A practical tutorial on S4 programming

Laurent Gatto*

Cambridge Center for Proteomics University of Cambridge

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^{*}lg390@cam.ac.uk

1 Introduction

This document¹ introduces the R object-oriented programming paradigm using the microarray as a use case. The introduction is purely practical and does not aim for an exhaustive guide to R object-oriented programming. We will concentrate on the S4 system and only mention the older S3 system and the recent S4 reference class infrastructure. See the appropriate literature, ?ReferenceClasses or our more thorough introduction to OO programming² and references therein for mote details.

In section 2, we present a solution on how to represent microarray data in R using simple data types and conclude with some issues with this implementation. In section 3, we introduce fundamental concepts of OO programming and introduce how OO programming is implemented in S4 (and S3) system.

2 The microarray example

We assume the reader is familiar with the concept of microarrays and the type of data that is obtained from such an experiment. Before embarking in any serious programming task, in particular when modelling data and defining data structures (using a OO class or not), to carefully think about how to best represent and store the data.

Exercise 1: Based on your understanding of microarrays and the kind of data that is to be used to computational analysis, think of what is going to be needed to describe an experiment and what the types of data structures available in R (data.frame, matrix, vector, ...) are most appropriate. Ideally, one would want everything (data, meta-data, ...) to be stored together as a single variables.

There are of course multiple valid solutions to the above question. Below are three pieces of information that consider essential along with their respective R data structure.

- We choose to represent the microarray results as a matrix of size n × m, where
 n is the number of probes on the microarray and m is the number of samples.
 The matrix that stores the intensities (these could also represent fold-changes)
 is named marray.
- The sample annotation (meta-data) is described using a data.frame with exactly m rows and any number of columns. It is named pmeta.

¹The latest version of this document is available on its github repository https://github.com/lgatto/S4-tutorial.

²https://github.com/lgatto/roo

• The feature (probe) annotation (meta-data) is described using a data.frame with exactly n rows and any number of columns. Let's call it fmeta.

We will manually use the same names for intensity matrix columns and the sample meta-data rows as well as the matrix rows and feature meta-data row. Finally, to keep these pieces of information together, they will all be combined into a list that will represent our microarray experiment.

```
> rownames(pmeta) <- colnames(marray) <- LETTERS[1:m]
> fmeta <- data.frame(geneId = 1:n,</pre>
                       pathway = sample(LETTERS, n, replace = TRUE))
+
> rownames(fmeta) <-</pre>
      rownames(marray) <- paste0("probe", 1:n)</pre>
> maexp <- list(marray = marray,</pre>
+
                fmeta = fmeta,
                pmeta = pmeta)
> rm(marray, fmeta, pmeta)
> str(maexp)
List of 3
 $ marray: num [1:10, 1:6] 6.87 10.92 5.82 17.98 11.65 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:10] "probe1" "probe2" "probe3" "probe4" ...
  ....$ : chr [1:6] "A" "B" "C" "D" ...
 $ fmeta : 'data.frame': 10 obs. of 2 variables:
  ..$ geneId : int [1:10] 1 2 3 4 5 6 7 8 9 10
  ..$ pathway: Factor w/ 8 levels "E", "F", "L", "M", ..: 8 4 4 1 7 3 5 2 2 6
 $ pmeta :'data.frame': 6 obs. of 2 variables:
  ..$ sampleId : int [1:6] 1 2 3 4 5 6
  ..$ condition: Factor w/ 2 levels "MUT", "WT": 2 2 2 1 1 1
```

We can access the respective elements of our microarray experiment with the \$ operator.

```
> maexp$pmeta
  sampleId condition
         1
Α
                   WT
          2
В
                   WT
С
         3
                   WT
D
         4
                  MUT
Ε
         5
                  MUT
F
         6
                  MUT
> summary(maexp$marray[, "A"])
   Min. 1st Qu. Median
                           Mean 3rd Qu.
                                              Max.
           7.27
                   11.30
   5.82
                            10.70
                                    12.80
                                             18.00
> wt <- maexp$pmeta[, "condition"] == "WT"</pre>
> maexp$marray["probe8", wt]
     Α
             В
                    C
13.692 14.719 2.646
> maexp[["marray"]]["probe3", !wt]
          Ε
    D
11.94 13.48 11.71
```

Exercise 2: But what if we want to subset the experiment. How would we extract the 10 first probes for the 3 first samples?

