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# Data Manipulation with R

Perform group-wise data manipulation and deal with large datasets using R efficiently and effectively

Jaynal Abedin



**BIRMINGHAM - MUMBAI** 

#### Data Manipulation with R

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Firstly, I would like to thank my teachers, who introduced me to this wonderful open source technology during my undergraduate years. Then I would like to express my gratitude to my family, friends, colleagues, and well wishers, who always motivated me to contribute to this technology. Last but not least, I would like to express my deepest gratitude to Packt Publishing and its team, who gave me the opportunity to write this book. Finally, I am grateful to all of the reviewers for their time and constructive suggestions.

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Dedicated to my late grandmother
-Jaynal Abedin

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# **Preface**

This book, *Data Manipulation with R*, is aimed at giving intermediate to advanced level users of R (who have knowledge about datasets) an opportunity to use state-of-the-art approaches in data manipulation. This book will discuss the types of data that can be handled using R and different types of operations for those data types. Upon reading this book, readers will be able to efficiently manage and check the validity of their datasets with the effective use of R programming, including specialized packages for data management. Readers will come to know about the split-apply-combine strategy, which is the state-of-the-art approach in data management. This book ends with an introduction to how R can be utilized with different database software.

#### What this book covers

Chapter 1, R Data Types and Basic Operations, discusses the different types of data used in R and their basic operations. Before introducing the data types in this chapter, we will highlight what an object in R is and its mode and class. The mode of an object could be either numeric, character, or logical, whereas its class could be vector, factor, list, data frame, matrix, array, or others. This chapter also highlights how to deal with objects in different modes and how to convert from one mode to another and what caution should be taken during conversion. Missing values in R and how to represent missing character and numeric data types are also discussed here. Along with the data types and basic operations, this chapter sheds light on another important aspect, which is almost never mentioned in other text books — the object naming convention in R. We talk about popular object-naming conventions used in R.

Chapter 2, Basic Data Manipulation, introduces some special features that we need to consider during data acquisition. Then, an important aspect of factor manipulation will be discussed, especially when subsetting a factor variable and how to remove unused factor levels. Date processing is also covered using an efficient R package: lubridate. Dealing with the date variable using the lubridate package is much more efficient than any other existing packages that are designed to work with the date variable. Also, string processing will be highlighted and the chapter ends with a description of subscripting and subsetting.

Chapter 3, Data Manipulation Using plyr, introduces the state-of-the-art approach called split-apply-combine to manipulate datasets. Data manipulation is an integral part of data cleaning and analysis. For large data, it is always preferable to perform the operations within the subgroup of a dataset to speed up the process. In R, this type of data manipulation can be done with base functionality, but for large data it requires considerable amount of coding and eventually takes more processing time. In the case of large datasets, we can split the data and perform the manipulation or analysis and then again combine them into a single output. This chapter contains a discussion on the different functions in the plyr package that are used for group-wise data manipulation and also for data analysis.

Chapter 4, Reshaping Datasets, deals with the orientation of datasets. Reshaping data is a common and tedious task in real-life data manipulation and analysis. A dataset might come with different levels of grouping and we need some reorientation to perform certain types of analysis. To perform statistical analysis, we sometimes require wide data and sometimes long data, and in that case we need to be able to fluently and fluidly reshape data to meet the requirements. Important functions from the reshape package will be discussed in this chapter with examples.

Chapter 5, R and Databases, talks about dealing with database software and R. One of the major problems in R is that its memory is bound by RAM, and that is why working with a dataset requires the data to be smaller than its memory. But in reality, the dataset is larger than the capacity of RAM and sometimes the length of arrays or vectors exceeds the maximum addressable range. To overcome these two limitations, R can be utilized with databases. Interacting with databases using R and dealing with large datasets with specialized packages and data manipulation with sqldf will be discussed with examples in this chapter.

Bibliography, provides a list of citations used in the book.

# What you need for this book

Readers are expected to have basic knowledge of R and some knowledge of statistical data. To run the examples from this book, R should be installed, and it can be found at http://www.r-project.org. The example files are produced on R 2.15.2 and R 3.0.1.

#### Who this book is for

This book is for intermediate to advanced level users of R who have knowledge about datasets. Also, this book is for those who regularly deal with different research data, including but not limited to public health, business analysis, and the machine-learning community.

#### **Conventions**

In this book, you will find a number of styles of text that distinguish between different kinds of information. Here are some examples of these styles, and an explanation of their meaning.

Code words in text, database table names, folder names, filenames, file extensions, pathnames, dummy URLs, user input, and Twitter handles are shown as follows: "Once we have an R object we can easily assess its mode by using mode ()."

A block of code is set as follows:

```
num.obj <- seq(from=1,to=10,by=2)
logical.obj<-c(TRUE,TRUE,FALSE,TRUE,FALSE)
character.obj <- c("a","b","c")

is.numeric(num.obj)
[1] TRUE

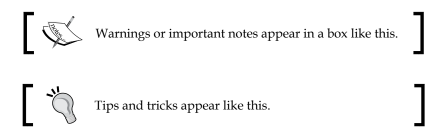
is.logical(num.obj)
[1] FALSE

is.character(num.obj)
[1] FALSE</pre>
```

When we wish to draw your attention to a particular part of a code block, the relevant lines or items are set in bold:

```
# Calling xlsx library
library(xlsx)
# importing xlsxanscombe.xlsx
anscombe xlsx <- read.xlsx2("xlsxanscombe.xlsx", sheetIndex=1)</pre>
```

**New terms** and **important words** are shown in bold. Words that you see on the screen, in menus or dialog boxes for example, appear in the text like this: "Click on the **Add...** button and select an appropriate ODBC driver and then locate the desired file and give a data source name."



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# R Data Types and Basic Operations

R is an object-oriented programming language that is a variation of the S language, and was written by Ross Ihaka and Robert Gentleman (hence the name **R**), the R Core Development Team, and an army of volunteers. What can we do using R? The answer is we can do anything we can think of that is logical and/or structural. With R, we can perform data processing, write functions, produce graphs, perform complex data analysis, and also produce our own customized packages (a collection of functions for performing specified tasks) to solve specific problems. We can develop up-to-date statistical techniques through R packages, and most importantly, R is open source and a freely available software and it will remain free.

Assuming you have preliminary knowledge on where to get R and how to install it, we will discuss R data types and different operations related to data types. But before introducing data types, we will briefly discuss R objects, modes, and classes because whenever we work in R, we have to deal with these three terminologies frequently. In this chapter, we are going to discuss the following:

- Modes and classes of R objects
- R object structure and mode conversion
- Vector
- Factor and its types
- Data frames, matrices, and arrays
- Lists
- Missing values in R

# Modes and classes of R objects

Whatever we do in R, R stores as objects. An R object is anything that can be assigned to a variable of interest. This could be a single number or a set of numbers, characters, and special characters; for example, TRUE, FALSE, NA, NAN, and Inf. Also, these can be the things that are already defined in R as functions, such as seq (to generate a sequence of numbers with a specified increment), names (to extract names such as variable names from a dataset), row.names (to extract the row names of the data, if any), or col.names (this is equivalent to names and it extracts column names from a matrix or data frame). Some of the examples of R objects are as shown in the following code:

```
# Constant
2
[1] 2
"July"
[1] "July"
NULL
NULL
NA
[1] NA
NaN
[1] NaN
Inf
[1] Inf
# Object can be created from existing object
# to make the result reproducible means every time we run the
# following code we will get the same results # we need to set
# a seed value
set.seed(123)
rnorm(9)+runif(9)
[1] -0.2325549  0.7243262  2.4482476  0.7633118  0.7697945
2.7093348 1.1166220 -0.5565308 -0.1427868
```

# "

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One important thing about objects in R is that if we do not assign an object to any variable, we will not able to re-use it and it does not store the object internally. In the preceding example, all are different objects, but they are not assigned to any variable so they are not stored and we cannot use them later until we enter the object's value itself. So whenever we deal with an object, we will assign it to an appropriate variable, and interestingly the assigned variable is also an object in R!

