# Python 6

# **Regular Expressions**

Regular Expressions is a language for pattern matching. Many different computer languages incorporate regular expressions as well as some unix commands like grep and sed. So far we have seen a few functions for finding exact matches in strings, but this is not always sufficient.

Functions that utilize regular expressions allow for non-exact pattern matching.

These specialized functions are not included in the core of Python. We need to import them by typing import relat the top of your script

```
1 #!/usr/bin/env python3
2
3 import re
```

First we will go over a few examples then go into the mechanics in more detail.

Let's start simple and find an exact match for the EcoRI restriction site in a string.

Since we can search for control characters like a tab (\t) it is good to get in the habit of using the raw string function \( \text{r} \) when defining patterns.

Here we used the search() function with two arguments, 1) our pattern and 2) the string we want to search.

Let's find out what is returned by the search() function.

Information about the first match is returned

How about a non-exact match. Let's seach for a methylation site that has to match the following critera:

- G or A
- followed by C
- followed by one of anything or nothing
- followed by a G

This could match any of these:

**GCAG** 

GCTG

GCGG

**GCCG** 

GCG

ACAG

ACTG

ACGG

ACCG

ACG

We could test for each of these, or use regular expressions. This is exactly what regular expressions can do for us.

Here you can see in the returned information that ACG starts at string postion 7 (nt 8). The first position following the end of the match is at string postion 10 (nt 11).

What about other potential matches in our DNA string? We can use findall() function to find all matches.

findall() returns a list of all the pieces of the string that match the regex.

A quick count of all the matching sites can be done by counting the length of the returned list.

```
1 >>> len (re.findall(r"[GA]C.?G",dna))
2 7
```

There are 7 methylation sites. Here we have another example of nesting. We call the findall() function, searching for all the matches of a methylation site. This function returns a list, the list is past to the len() function, which in turn returns the number of elements in the list.

Let's talk a bit more about all the new characters we see in the pattern.

The pattern in made up of atoms. Each atom represents **ONE** character.

### **Individual Characters**

Atom	Description
a-z, A-Z, 0-9 and some punctution	These are ordinary characters that match themselves
","	The dot, or period. This matches any single character execpt for the newline.

#### **Character Classes**

A group of characters to that are allowed to be matched one time. There are a few predefined classes, symbol that means a series of characters.

Atom	Description
	A bracketed list of characters, like [GA]. This indicates a single character can match any character in the bracketed list.
\d	Digits. Also can be written [ø-9]
\D	Not digits. Also can be written [^0-9]
\w	Word character. Also can be written [A-Za-z0-9_] Note underscore is part of this class
\W	Not a word character, or [^A-Za-z0-9_]
\s	White space character. Also can be written $\lceil \r \rceil$ . Note the space character after the first $\lceil$
\\$	Not whitespace. Also [^ \r\\t\n]

# **Anchors**

A pattern can be anhcored to a region in the string

Atom	Description
^	Matches the beginning of the string
\$	Matches the end of the string
\b	Matches a word boundary between w and w

# Examples:

```
1 |g..t
```

matches "gaat", "goat", and "gotta get a goat" (twice)

```
1 | g[gatc][gatc]t
```

matches "gaat", "gttt", "gatt", and "gotta get an agatt" (once)

```
1 \d\d\d-\d\d\d\\
```

matches 376-8380, and 5128-8181 but not 055-98-2818.

- matches 376-8380 and 376-83801 but not 5128-8181.
- only matches telephone numbers (without area code)
- 1 \bcat
- matches "cat", "catsup" and "more catsup please" but not "scat".
- 1 \bcat\b
- only text containing the word "cat".

# **Quantifiers**

Quantifiers quantify how many atoms are to be found. By default an atom matches only once. This behaviour can be modified following an atom with a quantifier.

