A gene is represented as a string of length n (where n is divisible by 4), composed of the letters n, nT, and G. It is considered to be *steady* if each of the four letters occurs exactly  $\frac{n}{4}$  times. For example, **GACT** and **AAGTGCCT** are both steady genes.

Bear Limak is a famous biotechnology scientist who specializes in modifying bear DNA to make it steady. Right now, he is examining a gene represented as a string *gene*. It is not necessarily steady. Fortunately, Limak can choose one (maybe empty) substring of *gene* and replace it with any string of the same length.

Modifying a large substring of bear genes can be dangerous. Given a string *gene*, can you help Limak find the length of the smallest possible substring that he can replace to make *gene* a steady gene?

Note: A substring of a string  $\boldsymbol{s}$  is a subsequence made up of zero or more contiguous characters of  $\boldsymbol{s}$ .

As an example, consider gene = ACTGAAAG. The substring AA just before or after G can be replaced with CT or TC. One selection would create ACTGACTG.

## **Function Description**

Complete the **steadyGene** function in the editor below. It should return an integer that represents the length of the smallest substring to replace.

steadyGene has the following parameter:

• gene: a string

## **Input Format**

The first line contains an interger n divisible by 4, that denotes the length of a string *gene*. The second line contains a string gene of length n.

### **Constraints**

- $4 \le n \le 500000$  n is divisible by 4
- $gene[i] \in [CGAT]$

## **Subtask**

• 4 < n < 2000 in tests worth 30% points.

#### **Output Format**

Print the length of the minimum length substring that can be replaced to make *gene* stable.

# **Sample Input**

GAAATAAA

## **Sample Output**

5

## **Explanation**

One optimal solution is to replace **AAATA** with **TTCCG** resulting in **GTTCCGAA**. The replaced substring has length **5**.