To assign an object in R to a variable, we can define the variable name in various ways, such as lowercase, uppercase, a combination of upper and lowercase, or even a combination of uppercase, lowercase, and a number and/or a dot; but there are some rules to define variable names. For example, the name cannot start with numbers; it will start with a character or underscore. There is no special character allowed in variable names, such as @, #, \$, and \*. Though R does not have a standard guideline for naming conventions, according to Bååth (in the paper *The State of Naming Conventions in R*, which can be found at http://journal.r-project.org/archive/2012-2/RJournal\_2012-2\_Baaaath.pdf), the most popular function naming convention is lower CamelCase while the most popular naming convention for arguments is period separated. For a variable name, we can use the same naming convention as that of arguments, but again there is no strict rule for naming conventions in R. The following table is reconstructed from the same article by Bååth to give you an idea of the different naming conventions used in R and their popularity:

Object type	Naming conventions	Percentage	
Function			
	lowerCamelCase	55.5	
	period.separated	51.8	
	underscore_separated	37.4	
	singlelowercaseword	32.2	
	_OTHER.conventions	12.8	
	UpperCamelCase	6.9	
Parameter (argument)			
	period.separated	82.8	
	lowerCamelCase	75.0	
	underscore_separated	70.7	
	singlelowercaseword	69.6	
	_OTHER.conventions	9.7	
	UpperCamelCase	2.4	

Once we store the R object into a variable, it is still treated as an R object. Each and every object in R has some attributes to describe the nature of the information contained in it. The mode and class are the most important attributes of an R object. Commonly encountered modes of an individual R object are numeric, character, and logical. When we work with data in R, problems might arise due to incorrect operations in incorrect object modes. So before working with data, we should study the mode; we need to know what type of operation is applicable.

The mode function returns the mode of R objects. The following example code describes how we can investigate the mode of an R object:

```
# Storing R object into a variable and then viewing the mode
num.obj <- seq(from=1,to=10,by=2)
mode(num.obj)
[1] "numeric"

logical.obj<-c(TRUE,TRUE,FALSE,TRUE,FALSE)
mode(logical.obj)
[1] "logical"

character.obj <- c("a","b","c")
mode(character.obj)
[1] "character"</pre>
```

For the numeric mode, R stores all numeric objects into either a 32-bit integer or double-precision floating point.

If an R object contains both numeric and logical elements, the mode of that object will be numeric and in that case the logical element automatically gets converted to numeric. The logical element TRUE converts to 1 and FALSE converts to 0. On the other hand, if any R object contains a character element along with both numeric and logical elements, it automatically converts to the character mode. Let's have a look at the following code:

The mode() function is not the only way to test R object modes; there are alternative ways too, which are is.numeric(), is.charater(), and is.logical(), as shown in the following code. The output of these functions is always logical.

```
num.obj <- seq(from=1,to=10,by=2)
logical.obj<-c(TRUE,TRUE,FALSE,TRUE,FALSE)
character.obj <- c("a","b","c")

is.numeric(num.obj)
[1] TRUE
is.logical(num.obj)
[1] FALSE
is.character(num.obj)
[1] FALSE</pre>
```

Other than these three modes (numeric, logical, and character) of objects, another frequently encountered mode is function; for example:

```
mode(mean)
[1] "function"
# Also we can test whether "mean" is function or not as follows
is.function(mean)
[1] TRUE
```

The class() function provides the class information of an R object. The primary purpose of the class() function is to know how different functions, including generic functions, will work. For example, with the class information, the generic function print or plot knows what to do with a particular R object. To assess the class information of the object created earlier, we can use the class() function. Let's have a look at the following code:

```
num.obj <- seq(from=1, to=10, by=2)
logical.obj<-c(TRUE, TRUE, FALSE, TRUE, FALSE)
character.obj <- c("a", "b", "c")

class(num.obj)
[1] "numeric"

class(logical.obj)
[1] "logical"

class(character.obj)
[1] "character"</pre>
```

Although we can easily assess the mode and class of an R object through mode () and class(), there is another collection of R commands that are also used to assess whether a particular object belongs to a certain class. These functions start with is., for example; is.numeric(), is.logical(), is.character(), is.list(), is.factor(), and is.data.frame(). As R is an object-oriented programming language, there are many functions (collectively known as generic functions) that will behave differently depending on the class of that particular object.

The mode of an object tells us how it's stored. It could happen that two different objects are stored in the same mode with different classes. How those two objects are printed using the print command is determined by its class; for example:

```
# Output omitted due to space limitation
num.obj <- seq(from=1,to=10,by=2)
set.seed(1234) # To make the matrix reproducible
mat.obj <- matrix(runif(9),ncol=3,nrow=3)
mode(num.obj)
mode(mat.obj)
class(num.obj)
class(mat.obj)
# prints a numeric object
print(num.obj)
# prints a matrix object
print(mat.obj)</pre>
```

Like character and numeric, there is another method you can use to store data when the data is categorical in nature. In categorical data, we usually have some unique values and their corresponding labels. To store this type of object in R, we use the class factor, which allows less storage location because it is required to store only unique levels once.

Interestingly, once we try to see the mode of a factor object, it always shows numeric even if it displays character data. For example:

```
character.obj <- c("a","b","c")
character.obj
[1] "a" "b" "c"

is.factor(character.obj)
[1] FALSE

# Converting character object into factor object using as.factor()
factor.obj <- as.factor(character.obj)</pre>
```

```
factor.obj
[1] a b c
Levels: a b c

is.factor(factor.obj)
[1] TRUE

mode(factor.obj)
[1] "numeric"

class(factor.obj)
[1] "factor"
```

We have to be careful when dealing with the factor class data in R. The important thing to remember is that for vectors (we will discuss vectors in the *Vector* section in this chapter), the class and mode will always be numeric, logical, or character. On the other hand, for matrices and arrays (we will discuss matrices and arrays in the *Factor and its type* section in this chapter), a class is always a matrix or array, but its mode can be numeric, character, or logical.

# R object structure and mode conversion

When we work with any statistical software, such as R, we rarely use single values for an object. We need to know how we can handle a collection of data values (for example, the age of 100 randomly selected diabetic patients) along with what type of objects need to store those data values. In R, the most convenient way to store more than one data value is vector (a collection of data values stored in a single object is known as a vector; for example, storing the ages of 100 diabetic patients in a single object). In fact, whenever we create an R object, it stores the values as a vector. It could be a single-element vector or multiple-element vector. The num.obj vector we have created in the previous section is a kind of vector comprising of numeric elements.