Quantifier	Description
?	atom matches zero or exactly once
*	atom matches zero or more times
+	atom matches one or more times
{3}	atom matches exactly 3 times
{2,4}	atom matches between 2 and 4 times, inclusive
[4,}]	atom matches at least 4 times

### Examples:

- 1 goa?t
- matches "goat" and "got". Also any text that contains these words.
- 1 g.+t
- matches "goat", "goot", and "grant", among others.

```
1 | g.*t
```

matches "gt", "goat", "goot", and "grant", among others.

```
1 \\^\d{3}-\d{4}$
```

matches US telephone numbers (no extra text allowed).

Something to think about.

1) What would be a pattern to recognize an email address? 2) What would be a pattern to recognize the ID portion of a sequence record in a FASTA file?

### Variables and Patterns

Variables can be used to store patterns.

```
1  >>> pattern = r"[GA]C.?G"
2  >>> len (re.findall(pattern,dna))
3  7
```

In this example we stored our methylation pattern in the variable named 'pattern' and used it as the first argument to findall.

#### **Either Or**

A pipe '|' can be used to indicated that either the pattern before or after the '|' can match. Enclose the two options in parenthesis.

```
1 | big bad (wolf|sheep)
```

This pattern must match a string that contains "big" followed by a space followed by "bad" followed by a space followed by either "wolf" or "sheep" This would match, "big bad wolf" Or "big bad sheep"

Something to think about. 1) What would a pattern to match 'ATG' followed by a C or a T look like?

# **Subpatterns**

Subpatterns, or parts of the pattern enclosed in parenthesis can be extracted and stored for later use.

```
1 Who's afraid of the big bad w(.+)f
```

This pattern has only one subpattern (.+)

You can combine parenthesis and quantifiers to quantify entire subpatterns

```
1 Who's afraid of the big (bad )?wolf\?
```

This matches "Who's afraid of the big bad wolf?". As well as "Who's afraid of the big wolf?". The 'bad ' is optional, it can be present 0 or 1 times in our string. This also shows how to literally match special characters. Use a '\' in to escape them.

Something to think about: How would you create a pattern to capture the ID in a sequence record of a FASTA file in a subpattern.

Example FASTA sequence record.

```
1 >ID Optional Descrption
2 SEQUENCE
3 SEQUENCE
4 SEQUENCE
```

### **Using Subpatterns Inside the Regular Expression Match**

This is helpful when you want to find a subpattern and then match the contents again. They can be used within the function call and used after the function call.

### Subpatterns within the function call

Once a subpattern matches, you can refer to it within the same regular expression. The first subpattern becomes \1, the second \2, the third \3, and so on.

```
1 Who's afraid of the big bad w(.)\1f
```

This would match "Who's afraid of the big bad woof" "Who's afraid of the big bad weef" "Who's afraid of the big bad waaf" But Not, "Who's afraid of the big bad wolf" Or, "Who's afraid of the big bad wife"

In a similar vein,

```
1 \b(\w+)s love \1 food\b
```

This pattern will match "dogs love dog food" But not "dogs love monkey food". We were able to use the subpattern within the regular expression by using  $\label{log}$  If there were more subpatterns they would be  $\label{log}$ ,  $\label{log}$ ,  $\label{log}$ ,  $\label{log}$ ,  $\label{log}$ 

# **Using Subpatterns Outside the Regular Expression**

Subpatterns can be retrieved after the <code>search()</code> function call, or outside the regular expression, by using the <code>group()</code> method. This is a method and it belongs to the object that is returned by the <code>search()</code> function.

The subpatterns are retrieved by a number. This will be the same number that could be used within the regular expression, i.e.,

- \1 within the subpattern can be used outside with search\_found\_obj.group(1)
- \2 within the subpattern can be used outside with search found obj.group(2)
- \3 within the subpattern can be used outside with search\_found\_obj.group(3)
- and so on

### Example:

1) This pattern will recognize a consensus transcription start site (TATTAT) 2) And store the 50 base pairs upstream of the site 3) And the 25 base pairs downstream of the site