We have to manually subset the individual elements of our list, making sure that the number of rows of the marray and fmeta elements remain identical as well as the number of columns of marray and the number of columns of pmeta.

```
> x <- 1:5
> y <- 1:3
> marray2 <- maexp$marray[x, y]
> fmeta2 <- maexp$fmeta[x, ]
> pmeta2 <- maexp$pmeta[y, ]
> maexp2 <- list(marray = marray2, fmeta = fmeta2, pmeta = pmeta2)
> rm(marray2, fmeta2, pmeta2)
> str(maexp2)
```

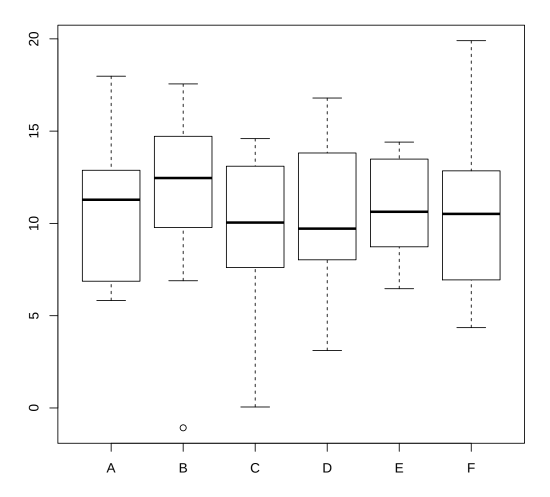


Figure 1: Boxplot representing the intensity distributions of the 10 probes for the 6 samples.

```
List of 3
$ marray: num [1:5, 1:3] 6.87 10.92 5.82 17.98 11.65 ...
..- attr(*, "dimnames")=List of 2
....$ : chr [1:5] "probe1" "probe2" "probe3" "probe4" ...
....$ : chr [1:3] "A" "B" "C"
$ fmeta : 'data.frame': 5 obs. of 2 variables:
..$ geneId : int [1:5] 1 2 3 4 5
..$ pathway: Factor w/ 8 levels "E","F","L","M",..: 8 4 4 1 7
$ pmeta : 'data.frame': 3 obs. of 2 variables:
..$ sampleId : int [1:3] 1 2 3
```

The above solution does not provide a clean syntax. As a user, we have to know the names or positions of the respective elements of the microarray list elements to directly access the parts of interest. Finally, a simple operation like sub-setting the microarray experiment is very cumbersome and prone to errors.

3 Using OO programming

Object-oriented programming is based on two important concepts, abstraction and encapsulation. We want to represent the microarray concept in a way that makes most sense to the users without distracting them with unnecessary technicalities. These technicalities refer to the underlying implementation. Do the users really need to know that we used a list and that the first element, called marray is the matrix? We want the users to comprehend microarrays in R like they know them in real life, i.e. manipulate the abstract concept microarray while keeping all the underlying technical details, the implementation, hidden, or encapsulated.

These goals are achieved in two steps. First, we defined a class that represents (abstracts) the concept of a microarray. This is very similar to what we have done with the list above (the S3 system does use list), but we will use a more elaborated approach that, although more verbose, provides numerous benefits that will be described in the next sections. The class represents a data container and is defined on its own. An instance of a specific class, that contains data arranged in the specific container, is called an object.

Once we have created a class, we will want to defined a set of specific behaviours, that make sense in the eyes of the users. These behaviours will be implemented by special functions, called methods. Methods are functions that tune their behaviour based on the class of their input. You have already observed this in your every day usage of R: whether we ask to produce the boxplot of a matrix (for example boxplot(maexp[[1]])) or provide a data.frame and a formula like boxplot(sampleId ~ condition, data = maexp[[3]]), R automatically does the right thing.

It now becomes obvious that we have two different kind of roles. The *developer* is the one that creates the class and knows the implementation and the *user* is the one that uses the class without knowing, or needing to know, its actual underlying representation.

