is defined to be the number of edges on the path from u to v. If there is a nonempty subset C of clients such that the distance from each u in C to S is greater than k, then replicas of the VOD system have to be placed in some servers so that the distance from each client to the nearest VOD server (the original VOD system or its replica) is k or less.

Given a tree network, a server S which has VOD system, and a positive integer k, find the minimum number of replicas necessary so that each client is within distance k from the nearest server which has the original VOD system or its replica.

For example, consider the following tree network.



In the above tree, the set of clients is $\{1, 6, 7, 8, 9, 10, 11, 13\}$, the set of servers is $\{2, 3, 4, 5, 12, 14\}$, and the original VOD server is located at node 12.

For k = 2, the quality of service is not guaranteed with one VOD server at node 12 because the clients in $\{6, 7, 8, 9, 10\}$ are away from VOD server at distance > k. Therefore, we need one or more replicas. When one replica is placed at node 4, the distance from each client to the nearest server of $\{12, 4\}$ is less than or equal to 2. The minimum number of the needed replicas is one for this example.

Input

Your program is to read the input from standard input. The input consists of T test cases. The number of test cases (T) is given in the first line of the input. The first line of each test case contains an integer n ($3 \le n \le 1,000$) which is the number of nodes of the tree network. The next line contains two integers s ($1 \le s \le n$) and k ($k \ge 1$) where s is the VOD server and k is the distance value for ensuring the quality of service. In the following n-1 lines, each line contains a pair of nodes which represent an edge of the tree network.

Output

Your program is to write to standard output. Print exactly one line for each test case. The line should contain an integer that is the minimum number of the needed replicas.

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Sample Input

2 14

12 2

1 2

2 3

3 4

4 5

5 6

7 5

8 5

4 9

10 3

2 12

12 14

13 14

14 11

14

3 4

1 2

2 3

3 4

4 5

5 6

7 5

8 5

4 9

10 3

2 12

12 14

13 14

14 11

Sample Output

1

0