If you want to find the upstream and downstream sequence of ALL 'TATTAT' sites, use the findall() function.

```
dna="ACAAAATACGTTTTGTAAATGTTGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCAC
    CGGTTTCCAAAGACAGTCTTCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTC
    CAAAGACAGTCTTCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGACAAAATACGTTTTGTAAATGT
    TGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCACCGGTTTCCAAAAGACAGTCTTCTAATTC
    CTCATTAGTAATAAGTAAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTCCAAAAGACAGTCTTCTAATTCCTCATTA
    GTAATAAGTAAAATGTTTATTGTTGTAGCTCTGG"

2    >>> found = re.findall( r"(.{50})TATTAT(.{25})" , dna )
3    >>> print(found)
4    [('TCTAATTCCTCATTAGTAATAAGTAAAAATGTTTATTGTTGTAGCTCTGGA',
    'CCGGTTTCCAAAGACAGTCTTCTAA'),
    ('TCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGA',
    'CCGGTTTCCAAAGACAGTCTTCTAA')]
```

The subpatterns are stored in tuples within a list. More about this type of data structure later.

Another option for retrieving the upstream and downstream subpatters is to put the findall() in a for loop

```
>>>
   dna="ACAAAATACGTTTTGTAAATGTTGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCAC
   CGGTTTCCAAAGACAGTCTTCTAATTCCTCATTAGTAAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTC
   CAAAGACAGTCTTCTAATTCCTCATTAGTAAATAAGTAAAATGTTTATTGTTGTAGCTCTGGACAAAATACGTTTTGTAAATGT
   TGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCACCGGTTTCCAAAGACAGTCTTCTAATTC
   CTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTCCAAAGACAGTCTTCTAATTCCTCATTA
   GTAATAAGTAAAATGTTTATTGTTGTAGCTCTGG"
2
   >>> for (upstream, downstream) in re.findall( r"(.{50})TATTAT(.{25})" , dna ):
         print("upstream:" , upstream)
3
         print("downstream:" , downstream)
4
   . . .
5
   upstream: TCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGA
6
7
   downstream: CCGGTTTCCAAAGACAGTCTTCTAA
   upstream: TCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGA
  downstream: CCGGTTTCCAAAGACAGTCTTCTAA
9
```

- 1) This code executes the findall() function once
- 2) The subpatterns are returned
- 3) The subpatterns are stored in the variables upstream and downstream
- 4) The for block of code is executed
- 5) The findall() searches again
- 6) A match is found
- 7) New subpatterns are returned and stored in the variables upstream and downstream 8) The for block of code gets executed again
- 9) The findall() searches again, but no match is found
- 10) The for loop ends

Another way to get this done is with an iterator, use the finditer() function in a for loop. This allows you to not store all the matches in memory. finditer() also allows you to retrieve the postion of the match.

```
>>>
1
   dna="ACAAAATACGTTTTGTAAATGTTGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCAC
   CGGTTTCCAAAGACAGTCTTCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTC
   CAAAGACAGTCTTCTAATTCCTCATTAGTAAATAAGTAAAATGTTTATTGTTGTAGCTCTGGACAAAATACGTTTTGTAAATGT
   TGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCACCGGTTTCCAAAGACAGTCTTCTAATTC
   CTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTCCAAAGACAGTCTTCTAATTCCTCATTA
   GTAATAAGTAAAATGTTTATTGTTGTAGCTCTGG"
   >>> for match in re.finditer(r"(.{50})TATTAT(.{25})" , dna):
3
         print("upstream:" , match.group(1))
4
         print("downstream:" , match.group(2))
   . . .
5
6
   upstream: TCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGA
   downstream: CCGGTTTCCAAAGACAGTCTTCTAA
8
   upstream: TCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGA
   downstream: CCGGTTTCCAAAGACAGTCTTCTAA
```

- 1) This code executes finditer() function once.
- 2) The match object is returned. A match object will have all the information about the match
- 3) In the for block we call the <code>group()</code> method on the first match object returned
- 4) We print out the first and second subpattern using the <code>group()</code> method
- 5) The finditer() function is executed a second time and a match is found
- 6) The second match object is returned
- 7) The second subpatterns are retrieved from the match object using the <code>group()</code> method
- 8) The finditer() functin is executed again, but no matches found, so the loop ends

