## 1224 - DNA Prefix

Given a set of **n** DNA samples, where each sample is a string containing characters from {**A**, **C**, **G**, **T**}, we are trying to find a subset of samples in the set, where the length of the longest common prefix multiplied by the number of samples in that subset is maximum.

To be specific, let the samples be:

ACGT ACGTGCGT ACCGTGC ACGCCGT

If we take the subset {ACGT} then the result is 4 (4 \* 1), if we take {ACGT, ACGTGCGT, ACGCCGT} then the result is 3 \* 3 = 9 (since ACG is the common prefix), if we take {ACGT, ACGTGCGT, ACCGTGC, ACGCCGT} then the result is 2 \* 4 = 8.

Now your task is to report the maximum result we can get from the samples.

## Input

Input starts with an integer  $T \leq 10$ , denoting the number of test cases.

Each case starts with a line containing an integer n ( $1 \le n \le 50000$ ) denoting the number of DNA samples. Each of the next n lines contains a non empty string whose length is not greater than 50. And the strings contain characters from  $\{A, C, G, T\}$ .

## Output

For each case, print the case number and the maximum result that can be obtained.

Sample Input	Output for Sample Input
3	Case 1: 9
4	Case 2: 66
ACGT	Case 3: 20
ACGTGCGT	
ACCGTGC	
ACGCCGT	
3	
CGCGCGCGCGCCCCGCCCC	
CGCGCGCGCCCCGCCCCAC	
CGCGCGCGCCCCCCCCCCCC	
2	
CGCGCCGCGCGCGCGC	
GGCGCCGCGCGCGCTC	

## **Note**

Dataset is huge. Use faster I/O methods.