

# Counting Point Mutations

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## Problem

Given two strings  $s$  and  $t$  of equal length, the **Hamming distance** between  $s$  and  $t$ , denoted  $d_H(s, t)$ , is the number of corresponding symbols that differ in  $s$  and  $t$ .

- **Given:** Two DNA strings  $s$  and  $t$  of equal length (not exceeding 1 kbp).
- **Return:** The Hamming distance  $d_H(s, t)$

## Sample Dataset

```
GAGCCTACTAACGGGAT
CATCGTAATGACGGCCT
```

## Sample Output

```
7
```

## Intuition

The Hamming distance function requires two strings of equal length, thus we first assert that  $|s| = |t|$  where  $|s|$  denotes the length of string  $s$  and  $|t|$  the length of string  $t$ . We initialize a variable `mismatches` to zero; this will be used for keeping track of the number of mismatches as we compare the base of each string at a specific index. As with other string algorithms, we iterate over the indices from 0 up to (but not including)  $|s|$ . If  $s_i = t_i$ , the value of `mismatches` is incremented by one. Otherwise we proceed to the following index for the next comparison.

## Solution

```
def hamming_distance(s: str, t: int) -> int:
    """Compute the number of index-specific mismatches between two strings."""
    assert len(s) == len(t), "Both strings must be of equal length"
    mismatches = 0
    for i in range(len(s)):
        if s[i] != t[i]:
            mismatches += 1
    return mismatches
```

```
s = "GAGCCTACTAACGGGAT"
t = "CATCGTAATGACGGCCT"
result = hamming_distance(s, t)
print(result)
```

```
7
```

## Bibliography