4 The MArray class

We can define a class with the setClass function. Our class is defined by a name, MArray, and a content. The different elements of an S4 class are called slots³.

```
> MArray <- setClass("MArray",
+ slots = c(marray = "matrix",
+ fmeta = "data.frame",
+ pmeta = "data.frame"))</pre>
```

The setClass function returns a special function called a constructor, that can be used to create an instance of the class.

```
> ## an empty object
> MArray()
An object of class "MArray"
Slot "marray":
<0 x 0 matrix>
Slot "fmeta":
data frame with 0 columns and 0 rows
Slot "pmeta":
data frame with 0 columns and 0 rows
> ma <- MArray(marray = maexp[[1]],</pre>
               pmeta = maexp[["pmeta"]],
               fmeta = maexp[["fmeta"]])
> class(ma)
[1] "MArray"
attr(,"package")
[1] ".GlobalEnv"
> ma
An object of class "MArray"
Slot "marray":
```

³Note that the usage of slots to define the representation of the class is the preferred way to define a class; the representation function is deprecated from version 3.0.0 and should be avoided.

```
В
                             С
                               D
                                           Ε
probe1
         6.868 17.559 14.59489 16.793
                                       9.177 11.991
       10.918 11.949 13.91068 9.486
                                       8.733
probe2
                                              6.940
probe3
        5.822 6.894 10.37282 11.938 13.485 11.706
       17.976 -1.073 0.05324 9.731 12.783
probe4
                                               4.353
        11.648 15.625 13.09913 3.115
                                       6.556 17.165
probe5
probe6
        5.898 9.775
                      9.71936 7.925
                                       6.463 19.902
probe7
        12.437 9.919 9.22102 8.029 11.823
                                              8.164
probe8
        13.692 14.719
                       2.64624 9.703 13.843
                                               4.779
probe9
        12.879 14.106 7.60925 15.500
                                       9.438 12.849
probe10 8.473 12.970 12.08971 13.816 14.406 9.325
Slot "fmeta":
        geneId pathway
probe1
             1
                     Ζ
probe2
             2
                     Μ
probe3
             3
                     Μ
probe4
             4
                     Ε
                     Τ
probe5
             5
probe6
             6
                     L
probe7
             7
                     N
probe8
             8
                     F
probe9
             9
                     F
                     Р
probe10
            10
Slot "pmeta":
  sampleId condition
Α
         1
                  WT
В
         2
                  WT
C
         3
                  WT
D
         4
                 MUT
Ε
         5
                 MUT
F
         6
                 MUT
```

To access individual slots, we need to use the ${\tt Q}$. This is equivalent of using the ${\tt \$}$ for a list.

```
> ma@pmeta
sampleId condition
```

Α	1	WT
В	2	WT
C	3	WT
D	4	MUT
E	5	MUT
F	6	MUT

But this is something we do not want a user to do. To access a slot like this, one needs to know its name, i.e. the underlying plumbing of the class. This breaks the notion of encapsulation. Instead, the developer will provide the user with specific accessor methods (see section 5.2) to extract (or update using a replace method, section 5.5) specific slots.

5 MArray methods

Before proceeding, we need to explain the concept of generic function. A generic function, or generic for short, is a function that dispatches methods to the appropriate class-specific implementation. A method do will implement behaviour for a specific class A, while another implementation of do, will define another behaviour for class B. The generic do is the link between the class and its dedicated implementation. If we have do(a) (where a is of class A), than the generic will make sure that the A-specific code of do will be executed.

Before we define a method with setMethod, we will always want to first check if such a method does not exists (in which case there is already a generic function), as illustrated with the show method in section 5.1. If it is the case, we write our new methods. If not, we first create the generic with setGeneric and then proceed with the method.

