

It is 2178, and alien life has been discovered on a distant planet. There seems to be only one species on the planet and they do not reproduce as animals on Earth do. Even more amazing, the genetic makeup of every single organism is identical!

The genetic makeup of each organism is a single sequence of nucleotides. The nucleotides come in three types, denoted by 'A' (Adenine), 'C' (Cytosine), and 'M' (Muamine). According to one hypothesis, evolution on this planet occurs when a new nucleotide is inserted somewhere into the genetic sequence of an existing organism. If this change is evolutionarily advantageous, then organisms with the new sequence quickly replace ones with the old sequence.

It was originally thought that the current species evolved this way from a single, very simple organism with a single-nucleotide genetic sequence, by way of mutations as described above. However, fossil evidence suggests that this might not have been the case. Right now, the research team you are working with is trying to validate the concept of "parallel evolution" — that there might actually have been two evolutionary paths evolving in the fashion described above, and eventually both paths evolved to the single species present on the planet today. Your task is to verify whether the parallel evolution hypothesis is consistent with the genetic material found in the fossil samples gathered by your team.

Input

The input file contains several test cases, each of them as described below.

The input begins with a number n ($1 \leq n \leq 4000$) denoting the number of nucleotide sequences found in the fossils. The second line describes the nucleotide sequence of the species currently living on the planet. Each of the next n lines describes one nucleotide sequence found in the fossils.

Each nucleotide sequence consists of a string of at least one but no more than 4 000 letters. The strings contain only upper-case letters 'A', 'C', and 'M'. All the nucleotide sequences, including that of the currently live species, are distinct.

Output

For each test case, display an example of how the nucleotide sequences in the fossil record participate in two evolutionary paths. The example should begin with one line containing two integers s_1 and s_2 , the number of nucleotide sequences in the fossil record that participate in the first path and second path, respectively. This should be followed by s_1 lines containing the sequences attributed to the first path, in chronological order (from the earliest), and then s_2 lines containing the sequences attributed to the second path, also in chronological order. If there are multiple examples, display any one of them. If it is possible that a sequence could appear in the genetic history of both species, your example should assign it to exactly one of the evolutionary paths.

If it is impossible for all the fossil material to come from two evolutionary paths, display the word 'impossible'.

Sample Input

```
5
AACCMMAA
ACA
MM
ACMAA
AA
A
3
ACMA
ACM
ACA
AMA
1
AM
MA
4
AAAAAA
AA
AAA
A
AAAAA
```

Sample Output

```
1 4
MM
A
AA
ACA
ACMAA
impossible
impossible
0 4
A
AA
AAA
AAAAA
```