

Prof. Dr. Boas Pucker

Python - Control structures



Availability of slides

- All materials are freely available (CC BY) after the lectures:
 - GitHub: https://github.com/bpucker/PyBo

Questions: Feel free to ask at any time



Feedback, comments, or questions: pucker[a]uni-...

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If & else

- Distinguish between two cases:
 - Analyzing table: row of interest / irrelevant row
- Action depends on result of comparison

```
a = 5 #define variable
#user inputs number
b = int( input("please enter number!") )
if b < a:#if b is smaller than a
    print("b is smaller than a")
else:
    print("b is NOT smaller than a")</pre>
```

please enter number!3
b is smaller than a



elif

- Distinguish between multiple cases
 - Annealing temperature for PCR: suitable / too low / too high
- Action depends on result of comparison

```
a = 5 #define variable
#user inputs number
b = int( input("please enter number!") )
if b < a:#if b is smaller than a
    print("b is smaller than a")
elif b == a:
    print("b is matching a")
else:
    print("b is bigger than a")</pre>
```

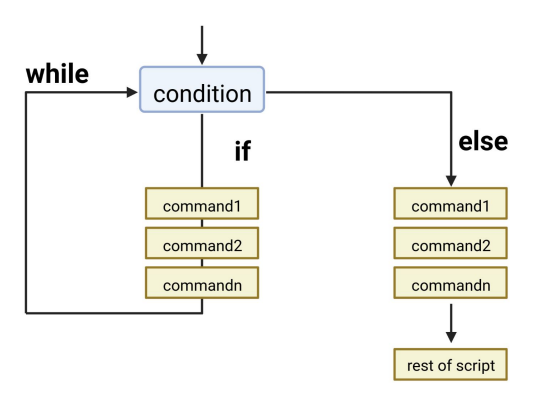


Exercises - Part3a

- 3.1) Write a script for guessing numbers!
- 3.2) Add tips (smaller/larger) during the guessing process!



Concept of loops





While loop (example)

```
pwhile a < 10: #checks if a is smaller than 10</pre>
 3
          print(str( a ) + " is smaller than 10")
 4
          a += 1 \# a = a+1
         #something useful could happen here
print("a was increased by 1")
 5
 6
     print(str( a ) + "is larger than 10")
  Code is executed until the
condition for this loop becomes
            false
```



While infinite loop

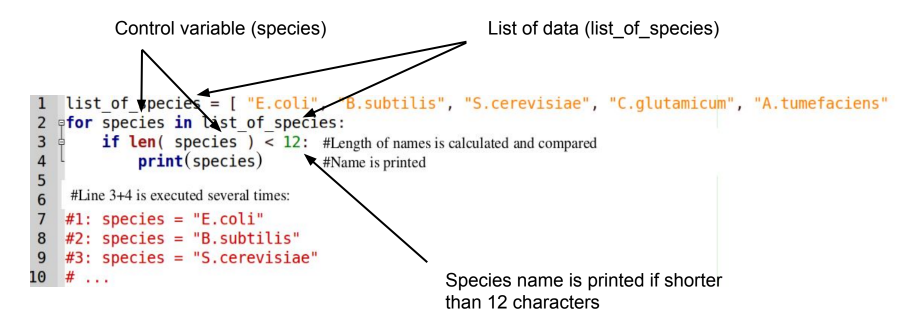
WARNING: this loop is infinite!

```
#infinite loop:
a = 0

while True: #always true
a += 1 #a = a+1
print(str( a ))
print("this line is never reached")
```



For loop





Exercises - Part3b

- 3.3) Write a function counting to 100 and printing all number which can be divided by 4 without any residue!
 - Info: 10%2 #modulo division in Python
- 3.4) Write a function counting down from 1000 to 0 and printing all numbers!
- 3.5) Generate a list of species names! Write a function that prints all species names starting with 'E'!
- 3.6) Expand this function to limit the printing to species names which are additionally shorter than 10 characters!
- 3.7) Expand this function to limit the printing to species names which are additionally ending with 'a'.



Exercises - Part3b (addition)

- 3X1: Print species names that match at least 2 of the following 3 conditions: (1) longer than 8 characters, (2) start with 'E', (3) end with 'a'
- 3X2: Print species names that are identical to the previous or next species name in the list.
- 3X3: Implement a function that can quickly sort the elements of a list alphabetically.



range()

```
list_of_species = ["E.coli", "B.subtilis", "S.cerevisiae", "C.glutamicum", "A.tumefaciens"]
length = len( list_of_species ) #length = 5

for i in range( length ): #starts at 0 and runs to i=4 (five values)

if len( list_of_species[ i ] ) < 12: #length of name is calculated and compared

print(list_of_species[ i ]) #name is printed

#i is taking five different values:
#1: i=0
#2: i=1
#4: i=3
#4: i=3
#5: i=4
#4: i=5 is never reached by range()</pre>
```



enumerate()



Exercises - Part3c

- 3.8) Write a script to print 50x 'here' and the current value of the control variable!
- 3.9) Write a script to walk through the species list and to print the character from the species where the index corresponds to the current control variable value!



Checking a list/string for a sublist/substring

Check if an element is contained in a list:

```
genes = ["g1", "g2", "g3", "g4", "g5"]
if "g3" in genes:
    print("g3 in genes")

g3 in genes
```

Check if a substring is contained in a string:

```
genome_seq = "ACACCGATTACTGGAGGTTACGTAATGGCCA"
primer = "ACTGGAGGTTACG"
if primer in genome_seq:
    print("primer binding site detected.")
else:
    print("no primer binding site detected")
```

primer binding site detected.



Break & pass

- Break allows you to exit a loop once a condition is met
- Pass allows you to skip elements in a loop

```
1 values = ["species1", "species2", "species3", "species4"]
2 for val in values:
3    if int( val[-1] ) < 3:
4        pass
5    elif int( val[-1] ) == 4:
6    break</pre>
```



Dictionaries

- Dictionary = data structure that allows very efficient access
- update() = adds new elements (dictionaries) to an existing dictionary

```
1 # dictionary:
2 my_dictionary = { 'key1': 'valueA', 'key2': 'valueB', 'key3': 'valueC' }
3 
4 # adding an element to a dictionary:
5 my_dictionary.update( { 'key4': 'valueD' } )
```



Try & except

- Try = run a code block until an error occurs
- Except = capture one or multiple errors and run alternative code

```
#Try & except:
my_dictionary = { 'key1': 'valueA', 'key2': 'valueB', 'key3': 'valueC' }

dict_keys = list( my_dictionary.keys() )
for key in dict_keys:
    try:
    value = my_dictionary[ key ]
    except KeyError:
    print( "invalid key: " + key )
```



Time for questions!