One of the simplest ways to create a vector in R is to use the c() function. For example:

```
# creating vector of numeric element with "c" function
num.vec <- c(1,3,5,7)
num.vec
[1] 1 3 5 7
mode(num.vec)
[1] "numeric"
class(num.vec)
[1] "numeric"
is.vector(num.vec)
[1] TRUE</pre>
```

If we create a vector with mixed elements (character and numeric), the resulting vector will be a character vector. For example:

```
# Vector with mixed elements
num.char.vec <- c(1,3,"five",7)
num.char.vec
[1] "1" "3" "five" "7"
mode(num.char.vec)
[1] "character"
class(num.char.vec)
[1] "character"
is.vector(num.char.vec)
[1] TRUE</pre>
```

We can create a big new vector by combining multiple vectors, and the resulting vector's mode will be character if any element of any vector contains a character. The vector could be named or without a name; in the previous example, vectors were without names. The following example shows how we can create a vector with the name of each element:

```
# combining multiple vectors
comb.vec <- c(num.vec,num.char.vec)
mode(comb.vec)
[1] "character"

# creating named vector
named.num.vec <- c(x1=1,x2=3,x3=5)
named.num.vec
x1 x2 x3
1 3 5</pre>
```

The name of the elements in a vector can be assigned separately using the names() command. In R, any single constant is also stored as a vector of the single element. For example:

```
# vector of single element
unit.vec <- 9
is.vector(unit.vec)
[1] TRUE</pre>
```

R has six basic storage types of vectors and each type is known as an atomic vector. The following table shows the six basic vector types, their mode, and the storage mode:

Type	Mode	Storage mode
logical	logical	logical
integer	numeric	integer
double	numeric	double
complex	complex	complex
character	character	character
raw	raw	raw

Other than vectors, there are different storage types available in R to handle data with multiple elements, which are matrix, dataframe, and list. We will discuss each of these types in the subsequent sections.

To convert the object mode, R has user friendly functions that can be depicted as follows: as.x. Here, x could be numeric, logical, character, list, data.frame, and so on. For example, if we need to perform a matrix operation that requires the numeric mode and the data is stored in some other mode, the operation cannot be performed. In that case, we need to convert that data into the numeric mode.

In the following example, we will see how we can convert an object's mode:

```
# creating a vector of numbers and then converting it to logical
# and character
numbers.vec <- c(-3,-2,-1,0,1,2,3)
numbers.vec
[1] -3 -2 -1 0 1 2 3
num2char <- as.character(numbers.vec)
num2char
[1] "-3" "-2" "-1" "0" "1" "2" "3"
num2logical <- as.logical(numbers.vec)
num2logical
[1] TRUE TRUE TRUE FALSE TRUE TRUE TRUE</pre>
```

# creating character vector and then convert it to numeric and logical

```
char.vec <- c("1", "3", "five", "7")
char.vec
[1] "1"
         "3"
                  "five" "7"
char2num <- as.numeric(char.vec)</pre>
Warning message:
NAs introduced by coercion
char2num
[1] 1 3 NA 7
char2logical <- as.logical(char.vec)</pre>
char2logical
[1] NA NA NA NA
# logical to character conversion
logical.vec <- c(TRUE, FALSE, FALSE, TRUE,</pre>
                                                TRUE)
logical.vec
     TRUE FALSE FALSE TRUE
logical2char <- as.character(logical.vec)</pre>
logical2char
[1] "TRUE" "FALSE" "FALSE" "TRUE"
                                      "TRUE"
```

Note that when we convert the numeric mode to the logical mode, only 0 (zero) gets false and all the other values get TRUE. Also, if we convert a character object to numeric, it produces numeric elements and NA (if any actual character is present), and a warning will be issued. Importantly, R does not convert a character object into a logical object, but if we try to do this, all the resulting elements will be NA. However, logical objects get successfully converted to character objects. Finally, we can say that any object can be converted to a character without any warning, but if we want to convert character objects to any other type, we have to be careful.

#### **Vector**

The R vector can be contiguous cells containing data. In R, the basic data storage type is vector. The vector itself could be numeric, character, and logical based on the elements. In fact, there are six types of vectors used in R. We can easily access elements of a vector through indexing. The following example shows how we can create a vector and access its individual elements and group of elements:

```
# creating a vector and accessing elements
vector1 <- c(1,3,5,7,9)
vector1
[1] 1 3 5 7 9</pre>
```

```
# accessing second elements of "vector1"
vector1[2]
[1] 3
# accessing three elements starting from second element
vector1[2:4]
[1] 3 5 7
# another way of creating vector. Here "from" is the starting point
# of the vector and "to" is the end point of the vector and "by" is
# increment
vector2 <- seq(from=2, to=10, by=2)
is.vector(vector2)
[1] TRUE</pre>
```

# Factor and its types

A factor is another important data type in R, especially when we deal with categorical variables. In an R vector, there is no limit on the number of distinct elements, but in factor variables, it takes only a limited number of distinct elements. This type of variable is usually referred to as a categorical variable during data analysis and statistical modeling. In statistical modeling, the behavior of a numeric variable and categorical variable is different, so it is important to store the data correctly to ensure valid statistical analysis.

In R, a factor variable stores distinct numeric values internally and uses another character set to display the contents of that variable. In other software, such as Stata, the internal numeric values are known as *values* and the character set is known as *value labels*. Previously, we saw that the mode of a factor variable is numeric; this is due to the internal values of the factor variable.

A factor variable can be created using the factor command; the only required input is a vector of values, which will return as a vector of factor values. The input can be numeric or character, but the levels of factor will always be a character. The following example shows how to create factor variables:

```
#creating factor variable with only one argument with factor()
factor1 <- factor(c(1,2,3,4,5,6,7,8,9))
factor1
[1] 1 2 3 4 5 6 7 8 9
Levels: 1 2 3 4 5 6 7 8 9
levels(factor1)</pre>
```

```
[1] "1" "2" "3" "4" "5" "6" "7" "8" "9"
labels(factor)
[1] "1"
labels(factor1)
[1] "1" "2" "3" "4" "5" "6" "7" "8" "9"

#creating factor with user given levels to display factor2 <- factor(c(1,2,3,4,5,6,7,8,9),labels=letters[1:9]) factor2
[1] a b c d e f g h i
Levels: a b c d e f g h i
levels(factor2)
[1] "a" "b" "c" "d" "e" "f" "g" "h" "i"
labels(factor2)
[1] "1" "2" "3" "4" "5" "6" "7" "8" "9"</pre>
```

In a factor variable, the values themselves are stored as numeric vectors, whereas the labels store only unique characters, and it stores only once for each unique character. Factors can be ordered if the ordered=T command is specified, otherwise it inherits the order of the levels specified.

A factor could be numeric with numeric levels, but direct mathematical operations are not possible with this numeric factor. Special care should be taken if we want to use mathematical operations. The following example shows a numeric factor and its mathematical operation:

```
# creating numeric factor and trying to find out mean
num.factor <- factor(c(5,7,9,5,6,7,3,5,3,9,7))
num.factor
[1] 5 7 9 5 6 7 3 5 3 9 7
Levels: 3 5 6 7 9
mean(num.factor)
[1] NA
Warning message:
In mean.default(num.factor) :
argument is not numeric or logical: returning NA</pre>
```

From the preceding example, we see that we can create a numeric factor, but the mathematical operation is not possible. And when we tried to perform a mathematical operation, it showed us a warning and produced the result NA. To perform any mathematical operation, we need to convert the factor to its numeric counterpart. One can assume that we can easily convert the factor to numeric using the as.numeric() function, but if we use the as.numeric() function, it will only convert the internal values of the factors, not the desired values.

