# SimpleS3

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**Abstract** Writing functions in R is an important skill for an R programmer. Many R programmers are adept at creating their own functions, but do not use R's S3 methods. S3 methods allow for functions to be generalised across different classes and are easy to implement, once you know how. This guide is simple and targeted, serving to explain S3 methods so that users can create their own.

#### Note

This file is only a basic article template. For full details of *The R Journal* style and information on how to prepare your article for submission, see the Instructions for Authors.

#### Introduction

A standard principle of programming is DRY - Don't Repeat Yourself [ref]. Under this axiom, the copying and pasting of the same or similar code (copypasta), is avoided and replaced with a function. Having one function to replace several of the same or similar coded sections simplifies code maintenance as it means that only one section of code needs to be maintained, instead of several. This means that if the code breaks, then one simply needs to update the function, rather than finding all of the coded sections that are now broken.

The function summary() is an S3 method. When applied to an object of class data.frame, summary shows descriptive statistics (Mean, SD, etc.) for each variable. For example, iris is of class data.frame:

```
class(iris)
#> [1] "data.frame"
```

So applying summary to iris gives us summary information relevant to a dataframe

summary(iris)

```
Sepal.Width Petal.Length
                                                  Petal.Width
#>
    Sepal.Length
  Min. :4.300 Min. :2.000 Min. :1.000 Min. :0.100
#>
#>
   1st Qu.:5.100    1st Qu.:2.800    1st Qu.:1.600    1st Qu.:0.300
#>
   Median :5.800
                   Median :3.000 Median :4.350
                                                  Median :1.300
   Mean :5.843
                   Mean :3.057
                                  Mean :3.758
                                                  Mean :1.199
#>
                   3rd Qu.:3.300 3rd Qu.:5.100 3rd Qu.:1.800 Max. :4.400 Max. :6.900 Max. :2.500
#>
   3rd Qu.:6.400
   Max. :7.900
#>
#>
         Species
#> setosa :50
#>
  versicolor:50
   virginica:50
#>
#>
#>
#>
```

summary also performs differently when applied to different object. In fact, you can find all the classes that work with an S3 method by typing the following:

methods(summary)

#> [13] summary.lm

```
#> [15] summary.manova
                                      summary.matrix
#> [17] summary.mlm*
                                      summary.nls*
#> [19] summary.packageStatus*
                                      summary.PDF_Dictionary*
#> [21] summary.PDF_Stream*
                                      summary.POSIXct
#> [23] summary.POSIXlt
                                      summary.ppr*
#> [25] summary.prcomp*
                                      summary.princomp*
#> [27] summary.proc_time
                                      summary.srcfile
#> [29] summary.srcref
                                      summary.stepfun
#> [31] summary.stl*
                                      summary.table
#> [33] summary.tukeysmooth*
#> see '?methods' for accessing help and source code
   There's over 40 different methods!
   We can use summary on a linear model, for example:
lm_iris <- lm(Sepal.Length ~ Sepal.Width, data = iris)</pre>
summary(lm_iris)
#>
#> Call:
#> lm(formula = Sepal.Length ~ Sepal.Width, data = iris)
#>
#> Residuals:
#>
      Min
               1Q Median
                               3Q
#> -1.5561 -0.6333 -0.1120 0.5579 2.2226
#>
#> Coefficients:
       Estimate Std. Error t value Pr(>|t|)
#>
#> (Intercept) 6.5262 0.4789 13.63 <2e-16 ***
#> Sepal.Width -0.2234 0.1551 -1.44
                                            0.152
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Residual standard error: 0.8251 on 148 degrees of freedom
#> Multiple R-squared: 0.01382,
                                  Adjusted R-squared: 0.007159
#> F-statistic: 2.074 on 1 and 148 DF, p-value: 0.1519
```

summary produces a description of the linear model, describing how it was called (call), as well as the residuals, coefficients, t-values, p-values,  $R^2$ , and more. This output is **completely** different to the information output from summary used for the iris dataframe.

summary.loess\*

So how does the same function, summary perform differently for different objects? The answer is that R is sneaky, and *hides* information. There are in fact, many different summary functions. For example:

- summary.lm
- summary.data.frame
- summary.Date
- summary.matrix

Being an S3 method, summary calls the appropriate function based upon the class of the object it operates on. So using summary on an object of class "Date" will evoke the function, summary. Date. But all you need to do is type summary, and the S3 method does the rest.