# Get position of the subpattern with finditer()

The match object contains information about the match that can be retrieved with match methods like start() and end()

```
#!/usr/bin/env python3
1
2
3
    import re
    dna="ACAAAATACGTTTTGTAAATGTTGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCA
    {\sf CCGGTTTCCAAAGACAGTCTTCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTT}
    TCCAAAGACAGTCTTCTAATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGACAAAATACGTTTTGTAAA
    TGTTGTGCTGTTAACACTGCAAATAAACTTGGTAGCAAACACTTCCAAAAGGAATTCACCGGTTTCCAAAGACAGTCTTCTA
    ATTCCTCATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGGATATTATCCGGTTTCCAAAGACAGTCTTCTAATTCCT
    CATTAGTAATAAGTAAAATGTTTATTGTTGTAGCTCTGG"
    for found in re.finditer(r"(.{50})TATTAT(.{25})", dna):
7
      whole = found.group(0)
8
      up
9
             = found.group(1)
      down = found.group(2)
10
      up start = found.start(1) + 1 # need to convert from 0 to 1 notation
11
      up\_end = found.end(1) + 1
12
      dn_start = found.start(2) + 1
13
      dn end = found.end(2) + 1
14
15
      output = [ whole , up , str(up_start), str(up_end) , down , str(dn_start) ,
16
    str(dn_end) ]
17
18
      print( "\t".join(output) )
```

we can use these match object methods <code>group()</code>, <code>start()</code>, <code>end()</code> to get the string, start position, and end position of each subpattern.

**FYI:** match() function is another regular expression function that looks for patterns. It is similar to search but it only looks at the begining of the string for the pattern while search() looks in the entire string. Usually finditer(), search() and findall() will be more useful.

# **Subpatterns and Greediness**

By default, regular expressions are "greedy". They try to match as much as they can. Use the quantifier '?' to make the match not greedy. The not greedy match is called 'lazy'

```
1  >>> str = 'The fox ate my box of doughnuts'
2  >>> found = re.search(r"(f.+x)",str)
3  >>> print(found.group(1))
4  fox ate my box
```

The pattern f.+x does not match what you might expect, it matches past 'fox' all the way out to 'fox ate my box'.

The '.+' id greedy As many characters as possible are found that are between the 'f' and the 'x'.

Let's make this match lazy by using '?'

```
1  >>> found = re.search(r"(f.+?x)",str)
2  >>> print(found.group(1))
3  fox
```

The match is now lazy and will only match 'fox'

### **Practical Example: Codons**

Extracting codons from a string of DNA can be accomplished by using a subpattern in a findall() function. Remember the findall() function will return a list of the matches.

```
1  >>> dna = 'GTTGCCTGAAATGGCGGAACCTTGAA'
2  >>> codons = re.findall(r"(.{3})",dna)
3  >>> print(codons)
4  ['GTT', 'GCC', 'TGA', 'AAT', 'GGC', 'GGA', 'ACC', 'TTG']
```

Or you can use a for loop to do something to each match.

```
>>> for codon in re.findall(r"(.{3})",dna):
          print(codon)
2
    . . .
3
    . . .
4
    GTT
    GCC
    TGA
6
7
    AAT
    GGC
8
9
    GGA
10
    ACC
11
    TTG
12
    >>>
```

```
finditer() would also work in this for loop.
Each codon can be accessed by using the group() method.
```

### **Truth and Regular Expression Matches**

The search(), match(), findall(), and finditer() can be used in conditional tests. If a match is not found an empty list or 'None' is returned. These both are False.

```
>>> found=re.search( r"(.{50})TATTATZ(.{25})" , dna )
2
   >>> if found:
          print("found it")
3
   . . .
4
   ... else:
   . . .
          print("not found")
6
7
   not found
   >>> print(found)
8
9
   None
```