So the conversion must be done with levels of that factor variable; optionally, we can firstly convert the factor into a character using as.character() and then use as.numeric(). The following example describes the scenario:

```
num.factor <- factor(c(5,7,9,5,6,7,3,5,3,9,7))
num.factor
[1] 5 7 9 5 6 7 3 5 3 9 7
Levels: 3 5 6 7 9
#as.numeric() function only returns internal values of the factor
as.numeric(num.factor)
[1] 2 4 5 2 3 4 1 2 1 5 4
# now see the levels of the factor
levels(num.factor)
[1] "3" "5" "6" "7" "9"
as.character(num.factor)
[1] "5" "7" "9" "5" "6" "7" "3" "5" "3" "9" "7"
# now to convert the "num.factor" to numeric there are two method
# method-1:
mean(as.numeric(as.character(num.factor)))
[1] 6
# method-2:
mean(as.numeric(levels(num.factor)[num.factor]))
[1] 6
```

#### **Data frame**

A data frame is a rectangular arrangement of rows and columns of vectors and/or factors, such as a spreadsheet in MS Excel. The columns represent variables in the data and the rows represent observations or records. In other software, such as a database package, each column represents a field and each row represents a record. Dealing with data does not mean dealing with only one vector or factor variable, rather it is the collection of variables. Each column represents only one type of data: numeric, character, or logical, and each row represents case information across all columns. One important thing to remember about R data frames is that all vectors should be of the same length. In an R data frame, we can store different types of variables, such as numeric, logical, factor, and character. To create a data frame, we can use the data.frame() command. The following example shows how to create a data frame using different vectors and factors:

```
#creating vector of different variables and then creating data frame
var1 <- c(101,102,103,104,105)</pre>
```

```
var2 < -c(25, 22, 29, 34, 33)
var3 <- c("Non-Diabetic", "Diabetic", "Non-Diabetic", "Non-Diabetic",</pre>
"Diabetic")
var4 <- factor(c("male", "male", "female", "female", "male"))</pre>
# now we will create data frame using two numeric vectors one
# character vector and one factor
diab.dat <- data.frame(var1,var2,var3,var4)</pre>
diab.dat
  var1 var2
                     var3
                           var4
1 101 25 Non-Diabetic male
2 102
        22 Diabetic male
3 103 29 Non-Diabetic female
         34 Non-Diabetic female
                Diabetic
                           male
5 105
       3.3
```

Now if we see the class of individual columns of the newly created data frame, we will see that the first two columns' classes are numeric and the last two columns' classes are factor, though initially the class of var3 was character. One thing is obvious here—when we create data frames and any one of the column's classes is character, it automatically gets converted to factor, which is a default R operation. But there is one argument, stringsAsFactors=FALSE, that allows us to prevent the automatic conversion of character to factor during data frame creation. In the following example, we will see this:

```
#class of each column before creating data frame
class(var1)
[1] "numeric"
class(var2)
[1] "numeric"
class(var3)
[1] "character"
class(var4)
[1] "factor"
```

To access individual columns (variables) from a data frame, we can use a dollar (\$) sign along with the data frame name; for example, diab.dat\$var1. There are some other ways to access variables from a data frame, such as the following:

- Data frame name followed by double square brackets with variable names within quotation marks; for example, diab.dat[["var1"]]
- Data frame name followed by single square brackets with the column index; for example, diab.dat[,1]

Besides these, there is one other way that allows us to access each of the individual variables as separate objects. The R attach() function allows us to access individual variables as separate R objects. Once we use the attach() command, we need to use detach() to remove individual variables from the working environment. Let's have a look at the following code:

```
# class of each column after creating data frame
class(diab.dat$var1)
[1] "numeric"
class(diab.dat$var2)
[1] "numeric"
class(diab.dat$var3)
[1] "factor"
class(diab.dat$var4)
[1] "factor"
# now create the data frame specifying as.is=TRUE
diab.dat.2 <- data.frame(var1, var2, var3, var4, stringsAsFactors=FALSE)
diab.dat.2
var1 var2
                 var3 var4
  101 25 Non-Diabetic male
2 102
       2.2
              Diabetic
                          male
3 103 29 Non-Diabetic female
       34 Non-Diabetic female
4 104
5 105 33 Diabetic male
class(diab.dat.2$var3)
[1] "character"
```

#### **Matrices**

A matrix is also a two-dimensional arrangement of data but it can take only one class. To perform any mathematical operations, all columns of a matrix should be numeric. However, in data frames we can store numeric, character, or factor columns. To perform any mathematical operation, especially a matrix operation, we can use matrix objects. However, in data frames, we are unable to perform certain types of mathematical operations, such as matrix multiplication. To create a matrix, we can use the matrix() command or convert a numeric data frame to a matrix using as.matrix(). We can convert the data frame that we created earlier as diab. dat to a matrix using as.matrix(), but this is not suitable to perform mathematical operations, as shown in the following example:

```
# data frame to matrix conversion
mat.diab <- as.matrix(diab.dat)</pre>
```

```
mat.diab
    var1 var2 var3
                               var4
[1,] "101" "25" "Non-Diabetic" "male"
[2,] "102" "22" "Diabetic"
                              "male"
[3,] "103" "29" "Non-Diabetic" "female"
[4,] "104" "34" "Non-Diabetic" "female"
[5,] "105" "33" "Diabetic" "male"
class(mat.diab)
[1] "matrix"
mode(mat.diab)
[1] "character"
# matrix multiplication is not possible with this newly created matrix
t(mat.diab) %*% mat.diab
Error in t(mat.diab) %*% mat.diab :
requires numeric/complex matrix/vector arguments
# creating a matrix with numeric elements only
# To produce the same matrix over time we set a seed value
set.seed(12345)
num.mat <- matrix(rnorm(9), nrow=3, ncol=3)</pre>
num.mat
                                [,3]
                 [,2]
           [,1]
[1,] 0.5855288 -0.4534972 0.6300986
[2,] 0.7094660 0.6058875 -0.2761841
[3,] -0.1093033 -1.8179560 -0.2841597
class(num.mat)
[1] "matrix"
mode(num.mat)
[1] "numeric"
# matrix multiplication
t(num.mat) %*% num.mat
          [,1]
                     [,2]
                                [,3]
[1,] 0.8581332 0.36302951 0.20405722
[2,] 0.3630295 3.87772320 0.06350551
[3,] 0.2040572 0.06350551 0.55404860
```

# **Arrays**

An array is a multiply-subscripted data entry that allows the storing of data frames, matrices, or vectors of different types. Data frames and matrices are of two dimensions only, but an array could be of any number of dimensions. Sometimes, we need to store multiple matrices or data frames into a single object; in this case, we can use arrays to store this data. The following is a simple example to store three matrices of order 2x2 in a single array object:

```
mat.array=array(dim=c(2,2,3))
# To produce the same results over time we set a seed value
set.seed(12345)
mat.array[,,1]<-rnorm(4)
mat.array[,,2]<-rnorm(4)
mat.array[,,3]<-rnorm(4)
mat.array
, , 1
          [,1]
              [,2]
[1,] 0.5855288 -0.1093033
[2,] 0.7094660 -0.4534972
, , 2
           [,1] [,2]
[1,] 0.6058875 0.6300986
[2,] -1.8179560 -0.2761841
, , 3
           [,1]
                      [,2]
[1,] -0.2841597 -0.1162478
[2,] -0.9193220 1.8173120
```