5.1 The show method

The **show** method (it is a method, as it exhibits custom behaviour depending on the class of its argument) is a very helpful one. It allows to define custom summary view of an object when we type its name in the console, instead of having all its (possibly very long content) displayed.

```
> show
standardGeneric for "show" defined from package "methods"
function (object)
```

```
standardGeneric("show")
<bytecode: 0x33d6e28>
<environment: 0x3cdaf30>
Methods may be defined for arguments: object
Use showMethods("show") for currently available ones.
(This generic function excludes non-simple inheritance; see ?setIs)
> isGeneric("show")

[1] TRUE
> hasMethod("show")
[1] TRUE
```

As there is already a show generic function, we can immediately proceed with the method definition using the setMethod function. To do so we need a few things. First, we need to know for what class we implement the specific show method; this is the MArray class and will be passed as the signature argument in setMethod. We also need to know the argument names that are defined in the generic. These must match exactly, as we write a method for that sepcific generic. The arguments can be found by just typing the name of the generic (as in the previous) code chunk, look at its documentation or directly ask for the arguments with args(show). We see that there is only one argument, object (naming the first argument of a generic object is a widely applied convention). This is the same name that we will have to use when writing the defintion of our method.

5.2 Accessors

As mentioned above, we want to provide customised and controlled access to the class slots. This does not prevent us, as developers, to use the @ accessor, but does not force others to know the implementation details.

Let's create an accessor for the marray slot and call the accessor marray. There is not harm in naming the slot and its accessor but there is no constraint in doing so. There is no such method or generic; just typing marray with tell you that no such object is found. Below, we create a new generic function with setGeneric. We define the name of our new generic as well as the name of the argument(s) that will have to be re-used when defining our class-specific method.

```
> setGeneric("marray", function(object) standardGeneric("marray"))
[1] "marray"
```

We now proceed in the same way as previously, using **setMethod**. The definition of our method (i.e. the actual code that will be executed) is very basic and of course uses the **@**.

```
> setMethod("marray", "MArray",
            function(object) object@marray)
[1] "marray"
> marray(ma)
                             C
             Α
                    В
                                    D
                                           F.
                                                  F
         6.868 17.559 14.59489 16.793 9.177 11.991
probe1
probe2
        10.918 11.949 13.91068 9.486
                                       8.733
         5.822 6.894 10.37282 11.938 13.485 11.706
probe3
        17.976 -1.073 0.05324 9.731 12.783
probe4
        11.648 15.625 13.09913 3.115
                                      6.556 17.165
probe5
probe6
        5.898
                9.775
                      9.71936
                               7.925
                                      6.463 19.902
probe7
        12.437
                9.919
                      9.22102
                               8.029 11.823
                                              8.164
probe8
        13.692 14.719
                      2.64624 9.703 13.843
                                              4.779
        12.879 14.106 7.60925 15.500 9.438 12.849
probe9
probe10
        8.473 12.970 12.08971 13.816 14.406
```

If we change the underlying implementation by changing the name of the slot or using an environment instead of a matrix, the ma@marray is going to break.

However, when providing accessors, we can echo the changes in the accessor implementation without affecting the users' behaviour or existing scripts.

5.3 The sub-setting operation

Let's now encapsulate the sub-setting of an MArray object in a proper method so facilitate this simple operation. In R, the default subsetting operator is [.

5.4 The validity method

5.5 A replace method

5.6 The dim method

Let's also implement a dim method that will tell us the dimensions of the marray matrix. Let's start by looking at dim to see if it is a method at all.

```
function (x) .Primitive("dim")
```

6 Introspection

```
> slotNames(ma)

[1] "marray" "fmeta" "pmeta"

> getClass("MArray")

Class "MArray" [in ".GlobalEnv"]

Slots:

Name: marray fmeta pmeta
Class: matrix data.frame data.frame
```

Finding stuff with showMethods, getMethod().

7 Conclusion

A little overhead in the beginning for improved stability and usability in the future. This however implied to think about the needs of the users role before you write code.

Session information

All software and respective versions used to produce this document are listed below.

- R Under development (unstable) (2013-06-16 r62969), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_GB.UTF-8, LC_NUMERIC=C, LC_TIME=en_GB.UTF-8, LC_COLLATE=en_GB.UTF-8, LC_MONETARY=en_GB.UTF-8, LC_MESSAGES=en_GB.UTF-8, LC_PAPER=C, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_GB.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: codetools 0.2-8, knitr 1.2
- Loaded via a namespace (and not attached): digest 0.6.3, evaluate 0.4.3, formatR 0.7, stringr 0.6.2, tools 3.1.0