

Advanced R programming: solutions 2

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1 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.

```
methods("mean")  
  
## [1] mean,ANY-method      mean,Cohort-method  
## [3] mean.Date           mean.POSIXct  
## [5] mean.POSIXlt        mean.cohort  
## [7] mean.default         mean.difftime  
## see '?methods' for accessing help and source code
```

- Examine the source code of `mean`.

```
body("mean")
```

- What are the arguments of `mean`?

```
args("mean")  
  
## function (x, ...)  
## NULL
```

- Create a function called `mean.cohort` that returns a vector containing the mean weight and mean height.¹

```
mean.cohort = function(x, ...) {  
  m1 = mean(x$details[, 1], ...)  
  m2 = mean(x$details[, 2], ...)  
  return(c(m1, m2))  
}
```

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

2. Let's now make a similar function for the standard deviation

- Look at the arguments of the `sd` function.
- Create a 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.

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

```
sd = function(x, ...) UseMethod("sd")  
sd.default = function(x, ...) stats::sd(x, ...)  
sd.cohort = function(x, ...) {  
  s1 = sd(x$details[, 1], ...)  
  s2 = sd(x$details[, 2], ...)
```

```

    return(c(s1, s2))
}

```

3. Create a `summary` method for the `cohort` class. When the `summary` function is called on a `cohort` object it should call the base `summary` on the `details` element.

- Use the `body` function to check if the function is already a generic function.
- Use the `args` function to determine the arguments.
- Create a `summary.cohort` function

```

## summary is already a generic
body(summary)

## standardGeneric("summary")

## Match the args
args(summary)

## function (object, ...)
## NULL

## Function
summary.cohort = function(object, ...) summary(object$details, ...)

```

4. Create a `hist` method for the `cohort` class. When the `hist` function is called on a `cohort` object, it should produce a single plot showing two histograms - one for height and another for weight.

```

## hist is already a generic
body(hist)

## standardGeneric("hist")

## Match the args
args(hist)

## function (x, ...)
## NULL

## Function
hist.cohort = function(x, ...) {
  op = par(mfrow=c(1, 2))
  hist(x$details[,1], main="Weight")
  hist(x$details[,2], main="Height")
  par(op)
}

```

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

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

would return the first three rows of the data frame.

```
## Lots of methods available.
methods('[')

## [1] [,Cohort-method]      [,nonStructure-method]
## [3] [.AsIs]              [.Date]
## [5] [.Dlist]             [.POSIXct]
## [7] [.POSIXlt]           [.acf*]
## [9] [.bibentry*]         [.cohort]
## [11] [.data.frame]       [.difftime]
## [13] [.factor]            [.formula*]
## [15] [.fseq*]             [.getAnywhere*]
## [17] [.hexmode]           [.listof]
## [19] [.noquote]           [.numeric_version]
## [21] [.octmode]           [.pdf_doc*]
## [23] [.person*]          [.raster*]
## [25] [.roman*]           [.simple.list]
## [27] [.terms*]            [.ts*]
## [29] [.tskernel*]         [.warnings]
## see '?methods' for accessing help and source code

## Examine [.data.frame]
args('.data.frame')

## function (x, i, j, drop = if (missing(i)) TRUE else length(cols) ==
##   1)
## NULL

' [.cohort' = function(x, ...){
  x$details = x$details[...]
  x
}
```

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

```
cc[1,1] = 10
```

```
## Lots of methods available.
methods('[<-')

## [1] [<-,Cohort-method]      [<-,data.frame-method]
```

```

## [3] <- .Date           [<- POSIXct
## [5] <- .POSIXlt        [<- cohort
## [7] <- .data.frame     [<- factor
## [9] <- .numeric_version [<- raster*
## [11] <- .ts*
## see '?methods' for accessing help and source code

## Examine [.data.frame
args('[<-data.frame')

## function (x, i, j, value)
## NULL

'[<-cohort' = function(x, i, j, value){
  x$details[i, j] = value
  x
}
cc[1:3, ] = 55