#### list

A list object is a generic R object that can store other objects of any type. In a list object, we can store single constants, vectors of numeric values, factors, data frames, matrices, and even arrays. Recalling the vectors var1, var2, var3, and var4; the data frame created using these vectors; and also recalling the array created in the *Arrays* section, we will create a list object in the following example:

```
var1 <- c(101,102,103,104,105)</pre>
var2 <- c(25,22,29,34,33)</pre>
var3 <- c("Non-Diabetic", "Diabetic", "Non-Diabetic", "Non-Diabetic",</pre>
"Diabetic")
var4 <- factor(c("male", "male", "female", "female", "male"))</pre>
diab.dat <- data.frame(var1, var2, var3, var4)</pre>
mat.array=array(dim=c(2,2,3))
set.seed(12345)
mat.array[,,1]<-rnorm(4)
mat.array[,,2]<-rnorm(4)
mat.array[,,3]<-rnorm(4)
# creating list
obj.list <- list(elem1=var1,elem2=var2,elem3=var3,elem4=var4,elem5=di
ab.dat,elem6=mat.array)
obj.list
$elem1
[1] 101 102 103 104 105
$elem2
[1] 25 22 29 34 33
$elem3
[1] "Non-Diabetic" "Diabetic"
                                     "Non-Diabetic" "Non-Diabetic"
"Diabetic"
$elem4
                  female female male
[1] male
           male
Levels: female male
```

```
$elem5
                 var3
 var1 var2
                        var4
1 101 25 Non-Diabetic male
2 102
       22 Diabetic male
3 103 29 Non-Diabetic female
4 104 34 Non-Diabetic female
5 105
      33 Diabetic male
$elem6
, , 1
         [,1] [,2]
[1,] 0.5855288 -0.1093033
[2,] 0.7094660 -0.4534972
, , 2
          [,1] \qquad [,2]
[1,] 0.6058875 0.6300986
[2,] -1.8179560 -0.2761841
, , 3
          [,1] [,2]
[1,] -0.2841597 -0.1162478
[2,] -0.9193220 1.8173120
```

To access individual elements from a list object, we could use the name of that component or use double square brackets with the index of those elements. For example, obj.list[[1]] will give the first element of the newly created list object.

#### Missing values in R

Missing values are part of the data manipulation process and we will encounter some missing values in almost every dataset. So, it is important to know how R handles missing values and how they are represented. In R, a numeric missing value is represented by NA while character missing values are represented by <NA>. To test if there is any missing value present in a dataset (data frame), we can use is.na() for each column or we can use this function in combination with the any() function. The following example shows how we can see if there are any missing values present in a dataset:

```
v1 v2

1 1 M

2 NA F

3 0 <NA>
4 1 M

is.na(missing_dat$v1)

[1] FALSE TRUE FALSE FALSE is.na(missing_dat$v2)

[1] FALSE FALSE TRUE FALSE any(is.na(missing_dat))

[1] TRUE
```

#### **Summary**

In this chapter, we firstly talked very briefly about what R is. We did not cover where to get it and how to install it as we are assuming the reader will have some preliminary knowledge in those areas. Then we introduced what R objects are and their modes and classes. We also highlighted how we can convert modes of objects using R functions such as as.numeric and as.character. Finally, we discussed different R objects, such as vector, factor, data frame, matrix, and list. The chapter ended with an introduction to how missing values are represented and dealt with in R. In the next chapter, we will discuss data manipulation with different R objects in greater detail.

## **Basic Data Manipulation**

When preparing a dataset for statistical analysis, data processing and manipulations such as checking, cleaning, and creating new variables are two of the important tasks. In this chapter, the basics of data manipulation will be discussed with examples that will give us an idea about checking a dataset and cleaning it if necessary. This chapter will deal with the following topics:

- Importing datasets from different sources
- Factor manipulations
- Numeric variables and date processing
- Character manipulations
- Subscripting and subsetting datasets

#### Acquiring data

A dataset can be stored in a computer or any other storage device in different file formats. R provides the useful facility to access different file formats through different commands. Some of the commonly used file formats are:

- Comma separated values (\*.csv)
- Text file with Tab delimited
- MS Excel file (\*.xls or \*.xlsx)
- R data object (\*.RData)

Other than the file formats mentioned in the preceding list, the dataset can be stored in another statistical software format; for example, Stata, SPSS, or SAS. In R, using the foreign library, we can acquire a dataset from other statistical software. In the following examples, we will see how we can acquire data in R from different file formats.

Firstly, we will import a <code>.csv</code> file, <code>CSVanscombe.csv</code>. This file contains four pairs of numeric variables, (x1,y1) to (x4,y4). The noticeable feature of this file is that the actual data starts from the third row and the first two rows contain a brief description about the dataset. Now we will use <code>read.csv()</code> to import the file and store it in the <code>anscombe</code> object in R, which will be a data frame, as shown in the following code:

```
# Before running the following command we need to set the data
# location using setwd(). For example setwd("d:/chap2").
anscombe <- read.csv("CSVanscombe.csv",skip=2)</pre>
```

Note that in the preceding code, the skip=2 command is used, which tells R that the actual data starts from the third row.

If a .csv file contains both numeric and character variables and we use read. csv(), the character variables get automatically converted to the factor type. We can prevent character variables from this automatic conversion to factor by specifying stringsAsFactors=FALSE within the read.csv() command, as shown in the following code:

```
# import csv file that contains both numeric and character variable
# firstly using default and then using stringsAsFActors=FALSE

iris_a <- read.csv("iris.csv")
str(iris_a)
'data.frame': 150 obs. of 5 variables:
$ Sepal.Length: num 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
$ Sepal.Width: num 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
$ Petal.Length: num 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
$ Petal.Width: num 0.2 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
$ Species : Factor w/ 3 levels "setosa", "versicolor",..:
1 1 1 1 1 1 1 1 1 1 ...</pre>
```

In the following example, we will see the difference if we specify the stringsAsFactors = FALSE argument:

```
# Now using stringsAsFactors=FALSE
iris_b <- read.csv("iris.csv",stringsAsFactors=F)
'data.frame': 150 obs. of 5 variables:
$ Sepal.Length: num 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
$ Sepal.Width: num 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
$ Petal.Length: num 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
$ Petal.Width: num 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
$ Species : chr "setosa" "setosa" "setosa" "setosa" ...</pre>
```

We see that in the first data frame, the class of the Species variable is factor, whereas in the second data frame the class of the same variable is character. So we have to be careful while importing the .csv file with mixed variables.

Sometimes, it could happen that the file extension is \*.csv but the data is not comma separated; rather, the data supplier has used a semicolon (;) as a separator, or any other symbol. In that case, we can still use the read.csv() command, but this time we have to specify the separator. Let's look at the example with a semicolon-separated .csv file of the same iris data:

```
iris_semicolon <-
   read.csv("iris_semicolon.csv",stringsAsFactors=FALSE,sep=";")
str(iris_semicolon)
'data.frame': 150 obs. of 5 variables:
$ Sepal.Length: num 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
$ Sepal.Width: num 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
$ Petal.Length: num 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
$ Petal.Width: num 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...
$ Species : chr "setosa" "setosa" "setosa" "setosa" ...</pre>
```

Similarly, if the values are tab separated, we can use read.csv() with sep= "\t". Alternatively, we can use read.table(). The following is an example:

```
anscombe_tab <- read.csv("anscombe.txt",sep="\t")
anscombe_tab_2 <- read.table("anscombe.txt",header=TRUE)</pre>
```

Notice that here when we used read.table(), we have to specify whether the variable name is present or not using header=TRUE.

