

Examples:

Input: str = "01010101010"

Output: Yes

Input: str = "REC101"

Output: No

Input	Result
01010101010	Yes
010101 10101	No

Ex. No. : 8.1 Date:

Register No.: 230701092 Name:

Binary String

Coders here is a simple task for you, Given string str. Your task is to check whether it is a binary string or not by using python set.

```
s=input()
s1=set(s)
if (s1=={'0','1'}):
    print("Yes")
else:
    print("No")
```

Examples:

Input: t = (5, 6, 5, 7, 7, 8), K = 13

Output: 2 Explanation:

Pairs with sum K(=13) are $\{(5, 8), (6, 7), (6, 7)\}.$

Therefore, distinct pairs with sum K(=13) are $\{(5, 8), (6, 7)\}$.

Therefore, the required output is 2.

Input	Result
1,2,1,2,5	1
1,2	0

Ex. No. : 8.2 Date:

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Check Pair

Given a tuple and a positive integer k, the task is to find the count of distinct pairs in the tuple whose sum is equal to K.

```
def find_pairs_with_sum(numbers, target_sum):
  numbers_list = list(numbers)
  pairs = set()
  visited = set()
  for number in numbers list:
     complement = target_sum - number
     if complement in visited:
       pair = tuple(sorted((number, complement)))
       pairs.add(pair)
     visited.add(number)
  return pairs
numbers_input = input("")
target_sum = int(input(""))
numbers = tuple(map(int, numbers_input.split(',')))
pairs = find_pairs_with_sum(numbers, target_sum)
print(f"{len(pairs)}")
```

Input: s = "AAAAACCCCCCAAAAACCCCCCAAAAAGGGTTT"

Output: ["AAAAACCCCC","CCCCCAAAAA"]

Example 2:

Input: s = "AAAAAAAAAAA"

Output: ["AAAAAAAAAA"]

Input	Result
AAAAACCCCCAAAAACCCCCCAAAAAGGGTTT	AAAAACCCCC CCCCAAAAA

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DNA Sequence

The **DNA sequence** is composed of a series of nucleotides abbreviated as 'A', 'C', 'G', and 'T'.

For example, "ACGAATTCCG" is a **DNA sequence**.

When studying **DNA**, it is useful to identify repeated sequences within the DNA.

Given a string s that represents a **DNA sequence**, return all the **10-letter-long** sequences (substrings) that occur more than once in a DNA molecule. You may return the answer in **any order**.

```
def findRepeatedDnaSequences(s):
  if len(s) < 10:
     return []
  sequences = {}
  result = []
  for i in range(len(s) - 9):
     substring = s[i:i+10]
     if substring in sequences:
        sequences[substring] += 1
     else:
        sequences[substring] = 1
  for sequence, count in sequences.items():
     if count > 1:
        result.append(sequence)
  for sequence, count in sequences.items():
        if count > 1:
          result.append(sequence)
     for i in result:
        print(i)
  s1=input()
  findRepeatedDnaSequences(s1)
```

Input: nums = [1,3,4,2,2]

Output: 2

Example 2:

Input: nums = [3,1,3,4,2]

Output: 3

Input	Result
1 3 4 4 2	4

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Print repeated no

Given an array of integers nums containing n + 1 integers where each integer is in the range [1, n] inclusive. There is only **one repeated number** in nums, return this repeated number. Solve the problem using <u>set</u>.

```
def find_duplicate(nums):
    seen = set()
    for num in nums:
        if num in seen:
            return num
        seen.add(num)

# Input
I = input()
nums = list(map(int, I.split()))

# Output
print(find_duplicate(nums))
```

Sample Input:

5 4

12865

26810

Sample Output:

1 5 10

3

Sample Input:

5 5

 $1\ 2\ 3\ 4\ 5$

 $1\ 2\ 3\ 4\ 5$

Sample Output:

NO SUCH ELEMENTS

Input	Result
5 4 1 2 8 6 5 2 6 8 10	1 5 10 3

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Remove repeated

Write a program to eliminate the common elements in the given 2 arrays and print only the non-repeating elements and the total number of such non-repeating elements.

Input Format:

The first line contains space-separated values, denoting the size of the two arrays in integer format respectively.

The next two lines contain the space-separated integer arrays to be compared.

```
a=[int(i) for i in input().split()]
n1=[int(i) for i in input().split()]
n2=[int(i) for i in input().split()]
com=set(n1)&set(n2)
uni=set(n1)|set(n2)
tot=sorted(uni-com)
for i in tot:
    print(i,end=' ')
print('\n',end='')
print(len(tot))
```

Input: text = "hello world", brokenLetters = "ad"

Output:

1

Explanation: We cannot type "world" because the 'd' key is broken.

Input	Result
hello world ad	1

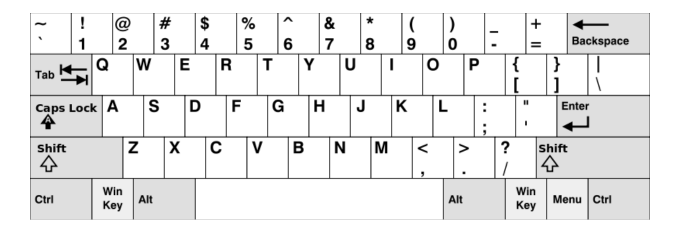
Ex. No.	:	8.6	Date:
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Malfunctioning Keyboard

There is a malfunctioning keyboard where some letter keys do not work. All other keys on the keyboard work properly.

Given a string text of words separated by a single space (no leading or trailing spaces) and a string brokenLetters of all distinct letter keys that are broken, return the number of words in text you can fully type using this keyboard.

```
a=[i for i in input().split()]
k=list(input())
s=set()
for i in a:
    n=[j for j in i]
    m=[z for z in k if z in n]
    s.update(m)
print(len(s))
```



Input: words = ["Hello","Alaska","Dad","Peace"]

Output: ["Alaska","Dad"]

Example 2:

Input: words = ["omk"]

Output: [] Example 3:

Input: words = ["adsdf","sfd"]
Output: ["adsdf","sfd"]

Input	Result
4 Hello Alaska Dad Peace	Alaska Dad

Ex. No. : 8.7 Date:

Register No.: Name:

American keyboard

Given an array of strings words, return the words that can be typed using letters of the alphabet on only one row of American keyboard like the image below.

In the American keyboard:

- the first row consists of the characters "qwertyuiop",
- the second row consists of the characters "asdfghjkl", and
- the third row consists of the characters "zxcvbnm".

```
n=int(input())
f=0
a=[input() for i in range(n)]
I1=['qwertyuiop','asdfghjkl','zxcvbnm']
I=[[i for i in i] for i in I1]
for i in a:
   n=[i for i in i.lower()]
  #print(sorted(set(I[1])|set(n))==sorted(set(I[1])))
   #print(set(I[1]),set(n))
   if set(n)|set(I[0]) = set(I[0]):
     f=1
      print(i)
      continue
   elif set(n)|set(I[1]) == set(I[1]):
     f=1
      print(i)
      continue
   elif set(n)|set(I[2]) == set(I[2]):
     f=1
      print(i)
      continue
if not f:
   print('No words')
```

