## Python 5

## **Dictionaries**

Dictionaries are another iterable, like a string and list. Unlike strings and lists, dictionaries are not a sequence, or in other words, they are **unordered** and the position is not important.

Dictionaries are a collection of key/value pairs. In Python, each key is separated from its value by a colon (:), the items are separated by commas, and the whole thing is enclosed in curly braces. An empty dictionary without any items is written with just two curly braces, like this: {}

Each key in a dictionary is unique, while values may not be. The values of a dictionary can be of any type, but the keys must be of an immutable data type such as strings, numbers, or tuples.

Data that is appropriate for dictionaries are two pieces of information that naturally go together, like gene name and sequence.

Key	Value
TP53	GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC
BRCA1	GTACCTTGATTTCGTATTCTGAGAGGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA

#### **Creating a Dictionary**

```
genes = { 'TP53' :
'GATGGGATTGGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC' , 'BRCA1' :
'GTACCTTGATTTCGTATTCTGAGAGGCTGCTTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA' }
```

Breaking up the key/value pairs over multiple lines make them easier to read.

#### **Accessing Values in Dictionaries**

To retrieve a single value in a dictionary use the value's key in this format dict[key]. This will return the value at the specified key.

```
>>> genes = { 'TP53' :
'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC' , 'BRCA1' :
'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCCTTGGTTTCCGTGGCAACGGAAAA' }
>>>
    genes['TP53']
GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC
```

The sequence of the gene TP53 is stored as a value of the key 'TP53'. We can access the sequence by using the key in this format dict[key]

The value can be accessed and passed directly to a function or stored in a variable.

```
>>> print(genes['TP53'])
GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC
>>>
>>> seq = genes['TP53']
>>> print(seq)
GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC
```

#### **Changing Values in a Dictionary**

Individual values can be changed by using the key and the assignment operator.

```
>>> genes = { 'TP53' :
    'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC' , 'BRCA1' :
    'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA' }
>>> print(genes)
{'BRCA1': 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA',
    'TP53': 'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC'}
>>> genes['TP53'] = 'atg'
>>> print(genes)
{'BRCA1': 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA',
    'TP53': 'atg'}
```

The contents of the dictionary have changed.

Other assignment operators can also be used to change a value of a dictionary key.

```
>>> genes = { 'TP53' :
'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC' , 'BRCA1' :
'GTACCTTGATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA' }
>>>
>>> genes['TP53'] +=
'TAGAGCCACCGTCCAGGGAGCAGGTAGCTGCTGGGCTCCGGGGACACTTTGCGTTCGGGCTGGGAGCGTG'
>>>
>>> print(genes)
{'BRCA1': 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA',
'TP53':
'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTCTAGAGCCACCGTCCAG
GGAGCAGGTAGCTGCTGGGGCTCCGGGGACACTTTGCGTTCGGGCAGCGTG'}
```

#### **Accessing Each Dictionary Key/Value**

Since a dictionary is an iterable object, we can iterate through its contents.

A for loop can be used to retrieve each key of a dictionary one a time:

```
>>> for gene in genes:
... print(gene)
...
TP53
BRCA1
```

Once you have the key you can retrieve the value:

```
>>> for gene in genes:
... seq = genes[gene]
... print(gene, seq[0:10])
...
TP53 GATGGGATTG
BRCA1 GTACCTTGAT
```

## **Building a Dictionary one Key/Value at a Time**

Building a dictionary one key/value at a time is akin to what we just saw when we change a key's value. Normally you won't do this. We'll talk about ways to build a dictionary from a file in a later lecture.

```
>>> genes = {}
>>> print(genes)
{}
>>> genes['Brca1'] =
'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA'
>>> genes['TP53'] =
'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC'
>>> print(genes)
{'Brca1': 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTCTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA',
'TP53': 'GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC'}
```

We start by creating an empty dictionary. Then we add each key/value pair using the same syntax as when we change a value.

dict[key] = new\_value

#### **Checking That Dictionary Keys Exist**

Python generates an error (NameError) if you try to access a key that does not exist.

```
>>> print(genes['HDAC'])
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
NameError: name 'HDAC' is not defined
```

#### **Dictionary Operators**

Operator	Description	
in	key in dict returns True if the key exists in the dictionary	
not in	key not in dict returns True if the key does not exist in the dictionary	

Because Python generates a NameError if you try to use a key that doesn't exist in the dictionary, you need to check whether a key exists before trying to use it.

