Advanced R programming: practical 2 Dr Colin Gillespie May 7, 2014

- 1 Rprofile
- 1. Create an .Rprofile file. Put in the line

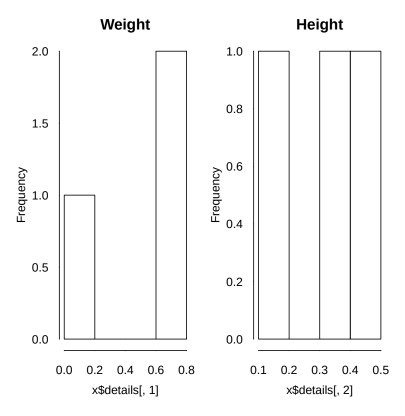
```
cat("Successfully loaded .Rprofile at", date(), "\n")
## Successfully loaded .Rprofile at Wed May 7 21:20:13 2014
```

Restart R. Does the welcome message appear.

- 2. Try adding my suggestions to your .Rprofile
- 2 S3 objects
- 1. Following the cohort example in the notes, suppose we want to create a mean method.
 - List all S₃ methods associated with the mean function.
 - Examine the source code of mean.
 - What are the arguments of mean?
 - Create a function called mean.cohort that returns a vector containing the mean weight and mean height.¹
- 2. Let's now make a similar function for the standard deviation
 - Look at the arguments of the sd function.
 - Create an function call sd.cohort that returns a vector containing the weight and height standard deviation.²
 - Create a default sd function. Look at cor.default in the notes for a hint.
- 3. Create a hist method for the cohort class. When the hist function is called on a cohort, it should produce a single plot showing two histograms one for height and another for weight.

¹ Ensure that you can pass in the standard mean arguments, i.e. na.rm.

² Ensure that you can pass in the standard sd arguments, i.e. na.rm.



4. Create a [method for the cohort class. This method should return a cohort object, but with the relevant rows sub setted. For example, if ç was a cohort object,

would return the first three rows of the data frame.

5. Create a verb"[<-" method for the cohort class. This method should allow us to replace values in the details data frame, i.e.

$$cc[1, 1] = 10$$

- S4 objects
- 1. Following the Cohort example in the notes, suppose we want to make a generic for the mean function.
 - Using the isGeneric function, determine if the mean function is an S4 generic. If not, use setGeneric to create an S4 generic.
 - Using setMethod, create a mean method for the Cohort class.³
- 2. Repeat the above steps for the sd function.
- 3. Create a hist method for the cohort class. When the hist function is called on a cohort, it should produce a single plot showing two histograms - one for height and another for weight.

I've intentionally mirrored the functions from section 2 of this practical to highlight the differences.

³ Be careful to match the arguments.

4. Create a [method for the cohort class. This method should return a cohort object, but with the relevant rows sub setted. For example, if ç was a cohort object,

```
cc[1:3, ]
```

would return the first three rows of the data frame.

5. Create a verb"[<-" method for the cohort class. This method should allow us to replace values in the details data frame.

Reference classes

The example in the notes created a random number generator using a reference class.

- Reproduce the randu generator from the notes and make sure that it works as advertised.4
- When we initialise the random number generator, the very first state is called the seed. Store this variable and create a new function called get_seed that will return the initial seed, i.e.

```
r = randu(calls = 0, seed = 10, state = 10)
r$r()
## [1] 0.0003052
r$get_state()
## [1] 655390
r$get_seed()
## [1] 10
```

• Create a variable that stores the number of times the generator has been called. You should be able to access this variable with the function get_num_calls

```
r = randu(calls = 0, seed = 10, state = 10)
r$get_num_calls()
## [1] 0
r$r()
## [1] 0.0003052
r$r()
## [1] 0.001831
r$get_num_calls()
## [1] 2
```

⁴ The reference class version, not the function closure generator.

Reference classes also have an initialise method - that way we would only specify the seed and would then initialise the other variables. I'll give you an example in the solutions.

«P2, echo=echo» @

Solutions

Solutions are contained within the course package

```
library("nclRadvanced")
vignette("solutions2", package = "nclRadvanced")
```