

Bioinformatique

Manipulations de séquences ADN, ARN, protéines,...

Compter les nucléotides d'une séquence ADN

```
<sxh python; title : Counting_DNA_Nucleotides-01.py> #!/usr/bin/env python # -*- coding: utf-8 -*-  
""" On dispose d'un exemple de chaîne ADN (constituée des symboles 'A', 'C', 'G', 'T') Le programme  
utilise plusieurs techniques pour donner les nombres d'occurrences respectifs des différentes bases  
""" adn =
```

```
"AGCTTTTCATTCTGACTGCAACGGGCAATATGTCTCTGTGTGGATTAAAAAAGAGTGTCTGATAGCAGC"
```

```
# utilisation d'une liste et de la méthode .count() bases=["A","C","G","T"] for base in bases:
```

```
    print adn.count(base),
```

```
print
```

```
# Variante : for c in 'ACGT':
```

```
    print adn.count(c),
```

```
print
```

```
# variante un peu moins lisible out = [] for c in 'ACGT':
```

```
    out.append(str(adn.count(c)))
```

```
print(' '.join(out))
```

```
# utilisation de la technique "list comprehension" count=[adn.count(c) for c in 'ACGT'] for val in  
count:
```

```
    print val,
```

```
print
```

```
# autre "list comprehension", avec impression formatée → version "one line" print "%d %d %d %d" %  
tuple([adn.count(X) for X in "ACGT"])
```

```
# count "à la main", sans utilisation de fonctions/librairie ACGT = "ACGT" count = [0,0,0,0] for c in  
adn:
```

```
    for i in range(len(ACGT)):  
        if c == ACGT[i]:  
            count[i] +=1
```

for val in count:

```
print val,
```

print

count "à la main", avec .index() ACGT = "ACGT" count = [0,0,0,0] for c in adn:

```
count[ACGT.index(c)] += 1
```

for val in count:

```
print val,
```

print

utilisation de la librairie collections from collections import defaultdict ncount = defaultdict(int) for c in adn:

```
ncount[c] += 1
```

print ncount['A'], ncount['C'], ncount['G'], ncount['T']

collections.Counter from collections import Counter for k,v in sorted(Counter(adn).items()): print v, print

avec un dictionnaire freq = {'A': 0, 'C': 0, 'G': 0, 'T': 0} for c in adn:

```
freq[c] += 1
```

print freq['A'], freq['C'], freq['G'], freq['T']

avec un dictionnaire et count(), impression différente dico={} for base in bases:

```
dico[base]=adn.count(base)
```

for key,val in dico.items():

```
print "{} = {}".format(key, val)
```

</sxh>

Références

- [Using biological databases to teach evolution and biochemistry](#)
- [Rosalind](#), plateforme d'apprentissage de la programmation en bioinformatique
- [GenBank](#)
- [Biopython](#)

- <https://en.wikipedia.org/wiki/Bioinformatics>
- https://en.wikipedia.org/wiki/Open_Bioinformatics_Foundation
- https://en.wikipedia.org/wiki/FASTA_format
- https://en.wikipedia.org/wiki/List_of_open-source_bioinformatics_software
- <http://www.amberbiology.com/>, "Python For The Life Sciences. A gentle introduction to Python for life scientists" (à paraître)

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