The best way to check whether a key exists is to use in

```
>>> gene = 'TP53'
>>> if gene in genes:
... print('found')
...
found
>>>
>>> if gene in genes:
... print(genes[gene])
...
GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCAAGACTGGCGCTAAAAGTTTTGAGCTTCTCAAAAGTC
>>>
```

#### Building a Dictionary one Key/Value at a Time using a loop

Now we have all the tools to build a dictionary one key/value using a for loop. This is how you will be building dictionaries more often in real life.

Here we are going to count and store nucleotide counts:

```
#!/usr/bin/env python3
# create a new empty dictionary
nt_count={}
# loop example from loops lecture
dna = 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA'
for nt in dna:
  # is this nt in our dictionary?
  if nt in nt count:
    # if it is, lets increment our count
    previous_count = nt_count[nt]
    new_count = previous_count + 1
    nt count[nt] = new count
  else:
    # if not, lets add this nt to our dictionary and make count = 1
    nt_count[nt] = 1;
print(nt_count)
```

```
{'G': 20, 'T': 21, 'A': 13, 'C': 16}
```

What is another way we could increment our count?

```
nt_count={}

dna = 'GTACCTTGATTTCGTATTCTGAGAGGCTGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA'

for nt in dna:
   if nt in nt_count:
      nt_count[nt] += 1
   else:
      nt_count[nt] = 1;

print(nt_count)
```

remember that count=count+1 is the same as count+=1

#### **Sorting Dictionary Keys**

If you want to print the contents of a dictionary, you should sort the keys then iterate over the keys with a for loop. Why do you want to sort the keys?

This will print keys in the same order every time you run your script. Dictionaries are unordered, so without sorting, you'll get a different order every time you run the script, which could be confusing.

### **Dictionary Functions**

Function	Description
len(dict)	returns the total number of key/value pairs
str(dict)	returns a string representation of the dictionary
type(variable)	Returns the type or class of the variable passed to the function. If the variable is dictionary, then it would return a dictionary type.

These functions work on several other data types too!

#### **Dictionary Methods**

Method	Description
dict.clear()	Removes all elements of dictionary dict
dict.copy()	Returns a shallow copy of dictionary dict. <u>Shallow vs. deep</u> copying only matters in multidimensional data structures.
<pre>dict.fromkeys(seq,value)</pre>	Create a new dictionary with keys from seq (Python sequence type) and values set to value.
dict.items()	Returns a list of (key, value) tuple pairs
dict.pop(key)	Removes the key:value pair and returns the value
dict.keys()	Returns list of keys
<pre>dict.get(key, default = None)</pre>	get value from dict[key], use default if not present
<pre>dict.setdefault(key, default = None)</pre>	Similar to get(), but will set dict[key] = default if key is not already in dict
dict.update(dict2)	Adds dictionary dict2's key-values pairs to dict
dict.values()	Returns list of dictionary dict's values

#### Sets

A set is another Python data type. It is essentially a dictionary with keys but no values.

- A set is unordered
- A set is a collection of data with no duplicate elements.
- Common uses include looking for differences and eliminating duplicates in data sets.

Curly braces {} or the set() function can be used to create sets.

Note: to create an empty set you have to use <code>set()</code>, not <code>{}</code> the latter creates an empty dictionary.

```
>>> basket = {'apple', 'orange', 'apple', 'pear', 'orange', 'banana'}
>>> print(basket)
{'orange', 'banana', 'pear', 'apple'}
```

Look, duplicates have been removed

Test to see if an value is in the set

```
>>> 'orange' in basket
True
>>> 'crabgrass' in basket
False
```

The in operator works the same with sets as it does with lists and dictionaries

Union, intersection, difference and symmetric difference can be done with sets

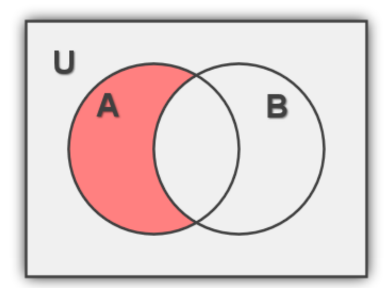
```
>>> a = set('abracadabra')
>>> b = set('alacazam')
>>> a
{'a', 'r', 'b', 'c', 'd'}
```

Sets contain unique elements, therefore, even if duplicate elements are provided they will be removed.

#### **Set Operators**

#### Difference

The difference between two sets are the elements that are unique to the set to the left of the — operator, with duplicates removed.

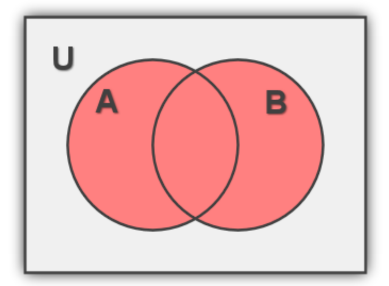


```
>>> a = set('abracadabra')
>>> b = set('alacazam')
>>> a - b
{'r', 'd', 'b'}
```

This results the letters that are in a but not in b

#### Union

The union between two sets is a sequence of the all the elements of the first and second sets combined, with duplicates removed.