If the dataset is stored in the \*.xls or \*.xlsx format, we have to use certain R packages to import those files; one of the packages is xlsx, which is designed to read files formatted as \*.xlsx. The following is an example to import the xlsxanscombe.xlsx file:

```
# Calling xlsx library
library(xlsx)
# importing xlsxanscombe.xlsx
anscombe xlsx <- read.xlsx2("xlsxanscombe.xlsx",sheetIndex=1)</pre>
```

In R, single or multiple data frames or other objects can be stored in the \*.RData format. This file format is convenient to store more than one dataset into a single file. To acquire a dataset for any other type of objects from the \*.RData file, we can use the load() function. The following is an example to load multiple datasets and a vector of R objects from a single \*.RData file:

```
# loading robjects.RData file
load("robjects.RData")
```

```
# to see whether the objects are imported correctly
objects()
"character.obj" "diab.dat" "logical.obj" "num.obj" "var1"
   "var2" "var3" "var4"
```

Note that the objects() command is used to look at all of the objects in the current R session. Now to see the mode and class of each object, we can easily use the mode() and class() function.

To import a Stata file into R, we need to call the foreign library and then use the read.dta() function. Similarly, if we want to import an SPSS data file, the corresponding command will be read.spss(); the output will always be a data frame. Here is an example of importing a Stata file:

```
library(foreign)
iris stata <- read.dta("iris stata.dta")</pre>
```

In this section, we saw that a dataset can be stored in different formats and R has some user friendly functionality to deal with each of them. The noticeable feature of this section is some of the arguments within the read.csv() function, such as skip, stringsAsFactors, and sep. To import any data correctly, we have to use these arguments carefully.

#### **Factor manipulation**

A variable that takes only a limited number of distinct values is usually known as a **categorical variable**, and in R, this is known as a **factor**. During data analysis, sometimes the factor variable plays an important role, particularly in studying the relationship between two categorical variables. In this section, we will see some important aspects of factor manipulation. When a factor variable is first created, it stores all its levels along with the factor. But if we take any subset of that factor variable, it inherits all its levels from the original factor levels. This feature sometimes creates confusion in understanding the results. Let us now see an example of this feature.

We will firstly create a factor variable from the datamanipulation character string with the English alphabet in lowercase as levels. Each letter of this string represents a value of that factor variable. Then, we will display the data with the table() function, where we will see lots of zero frequency corresponding to the letters that did not appear in the factor variable, as shown in the following code. We then drop those levels that are not part of the original factor variable and will display the data again.

```
# creating an R object whose value is "datamanipulation"
char.obj <- "datamanipulation"</pre>
```

Notice that there are only a few nonzero values in the table because the original factor variable does not have the entire alphabet as its value. Now we will drop the levels that do not appear in the original factor variable. To do so, we will create another factor variable from the original factor variable, as shown in the following code:

```
# re-creating factor variable from existing factor variable
factor.obj1 <- factor(factor.obj)

# Displaying levels of the new factor variable
levels(factor.obj1)
  [1] "a" "d" "i" "l" "m" "n" "o" "p" "t" "u"

# displaying data using table() function
table(factor.obj1)
factor.obj1
a d i l m n o p t u
4 1 2 1 1 2 1 1 2 1</pre>
```

The important feature to notice here is that we can drop unused factor levels by recreating factor variables from the original factor variable. This is most useful when we use a subset of a factor variable.

#### **Factors from numeric variables**

Numeric variables are convenient during statistical analysis, but sometimes we need to create categorical (factor) variables from numeric variables. We can create a limited number of categories from a numeric variable using a series of conditional statements, but this is not an efficient way to perform this operation. In R, cut is a generic command to create factor variables from numeric variables. In the following example, we will see how we can create factors from a numeric variable using a series of conditional statements. We will also use the cut command to perform the same task.

```
# creating a numeric variable by taking 100 random numbers
# from normal distribution
set.seed(1234) # setting seed to reproduce the example
numvar <- rnorm(100)</pre>
# creating factor variable with 5 distinct category
num2factor <- cut(numvar,breaks=5)</pre>
class(num2factor)
[1] "factor"
levels(num2factor)
[1] "(-2.35,-1.37]" "(-1.37,-0.389]" "(-0.389,0.592]"
"(0.592,1.57]" "(1.57,2.55]"
table(num2factor)
num2factor
(-2.35, -1.37] (-1.37, -0.389] (-0.389, 0.592] (0.592, 1.57] (1.57, 2.55]
                   43
                                                    13
                                                                   8
```

By default, the levels are produced using the actual range of values. Sometimes, the range of values are given a specific name for convenience. For example, the five categories of the preceding factor might be called the lowest group, lower-middle group, middle group, upper-middle group, and highest group, as shown in the following code:

```
2 lower middle group
                        43
3
        middle group
                        29
4
        upper middle
                        13
       highest group
                         8
# creating factor variable using conditional statement
num2factor <- factor(ifelse(numvar<=-1.37,1,</pre>
  ifelse(numvar<=-0.389,2,ifelse(numvar<=0.592,3,ifelse
    (numvar<=1.57,4,5)))),labels=c("(-2.35,-1.37]",
      "(-1.37,-0.389]", "(-0.389,0.592]",
        "(0.592,1.57]", "(1.57,2.55]"))
# displaying data using table function
table(num2factor)
num2factor
 (-2.35, -1.37] (-1.37, -0.389] (-0.389, 0.592] (0.592, 1.57] (1.57, 2.55]
```

Once we have converted the numeric variable to the factor variable and discarded the numeric variable, we cannot go back to the original numeric variable if needed. Therefore, we should be careful when converting the numeric variable to the factor variable.

#### **Date processing**

R can handle date variables in several ways. There are built-in R functions available to deal with date variables, and there are also some useful contributed packages available. The built-in R function as.Date() can handle only dates but not time, whereas the chron package, contributed by James and Hornik in 2008, can handle both date and time. However, it cannot deal with time zones. Using the POSIXct and POSIXlt class objects, we can deal with the inclusion of time zones. But there is another R package, lubridate, contributed by Grolemund and Wickham in 2011, that gives much more user friendly functionality to deal with date and time with time zone support. In this section, we will see how we can easily deal with date and time using the lubridate package and compare it with built-in R functions.

Like other statistical software, R also has a base date, and using that base date, R internally stores date objects. In R, dates are stored as the number of days elapsed since January 1, 1970. So if we convert any date object to its internal number, it will show the number of days. We can reformat the number into a date using the date class. The following are some examples:

```
# creating date object using built in as.Date() function
as.Date("1970-01-01")
```

```
[1] "1970-01-01"
# looking at the internal value of date object
as.numeric(as.Date("1970-01-01"))
[1] 0
# Second January 1970 is showing number of elapsed day is 1.
as.Date("1970-01-02")
[1] "1970-01-02"
as.numeric(as.Date("1970-01-02"))
[1] 1
```

Using the as.Date() function, we can easily create the date object; the typical format of the date object in this function is year, month, and then day. But we can also create a date object with other formats by specifying the format within the as.Date() function, as shown in the following example:

```
# creating date object specifying format of date
as.Date("Jan-01-1970",format="%b-%d-%Y")
[1] "1970-01-01"
```

Note that when specifying the format of the date, we have to give the format that is aligned with the input string. For the complete list of code that is used to specify date formats, users are directed to the help documentation of the strptime function. Users can access the complete list by just typing in help(strptime) in the R console.