```

2 S4 objects

- 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.

```

isGeneric("mean")
## [1] TRUE

setGeneric("mean")
## [1] "mean"

```

- Using `setMethod`, create a `mean` method for the `Cohort` class.³

```

setMethod("mean", signature=c("Cohort"),
          definition=function(x, ...) {
            m1 = mean(x@details[, 1], ...)
            m2 = mean(x@details[, 2], ...)
            return(c(m1, m2))
          }
)
## [1] "mean"

```

- Repeat the above steps for the `sd` function.

```

isGeneric("sd")
## [1] TRUE

```

I've intentionally mirrored the functions from section 1 of this practical to highlight the differences.

³ Be careful to match the arguments.

```
setGeneric("sd")

## [1] "sd"

setMethod("sd", signature=c("Cohort"),
            definition=function(x, na.rm=FALSE) {
              m1 = sd(x@details[, 1], na.rm=na.rm)
              m2 = sd(x@details[, 2], na.rm=na.rm)
              return(c(m1, m2))
            }
)

## [1] "sd"
```

3. Create a `summary` method for the `cohort` class

- Use `isGeneric` to determine if an S4 generic exists.
- Use `setGeneric` to set the generic method (if necessary).
- Create an S4 summary method.

```
isGeneric("summary")

## [1] TRUE

setGeneric("summary")

## [1] "summary"

setMethod("summary", signature=c("Cohort"),
            definition=function(object, ...) {
              summary(object@details)
            }
)

## [1] "summary"
```

4. 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.

```
isGeneric("hist")

## [1] TRUE

setGeneric("hist")

## [1] "hist"
```

```

setMethod("hist", signature=c("Cohort"),
          definition=function(x, ...) {
            op = par(mfrow=c(1, 2))
            hist(x@details[,1], main="Weight", ...)
            hist(x@details[,2], main="Height", ...)
            par(op)
          }
)

## [1] "hist"

```

5. Create a `[` method for the cohort class. This method should return a cohort object, but with the relevant rows sub setted.

```

isGeneric("[")  
  

## [1] TRUE  
  

getGeneric('["')  
  

## standardGeneric for "[" defined from package "base"  

##  

## function (x, i, j, ..., drop = TRUE)  

## standardGeneric("[", .Primitive("["))  

## <bytecode: 0x1f9ba80>  

## <environment: 0x1f90a60>  

## Methods may be defined for arguments: x, i, j, drop  

## Use showMethods("[") for currently available ones.  
  

## Can you determine what drop does?  

setMethod("[", signature=c("Cohort"),
          definition=function(x, i, j, ..., drop = TRUE) {
            x@details = x@details[i, j, ..., drop=drop]
            x
          }
)

## [1] "["

```

6. Create a `<-` method for the cohort class. This method should allow us to replace values in the `details` data frame.

```

isGeneric("[<-")  
  

## [1] TRUE  
  

setGeneric('[<-')

## [1] "[<-

```

```

setMethod("[<-", signature=c("Cohort"),
  definition=function(x, i, j, value) {
    x@details[i, j] = value
    x
  }
)

## [1] "[<-

coh_s4[1,]= 5

```

3 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] 0.0003051898

r$get_state()

## [1] 655390

r$get_seed()

## [1] 10

```

##Solutions - see below

- 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] 0.0003051898

r$r()

```

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.

```
## [1] 0.001831097
r$get_num_calls()
## [1] 2
```

```
## Solutions ##
randu = setRefClass("randu",
                     fields = list(calls = "numeric",
                                   seed="numeric",
                                   state="numeric"))
randu$methods(get_state = function() state)
randu$methods(set_state = function(initial) state <- initial)
randu$methods(get_seed = function() seed)
randu$methods(get_num_calls = function() calls)
randu$methods(r = function() {
  calls <- calls + 1
  state <- (65539*state) %% 2^31
  return(state/2^31)
})
```

Solutions

Solutions are contained within the course package

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