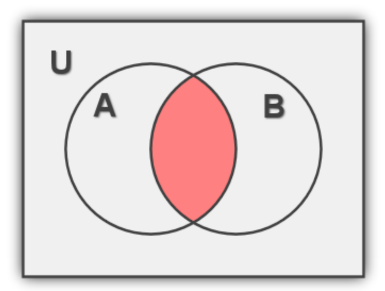


```
>>> a = set('abracadabra')
>>> b = set('alacazam')
>>> a | b
{'a', 'c', 'r', 'd', 'b', 'm', 'z', 'l'}
```

This returns letters that are in a or b both

#### Intersection

The intersection between two sets is a sequence of the elements which are in both sets, with duplicates removed.

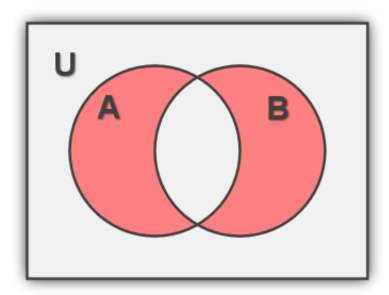


```
>>> a = set('abracadabra')
>>> b = set('alacazam')
>>> a & b
{'a', 'c'}
```

This returns letters that are in both a and b

#### **Symmetric Difference**

The symmetric difference is the elements that are only in the first set plus the elements that are only in the second set, with duplicates removed.



```
>>> a = set('abracadabra')
>>> b = set('alacazam')
>>> a ^ b
{'r', 'd', 'b', 'm', 'z', 'l'}
```

This returns the letters that are in a or b but not in both (also known as exclusive or)

#### **Set Functions**

Function	Description	
all()	returns True if all elements of the set are true (or if the set is empty).	
any()	returns True if any element of the set is true. If the set is empty, return False.	
returns an enumerate object. It contains the index and value of all the items of as a pair.		
len()	len() returns the number of items in the set.	
max() returns the largest item in the set.		
min()	returns the smallest item in the set.	
sorted()	returns a new sorted list from elements in the set (does not alter the original set).	
sum()	returns the sum of all elements in the set.	

#### **Set Methods**

Method	Description
set.add(new)	adds a new element
set.clear()	remove all elements
set.copy()	returns a shallow copy of a set
set.difference(set2)	returns the difference of set and set2
<pre>set.difference_update(set2)</pre>	removes all elements of another set from this set
set.discard(element)	removes an element from set if it is found in set. (Do nothing if the element is not in set)
<pre>set.intersection(sets)</pre>	return the intersection of set and the other provided sets
<pre>set.intersection_update(sets)</pre>	updates set with the intersection of set and the other provided sets
<pre>set.isdisjoint(set2)</pre>	returns True if set and set2 have no intersection
set.issubset(set2)	returns True if set2 contains set
<pre>set.issuperset(set2)</pre>	returns True if set contains set2
set.pop()	removes and returns an arbitrary element of set.
set.remove(element)	removes element from a set.
<pre>set.symmetric_difference(set2)</pre>	returns the symmetric difference of set and set2
<pre>set.symmetric_difference_update(set2)</pre>	updates set with the symmetric difference of set and set2
set.union(sets)	returns the union of set and the other provided sets
set.update(set2)	update set with the union of set and set2

#### Build a dictionary of NT counts using a set and loops

Let us put a twist on our nt count script. Let's use a set to find all the unique nts, then use the string count() method to count the nucleotide instead if incrementing the count as we did earlier.

Code:

```
#!/usr/bin/env python3

# create a new empty dictionary
nt_count = {}

# get a set of unique characters in our DNA string

dna = 'GTACCNTTGATTTCGTATTCTGAGAGGCTGCTTAGCGGTAGCCCCTTGGTTTCCGTGGCAACGGAAAA'
unique = set(dna)

print('unique nt: ', unique) ## {'C', 'A', 'G', 'T', 'N'}

# iterate through each unique nucleotide
for nt in unique:
    # count the number of this unique nt in dna
    count = dna.count(nt)

# add our count to our dict
nt_count[nt] = count

print('nt count:', nt_count)
```

Output:

```
unique nt: {'N', 'C', 'T', 'G', 'A'}
nt count: {'G': 20, 'T': 21, 'A': 13, 'C': 16, 'N': 1}
```

We have the count for all NTs even ones we might not expect.

# <u>Link to Python 5 Problem Set</u>