None is False so the else block is executed and "not found" is printed

#### Nest it!

```
1
   >>>
   >>> if re.search( r"(.{50})TATTATZ(.{25}))" , dna ):
3
          print("found it")
   ... else:
4
          print("not found")
5
   . . .
6
  . . .
7
   not found
  >>> print(found)
8
  None
```

### **Using Regular expressions in substitutions**

Earlier we went over how to find an **exact pattern** and replace it using the <code>replace()</code> method. To find a pattern, or inexact match, and make a replacement the regular expression <code>sub()</code> function is used. This function takes the pattern, the replacement, the string to be searched, the number of times to do the replacement, and flags.

```
1  >>> str = "Who's afraid of the big bad wolf?"
2  >>> re.sub(r'w.+f' , 'goat', str)
3  "Who's afraid of the big bad goat?"
4  >>> print(str)
5  Who's afraid of the big bad wolf?
```

The sub() function returns "Who's afraid of the big bad goat?"

The value of variable str has not been altered

The new string can be stored in a new variable for later use.

Let's save the returned new string in a variable

```
1     >>> str = "He had a wife."
2     >>> new_str = re.sub(r'w.+f' , 'goat', str)
3     >>> print(new_str)
4     He had a goate.
5     >>> print(str)
6     He had a wife.
```

The characters between 'w' and 'f' have been replaced with 'goat'.

The new string is saved in new\_str

### Using Regular expressions in substitutions

Sometimes you want to find a pattern and use it in the replacement.

```
1  >>> str = "Who's afraid of the big bad wolf?"
2  >>> new_str = re.sub(r"(\w+) (\w+) wolf" , r"\2 \1 wolf" , str)
3  >>> print(new_str)
4  Who's afraid of the bad big wolf?
```

We found two words before 'wolf' and swapped the order. \2 refers to the second subpattern \1 refers to the first subpattern

Something to think about.

How would you use regular expressions to find all occurances of 'ATG' and replace with '-M-' in this sequence 'GCAGAGGTGATGGACTCCGTAATGGCCAAATGACACGT'?

# Using subpatterns in the replacement

Sometimes you want to find a pattern and use it in the replacement.

```
1  >>> str = "Who's afraid of the big bad wolf?"
2  >>> new_str = re.sub(r"(\w+) (\w+) wolf" , r"\2 \1 wolf" , str)
3  >>> print(new_str)
4  Who's afraid of the bad big wolf?
```

We found two words before 'wolf' and swapped the order. \2 refers to the second subpattern \1 refers to the first subpattern

Something to think about.

How would you use regular expressions to find all occurances of 'ATG' and replace with '-M-' in this sequence 'GCAGAGGTGATGGACTCCGTAATGGCCAAATGACACGT'?

# **Regular Expression Option Modifiers**

Modifier	Description
re.I re.IGNORECASE	Performs case-insensitive matching.
re.MULTILINE	Makes \$ match the end of a line (not just the end of the string) and makes ^ match the start of any line (not just the start of the string).
re.S re.DOTALL	Makes a period (dot) match any character, including a newline.
re.U	Interprets letters according to the Unicode character set. This flag affects the behavior of \w, \W, \b, \B.
re.X VERBOSE	This flag allows you to write regular expressions that look nicer and are more readable by allowing you to visually separate logical sections of the pattern and add comments. Whitespace within the pattern is ignored, except when in a character class or when preceded by an unescaped backslash. When a line contains a # that is not in a character class and is not preceded by an unescaped backslash, all characters from the leftmost such # through the end of the line are ignored.

```
1  >>> dna = "atgcgtaatggc"
2  >>> re.search(r"ATG",dna)
3  >>>
4  >>> re.search(r"ATG",dna , re.I)
5  <_sre.SRE_Match object; span=(0, 3), match='atg'>
6  >>>
```

We can make our search case insenstive by using the re.I or re.IGNORECASE flag.

You can use more than one flag by concatenating them with []. re.search(r"ATG",dna, re.I|re.M)