To further illustrate, using summary on the iris data will actually call the function summary.data.frame, since iris is of class data.frame. We can find the class of an object using class

```
class(iris)
#> [1] "data.frame"
summary.data.frame(iris)
#> Sepal.Length Sepal.Width Petal.Length Petal.Width
#> Min. :4.300 Min. :2.000 Min. :1.000 Min. :0.100
#> 1st Qu.:5.100 1st Qu.:2.800 1st Qu.:1.600 1st Qu.:0.300
```

```
#> Median :5.800
                  Median :3.000 Median :4.350
                                                Median :1.300
                  Mean :3.057 Mean :3.758
#> Mean :5.843
                                                Mean :1.199
                  3rd Qu.:3.300 3rd Qu.:5.100
   3rd Qu.:6.400
#>
                                                3rd Qu.:1.800
                  Max. :4.400 Max. :6.900 Max. :2.500
#> Max. :7.900
#>
         Species
#> setosa :50
#> versicolor:50
#> virginica:50
#>
#>
#>
   which is the same as
summary(iris)
#>
   Sepal.Length
                  Sepal.Width
                                 Petal.Length
                                                Petal.Width
#>
   Min. :4.300
                  Min. :2.000
                                 Min. :1.000
                                                Min. :0.100
#>
                  1st Qu.:2.800
                                 1st Qu.:1.600
   1st Qu.:5.100
                                                1st Qu.:0.300
#> Median :5.800
                  Median :3.000
                                 Median :4.350
                                                Median :1.300
#> Mean :5.843
                                 Mean :3.758
                  Mean :3.057
                                                Mean :1.199
#>
   3rd Qu.:6.400
                  3rd Qu.:3.300
                                 3rd Qu.:5.100
                                                3rd Qu.:1.800
#> Max. :7.900
                  Max. :4.400 Max. :6.900 Max. :2.500
#>
         Species
#> setosa :50
#> versicolor:50
#> virginica :50
#>
#>
#>
   And using summary on the linear model object, lm_iris performs:
summary.lm(lm_iris)
#>
#> Call:
#> lm(formula = Sepal.Length ~ Sepal.Width, data = iris)
#>
#> Residuals:
              1Q Median
#> Min
                              3Q
#> -1.5561 -0.6333 -0.1120 0.5579 2.2226
#>
#> Coefficients:
             Estimate Std. Error t value Pr(>|t|)
#>
#> (Intercept) 6.5262 0.4789 13.63 <2e-16 ***
#> Sepal.Width -0.2234
                          0.1551 -1.44
                                           0.152
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#> Residual standard error: 0.8251 on 148 degrees of freedom
#> Multiple R-squared: 0.01382, Adjusted R-squared: 0.007159
#> F-statistic: 2.074 on 1 and 148 DF, p-value: 0.1519
   the same as
summary(lm_iris)
#> Call:
#> lm(formula = Sepal.Length ~ Sepal.Width, data = iris)
#>
#> Residuals:
#>
     Min
               1Q Median
                              30
                                    Max
#> -1.5561 -0.6333 -0.1120 0.5579 2.2226
```

```
#> Coefficients:
#> Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 6.5262 0.4789 13.63 <2e-16 ***
#> Sepal.Width -0.2234 0.1551 -1.44 0.152
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Residual standard error: 0.8251 on 148 degrees of freedom
#> Multiple R-squared: 0.01382, Adjusted R-squared: 0.007159
#> F-statistic: 2.074 on 1 and 148 DF, p-value: 0.1519
```

