1334 - Genes in DNA

Your friend is a biologist. He has just sequenced a DNA and wants to know about contribution of different genes in that DNA. Both Gene and DNA can be represented by a sequence of letters or strings. Given the sequence of a DNA **D** and a Gene **G**; your friend uses following method to calculate the contribution.

- Generate a list P of proper non-empty prefixes of G and another S of proper non-empty suffixes of G [1]. Additionally let the L is list of all strings that is concatenation of a prefix and a suffix. So if G = ACCT then P = A, AC, ACC and S = T, CT, CCT and L = AT, ACT, ACCT, ACCT, ACCCT, ACCCT, ACCCT. If |G| = n then it is obvious that size of L is (n 1)².
- 2. For each element of **L**, count number of times it occurs as substring in **D**. Contribution of Gene **G** in DNA **D** is total of these values. For example if **D** = **ACTACCTACCCT** then

AT	0
ACT	1
ACCT	1
ACT	1
ACCT	1
ACCCT	0
ACCT	1
ACCCT	0
ACCCCT	1
Total	6

As this process is very clumsy he wants to automate this process. As he is not a programmer, he needs your help. He will be very grateful if you kindly write him a program which will read the sequence of the DNA and the Gene, and will calculate contribution of the Gene in the DNA.

Input

Input starts with an integer $T \leq 20$, denoting the number of test cases.

Each case contains two lines. The first line contains a string denoting the sequence of DNA, and the second line contains another string denoting the Gene. The length of each string is less than **50000** and consists of only **A**, **C**, **T** and **G**.

Output

For each case, print the case number and the contribution, as described above.

Sample Input	Output for Sample Input
3	Case 1: 6
ACTACCTACCCCT	Case 2: 4
ACCT	Case 3: 8
AAA	
AAAA	
AAAA	
AAA	

Note

- 1. Proper prefix (suffix) of a string $\bf S$ is a prefix (suffix) of length smaller than $|\bf S|$. Here $|\bf S|$ denotes length of $\bf S$.
- 2. Dataset is huge, use faster I/O methods.