Class 6: R functions

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1. Function basics

Let's start writing our first silly function to add some numbers:

Every R function has 3 things:

- name (we get to pick this)
- input arguments (there can be loads of these separated by a comma)
- the body (the R code that does the work)

```
add <- function(x, y=10, z=0) {
    x + y + z
}
```

I can just use this function like any other function as long as R knows about it (i.e. run the code chunk)

```
add(1, 100)
```

[1] 101

```
add( x=c(1,2,3,4), y =100)
```

[1] 101 102 103 104

```
add(1)
```

[1] 11

Functions can have "required" input arguments and "optional" input arguments. The optional arguments are defined with an equals default value (y=10) in the function defination.

```
add(x=1,y=100,z=10)
```

[1] 111

Q. Write a function to return a DNA sequence of a user specified length. Call it generate_dna()

The sample() function can help here

```
#generate_dna <- function(size=5){ }
students <- c("jeff", "jeremy", "peter")
sample(students, size = 5, replace = TRUE)</pre>
```

```
[1] "peter" "jeff" "peter" "jeff" "jeremy"
```

2. Generate DNA sequence

Now work with bases rather than students

```
bases <- c("A","C", "G", "T")
sample(bases, size = 10, replace = TRUE)</pre>
```

```
[1] "C" "A" "T" "A" "C" "T" "T" "T" "A"
```

Now I have a working 'snippet' of code T can use this as the body of my first function version here:

```
generate_dna <- function(size=5) {
  bases <- c("A","C", "G", "T")
  sample(bases, size = size, replace = TRUE)
}</pre>
```

```
generate_dna()
```

```
[1] "G" "T" "A" "A" "A"
```

I want the ability to return a sequence like "AGTACCTG" i.e. a one element vector where the bases are all together.

```
generate_dna <- function(size=5, together=TRUE) {
  bases <- c("A","C", "G", "T")
  sequence <- sample(bases, size = size, replace = TRUE)

if(together) {
  sequence <- paste(sequence, collapse = "")
  }
  return(sequence)
}</pre>
```

```
generate_dna()
```

[1] "TGTCT"

```
generate_dna(together = F)
```

```
[1] "A" "T" "T" "A" "A"
```

3. Generate Protein Function

Q. Write a protein sequence generating function that will return sequences of a user specified length?

We can get the set of 20 natural amino-acids from the bio3d package.

```
aa <- bio3d::aa.table$aa1[1:20]
```

```
generate_protein <- function(size=6, together=TRUE) {

##Get the 20 amino acids as a vector
aa <- bio3d::aa.table$aa1[1:20]
sequence <- sample(aa, size = size, replace = TRUE)

## Optionally return a single element string
if(together) {
   sequence <- paste(sequence, collapse = "")
}
return(sequence)
}</pre>
```

```
generate_protein()
```

- [1] "RHFEQI"
 - Q. Generate random protein sequences of length 6 to 12 amino acids.

```
generate_protein(7)
```

[1] "TINYWLP"

```
generate_protein(8)
```

[1] "QCMTMSQA"

```
generate_protein(9)
```

[1] "VWTLPPQRR"

```
# generate_protein(size=6:12)
```

We can fix this inability to generate multiple sequences by either editing and adding to the function body code (e.g. a for loop) or by using the R apply family of utility functions.

```
sapply(6:12, generate_protein)
[1] "QTLGSW"
                                     "IFQAKVTD"
                                                     "CPIEERQEY"
                                                                     "PGQGNSEYTH"
                    "CPKIGVF"
[6] "MWWDFPDCRVS"
                    "CRGDCTQPHPVW"
It would be cool and useful if I could get FASTA format output
ans <- sapply(6:12, generate_protein)</pre>
ans
[1] "HVKEHK"
                                                     "RFGKTYGSF"
                    "DGWCLAR"
                                     "YYQSLVYQ"
                                                                     "NYACAYHLAR"
[6] "QYCNSYLCRWV"
                    "INSEKLNDSCCA"
cat(ans, sep="\n")
HVKEHK
DGWCLAR
YYQSLVYQ
RFGKTYGSF
NYACAYHLAR
QYCNSYLCRWV
INSEKLNDSCCA
I want this to look like FASTA format with an ID line. e.g.
>ID.6
YYKMTW
>ID.7
ERFAPFW
>ID.8
RYYCSTLL
The functions paste() and cat() can help us here...
```

```
cat(paste(">ID.", 6:12, "\n", ans, sep=""), sep="\n")
```

```
>ID.6
HVKEHK
>ID.7
DGWCLAR
>ID.8
YYQSLVYQ
>ID.9
RFGKTYGSF
>ID.10
NYACAYHLAR
>ID.11
QYCNSYLCRWV
>ID.12
INSEKLNDSCCA
```

```
id.line <- paste(">ID.", 6:12, sep="")
id.line
```

```
[1] ">ID.6" ">ID.7" ">ID.8" ">ID.9" ">ID.10" ">ID.11" ">ID.11" ">ID.12"
```

```
id.line <- paste(">ID.", 6:12, sep="")
seq.line <- paste(id.line, ans, sep="\n")
cat(seq.line, sep="\n", file ="myseq.fa")</pre>
```

Q. Determine if these sequences can be found in nature or are they unique

I BLASTp searched my FASTA format sequences against NR and found that length 6, 7, 8 are not unique and can be found in the databases with 100% coverage and 100% identity.

Random sequences of length 9 and above are unique and can't be found in the databases.