One could coerce a different method upon a different class, for example using summary.data.frame on an "lm" object:

summary.data.frame(lm\_iris)

```
coefficients
                    residuals
                                    effects
                                                        rank
#>
   Min. :-0.2234 Min. :-1.5561
                                 Min. :-71.56593
                                                   Min.
                                                        : 2
#>
   1st Qu.: 1.4640
                  1st Qu.:-0.6333
                                  1st Qu.: -0.65192
                                                   1st Qu.:2
                  Median :-0.1120
  Median : 3.1514
#>
                                  Median : -0.00897
                                                   Median :2
#> Mean : 3.1514 Mean : 0.0000
                                 Mean : -0.42040 Mean :2
  3rd Qu.: 4.8388
                  3rd Qu.: 0.5579 3rd Qu.: 0.61051
#>
                                                   3rd Qu.:2
#> Max. : 6.5262 Max. : 2.2225 Max. : 2.15225 Max. :2
                  assign qr.Length qr.Class qr.Mode df.residual
#> fitted.values
#> Min. :5.543 Min. :0.00 300
                                                     Min. :148
                                  -none- numeric
#>
  1st Qu.:5.789 1st Qu.:0.25 2
                                     -none- numeric
                                                      1st Qu.:148
#> Median :5.856 Median :0.50
                                     -none- numeric Median :148
#> Mean :5.843 Mean :0.50
                                     -none- numeric Mean :148
#> 3rd Qu.:5.901 3rd Qu.:0.75 1
                                     -none- numeric 3rd Qu.:148
  Max. :6.080 Max. :1.00
                                                      Max. :148
#>
#>
   xlevels
                call
                             terms
#>
  Length:0
             Length:3
                          Length:3
  Class:list Class:call Class1:terms
#>
  Mode :list Mode :call Class2:formula
#>
                           Mode :call
#>
#>
#>
   model.Sepal.Length model.Sepal.Width
#>
                   Min. :2.000000
#>
   Min. :4.300000
   1st Qu.:5.100000
                    1st Qu.:2.800000
#>
#>
   Median :5.800000
                    Median :3.000000
   Mean :5.843333
                    Mean :3.057333
#>
#>
   3rd Qu.:6.400000
                    3rd Qu.:3.300000
  Max. :7.900000
                    Max. :4.400000
#>
```

However the output may be a bit confusing.

To summarize, the important feature of S3 methods worth noting is that only the **first part**, summary, is required to be used on these objects of different classes.

### Why hide the text?

Hiding the trailing text after the . avoids the need to use a different summary function for every class. This means that one does not need to remember to use summary.lm for linear models, or summary.data.frame for data frames, or summary.aProposterousClassOfObject. By using S3 methods, cognitive load is reduced - you don't have to think as much to remember what class an object is and the commands are more intuitive. To get a summary of most objects, use summary, to plot most objects, use plot. Perhaps the most nifty feature of all is that a user can create their own S3 methods using the same functions such as summary and plot. This means a user can create their own special class of object

```
test_class <- 1:10
class(test_class) <- "myclass"
class(test_class)</pre>
```

```
#> [1] "myclass"
```

and then write their own S3 method for it - e.g., summary.myclass or plot.myclass, each proiding appropriate summary information, or nice plots, for that object.

## How to make your own S3 method?

Creating your own S3 method is not particularly difficult and is usually highly practical. A use case scenario for creating an S3 method is now discussed.