The lubridate package provides intuitive functionality to deal with the date object in R. The following are some of the examples to create the date object using the lubridate package:

```
# loading lubridate package
library(lubridate)
# creating date object using mdy() function
mdy("Jan-01-1970")
"1970-01-01 UTC"
```

Note that the default time zone in the mdy, dmy, or ymd function is **Coordinated Universal Time** (UTC). One of the most interesting and important features of the lubridate package is that it can deal with heterogeneous formats. Heterogeneous formats means users can store date information in various ways; for example, second chapter due on 2013, august, 24, first chapter submitted on 2013, 08, 18, or 2013 aug 23. From this heterogeneous date, we can extract the valid date object that can be processed further within R using the lubridate package, as shown in the following code:

```
# creating heterogeneous date object
hetero_date <- c("second chapter due on 2013, august, 24",
"first chapter submitted on 2013, 08, 18", "2013 aug 23")</pre>
```

```
# parsing the character date object and convert to valid date
ymd(hetero_date)
[1] "2013-08-24 UTC" "2013-08-18 UTC" "2013-08-23 UTC"
```

Although the lubridate package can handle heterogeneous dates, the sequence of year, month, and day should be similar across all values within the same object, otherwise during date extraction there will be a missing value that will be generated along with a warning message. For example, if we alter the last date to 23 aug 2013, this will not get converted into a valid date, as shown in the following code:

```
hetero_date <- c("second chapter due on 2013, august, 24", "first chapter submitted on 2013, 08, 18", "23 aug 2013") ymd(hetero_date)
[1] "2013-08-24 UTC" "2013-08-18 UTC" NA
Warning message:
1 failed to parse.
```

During the date manipulation, sometimes we need to change the month only within an existing R date object. The following is an example of doing this using the default R function and also using the lubridate package:

```
# Creating date object using based R functionality
date <- as.POSIXct("23-07-2013",format = "%d-%m-%Y", tz = "UTC")
date
[1] "2013-07-23 UTC"
# extracting month from the date object
as.numeric(format(date, "%m"))
[1] 7
# manipulating month by replacing month 7 to 8
date <- as.POSIXct(format(date, "%Y-8-%d"), tz = "UTC")</pre>
date
[1] "2013-08-23 UTC"
# The same operation is done using lubridate package
date <- dmy("23-07-2013")
date
[1] "2013-07-23 UTC"
month (date)
[1] 7
month(date) <- 8
date
[1] "2013-08-23 UTC"
```

In a dataset, the variable might have both date and time information and we need to round them to the nearest day or month. The following example shows the date-rounding functionality; this example also displays how to convert the time zone:

```
# accessing system date and time
current time <- now()
current time
[1] "2013-08-23 23:43:01 BDT"
# changing time zone to "GMT"
current time gmt <- with tz(current time, "GMT")
current time gmt
[1] "2013-08-23 17:43:01 GMT"
# rounding the date to nearest day
round_date(current_time_gmt,"day")
[1] "2013-08-24 GMT"
# rounding the date to nearest month
round date(current time gmt, "month")
[1] "2013-09-01 GMT"
# rounding date to nearest year
round date(current time gmt, "year")
[1] "2014-01-01 GMT"
```

In this section, we saw that dealing with dates using the lubridate package is really user friendly and intuitive.

#### **Character manipulation**

In any statistical software, all the data is expected to be either numeric or at least a factor, but sometimes we have to work with character data. In the area of text mining, character or string manipulation is the most important. R has complete functionality to manipulate character (string) data for further analysis. Besides default R functionality, there is one contributed package to deal with character data, which is more user friendly and intuitive compared to the base R counterpart. Wickham developed the stringr package in 2010 to manipulate character data with some user friendly functions. In this section, we will introduce different functions and their counterparts in a table so that the readers are able to use the functions from the stringr package easily:

Base R functions	stringr functions
paste(): This function is used to concatenate a vector of characters with a default separator as a space.	<pre>str_c(): This has a functionality similar to paste(), but it uses empty as the default separator. It also silently removes zero-length arguments.</pre>
nchar(): This returns the number of characters in a character string.	<pre>str_length(): This is the same as nchar(), but it preserves NA. For example:</pre>
For NA, it returns 2, which is not expected. For example:	str_length(c("x","y",NA)) [1] 1 1 NA
nchar(c("x","y",NA)) [1] 1 1 2	
substr(): This extracts or replaces substrings in a character vector.	str_sub(): This is the equivalent of substr(), but it returns a zero-length vector if any of its inputs are of zero length. It also accepts negative positions, which are calculated from the left of the last character. The end position defaults to -1, which corresponds to the last character.
Unavailable	<pre>str_dup(): This is used to duplicate the characters within a string.</pre>
Unavailable	<pre>str_trim(): This is used to remove the leading and trailing whitespaces.</pre>
Unavailable 	<pre>str_pad(): This is used to pad a string with extra whitespaces on the left, right, or both sides.</pre>

Other than the functions listed in the preceding table, there are some other user friendly functions for pattern matching. Those functions are str\_detect, str\_locate, str\_extract, str\_match, str\_replace, and so on. To get more details about these functions, readers should refer to the stringr: modern, consistent string processing paper, by Wickham, which can be found at http://journal.r-project.org/archive/2010-2/RJournal\_2010-2\_Wickham.pdf.

#### Subscripting and subsetting

Subscripting and subsetting a dataset is an integral part of data manipulation. If we need to extract a smaller part of any R object (vector, data frame, matrix, or list) that contains more than one element, we need to use subscripts. Subscripting is an approach to access individual elements of an R object, for example, accessing particular element of a vector. Usually, numeric integers are used for subscripting, but logical vectors can also be used for the same purposes. In R, the subscript starts from 1, and if we specify any negative subscript, it omits that position from the source object.

The following is an example of an R vector with 10 elements and the effect of positive and negative subscripting:

```
# creating a 10 element vector
num10 <- c(3,2,5,3,9,6,7,9,2,3)
# accessing fifth element
num10[5]
[1] 9
# checking whether there is any value of num10 object greater
# than 6
num10>6
[1] FALSE FALSE FALSE FALSE TRUE FALSE TRUE TRUE FALSE FALSE
# keeping only values greater than 6
num10[num10>6]
[1] 9 7 9
# use of negative subscript removes first element "3"
num10[-1]
[1] 2 5 3 9 6 7 9 2 3
```

Note that the subscripted indexes are written within square brackets. For one-dimensional vectors, we use a single index to access elements, but for two-dimensional objects such as data frames or matrices, we have to use two-dimensional subscripts. In that case, we have to use double square brackets for indexing. The first index is for representing rows and the second is for representing columns; for example:

```
# creating a data frame with 2 variables
data_2variable <- data.frame(x1=c(2,3,4,5,6),x2=c(5,6,7,8,1))
# accessing only first row
data_2variable[1,]
    x1 x2
1    2    5
# accessing only first column
data_2variable[,1]
[1] 2 3 4 5 6
# accessing first row and first column
data_2variable[1,1]
[1] 2</pre>
```