The Residual Sums of Squares (RSS),  $\sum (Y_i - \hat{Y})^2$  is a useful metric for determining model accuracy for continuous outcomes. For example, for a Classification and Regression Tree

```
library(rpart)
fit.rpart <- rpart(Sepal.Width ~ Sepal.Length + Petal.Length + Petal.Width + Species, data = iris)
   The RSS is calculated as
print_rss <- sum(residuals(fit.rpart)**2)
   One might be inclined to write a function to perform this task
rss <- function(x){
   sum(residuals(x)**2)
}
rss(fit.rpart)
#> [1] 10.17245
   However, there are many different decision tree models that one would like to compare, say boosted regression trees (BRT), and random forests (RF). The same code will not work:
library(randomForest)
#> randomForest 4.6-12
#> Type rfNews() to see new features/changes/bug fixes.
```

In this case, one could write three functions, one for each decision tree method: "rss\_rpart", "rss\_brt", and "rss\_rf". But to avoid having three functions and instead use just one, one could place all three functions inside of one function, using an if-then-else clause to direct the object of the appropriate class to the appropriate method. This is what I shall call a "Poor man's S3 method".

```
\begin{Schunk} \begin{Sinput} dt_rss <- function (x){
if ("rpart" %in% class(x)) {
  result <- sum((residuals(x)**2))
  return(result)
}
else if ("gbm" %in% class(x)) {
  result <- sum(x$residuals**n2)
  return(result)
}
else if ("randomForest" %in% class(x)) {</pre>
```

```
temp <- xy - xpredicted
  result <- sum(temp**2)
  return(result)
  }
  else warning(paste(class(x), "is not of an rpart, gbm, or randomForest object")) } \end{Sinput}
\end{Schunk}
  Here it is in action

dt_rss(fit.rpart)

#> [1] 10.17245
  The RSS method works, and if it is applied to a class that is not known, a special message is provided

fit.lm <- lm(Sepal.Width ~ Species, data = iris)

dt_rss(fit.lm)

#> Warning in dt_rss(fit.lm): lm is not of an rpart, gbm, or randomForest
#> object
```

The "poor man's S3 method" does what it needs to do. However, the cluster of ifelses is more difficult to read, manage, and debug, and is not as elegant and not as modular as a real S3 method. So let us create an S3 method.

First define the S3 method with UseMethod()

```
rss <- function(x) UseMethod("rss")</pre>
```

This creates the building block of an S3 method, the "root", if you will.

Here we have specified that our method will be called rss. Now we need to create the special cases of rss - the methods rss.rpart, rss.gbm, and rss.randomForest, where the sections of code after rss. are the classes of object we want them to work on.

```
\begin{Schunk} \begin{Sinput} rss.rpart <- function(x){
    sum((residuals(x)**2))
}
rss.gbm <- function(x){
    sum(x$residuals**2)
}
rss.randomForest <- function(x){
    res <- xy - xpredicted
    sum(res**2)
} \end{Sinput} \end{Schunk}</pre>
```

A default method can also be created - rss.default - which, as the name suggests, is the default method when other classes are not present.

```
rss.default <- function(x, ...){

warning(paste("RSS does not know how to handle object of class", class(x), "and can only be used on classe"}
```

In this case a warning is issued, to let the user know that the object class they were using was not appropriate.

We can now apply the rss method to an rpart model

```
rss(fit.rpart)
#> [1] 10.17245
```

Also observe what happens when the object used is not of the decision tree classes

```
rss(lm.fit)
#> Warning in rss.default(lm.fit): RSS does not know how to handle object of
#> class function and can only be used on classes rpart, gbm, and randomForest
```

This guide to S3 methods was written to provide R users with the minimal amount of information to start building their own S3 methods. For a more complete treatment on S3 methods, see Advanced-R, R Packages, and this blog, this resource.

#### **Extras**

For the uninitiated, you may find the class of an object using the command, class(), on the object. For example:

```
x <- c(1, 2, 3, 4, 5)
x
#> [1] 1 2 3 4 5
class(x)
#> [1] "numeric"
    Here, showing that the object x is of class numeric.
    str() can also provide more information:
str(x)
#> num [1:5] 1 2 3 4 5
    In this case, revealing that x is numeric, showing its contents.
```

#### Idea

Idea: Make a cheatsheet / infographic for writing functions in R, and for making s3 methods. Let's call it "The Anatomy of S3 Methods"

```
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