Similar indexing is used for matrices. For the list object, the indexing is different than that of data frames or matrices. To get access to a list object, we have to use [[]] for indexing; for example, [[1]] to get the first element of a list. If the list is nested within another list, we need to use a series of double square brackets within double square brackets. The following example creates a list object and accesses its elements:

```
list_obj<- list(dat=data_2variable,vec.obj=c(1,2,3))
list_obj
$dat
    x1 x2
1    2    5
2    3    6
3    4    7
4    5    8
5    6    1

$vec.obj
[1] 1 2 3
# accessing second element of the list_obj objects
list_obj[[2]]
[1] 1 2 3</pre>
```

Now, if we want to get access to the individual elements of <code>list\_obj[[2]]</code>, we have to use the following command:

```
list_obj[[2]][1]
[1] 1
```

If the list object is named, we can get access to the elements of that list using the name as follows:

```
# accessing dataset from the list object
list_obj$dat
x1 x2
1  2  5
2  3  6
3  4  7
4  5  8
5  6  1
```

Subsetting is just storing subscripted objects. Once we extract any subscripted R object and store it in another variable, the newly created object is the subset of the original variable.

#### **Summary**

In this chapter, we have covered some of the special features that we need to consider during data acquisition. We also discussed the important aspect of factor manipulation, especially when subsetting a factor variable and how to remove unused factor levels. The processing of date variables was covered with the use of the lubridate package, with its user friendly and intuitive functions, and also string processing has been highlighted. The chapter ended with an explanation of the concepts of subscripting and subsetting. For more details on date processing and string manipulation, readers should refer to the string: modern, consistent string processing paper, by Wickham, which can be found at http://journal.r-project.org/archive/2010-2/RJournal\_2010-2\_Wickham.pdf., and the Dates and Times Made Easy with lubridate journal, by Grolemund and Wickham, which can be found at http://www.jstatsoft.org/v40/i03/paper.

In the next chapter, we will discuss data manipulation with the plyr package, where we will focus on the split-apply-combine strategy, a state-of-the-art approach in the group-wise data manipulation using R.

## Data Manipulation Using plyr

In any data analysis task, a majority of the time is dedicated to data cleaning and preprocessing. It is considered that sometimes about 80 percent of the effort is devoted to data cleaning before conducting actual analysis (*Exploratory Data Mining and Data Cleaning, Dasu T.* and *Johnson T.*). Data manipulation is an integral part of data cleaning and analysis. For large sets of data, it is always preferable to perform the operation within subgroups of a dataset to speed up the process. In R, this type of data manipulation can be done with base functionality, but for large-scale data it requires a considerable amount of coding and eventually takes more processing time. In the case of large-scale data, we can split the dataset, perform the manipulation or analysis, and then combine it into a single output again. This type of split using default R is not efficient, and to overcome this limitation, Wickham, in 2011, developed an R package called **plyr**, in which he efficiently implemented the **split-apply-combine** strategy. This chapter starts with the concept of split-apply-combine and is followed by the different functions and utilities of the plyr package.

#### The split-apply-combine strategy

Often, we require similar types of operations in different subgroups of a dataset, such as group-wise summarization, standardization, and statistical modeling. This type of task requires us to break up a big problem into manageable pieces, perform operations on each piece separately, and finally combine the output of each piece into a single piece of output (the paper *The Split-Apply-Combine Strategy for Data Analysis*, by Wickham, which can be found at http://www.jstatsoft.org/v40/i01/paper). To understand the split-apply-combine strategy intuitively, we could compare this with the **map-reduce** strategy for processing large amounts of data, recently popularized by Google. In the map-reduce strategy, the map step corresponds to split and apply and the reduce step consists of combining. The map-reduce approach is primarily designed to deal with a highly parallel environment where the work has been done by several hundreds or thousands of computers independently.

The split-apply-combine strategy creates an opportunity to see the similarities of problems across subgroups that were previously unconnected. This strategy can be used in many existing tools, such as the GROUP BY operation in SAS, PivotTable in MS Excel, and the SQL GROUP BY operator.

To explain the split-apply-combine strategy, we will use Fisher's iris data. This dataset contains the measurements in centimeters of these variables: sepal length and width, and petal length and width, for 50 flowers from each of the three species of iris. The species are *Iris setosa*, *Iris versicolor*, and *Iris virginica*. We want to calculate the mean of each variable and for each species separately. This can be done in different ways using a loop or without using one.

#### Split-apply-combine without a loop

In this section, we will see an example of the split-apply-combine strategy without using a loop. The steps are as follows:

- 1. Split the iris dataset into three parts.
- 2. Remove the species name variable from the data.
- 3. Calculate the mean of each variable for the three different parts separately.
- 4. Combine the output into a single data frame.

The code for this is as follows:

```
# notice that during split step a negative 5 is used within the
# code, this negative 5 has been used to discard fifth column of the
# iris data that contains "species" information and we do not need
# that column to calculate mean.

iris.set <- iris[iris$Species=="setosa",-5]
iris.versi <- iris[iris$Species=="versicolor",-5]
iris.virg <- iris[iris$Species=="virginica",-5]
# calculating mean for each piece (The apply step)
mean.set <- colMeans(iris.set)
mean.versi <- colMeans(iris.versi)
mean.virg <- colMeans(iris.virg)

# combining the output (The combine step)
mean.iris <- rbind(mean.set,mean.versi,mean.virg)

# giving row names so that the output could be easily understood
rownames(mean.iris) <- c("setosa", "versicolor", "virginica")</pre>
```

#### Split-apply-combine with a loop

The following example will calculate the same statistics as in the previous section, but this time we will perform this task using a loop. The steps are similar but the code is different. In each iteration, we will split the data for each species and calculate the mean for each variable and then combine the output into a single data frame, as shown in the following code:

```
# split-apply-combine using loop
# each iteration represents split
# mean calculation within each iteration represents apply step
# rbind command in each iteration represents combine step

mean.iris.loop <- NULL
for(species in unique(iris$Species))
{
   iris_sub <- iris[iris$Species==species,]
   column_means <- colMeans(iris_sub[,-5])
   mean.iris.loop <- rbind(mean.iris.loop,column_means)
}

# giving row names so that the output could be easily understood rownames(mean.iris.loop) <- unique(iris$Species)</pre>
```

An important fact to note in the split-apply-combine strategy is that each piece should be independent of the other. If the calculation in one piece is somehow dependent on the other, the split-apply-combine strategy will not work. This strategy is not applicable in running an average type of operation, where a current average is dependent on the previous one. This strategy is only applicable when the big problem can be broken up into smaller manageable pieces and we can perform the desired operation on each piece independently. For running average calculations, the split-apply-combine strategy is not suitable; we can use a loop instead. But if processing speed is a concern, we can write the code in some lower-level language such as C or ForTran.

#### **Utilities of plyr**

The plyr package is a set of tools for a common set of problems. We want to split the big problem into smaller pieces, apply functions, and then combine all the outputs back together. The example we presented using the iris data is one where we applied this strategy. Though it is already possible to perform split-apply-combine operations with base R, such as the split and apply family of functions, plyr makes things much easier and intuitive with its consistent naming convention, various types of input-output processing, and built-in error recovery and informative messages. In general, plyr provides a replacement for the for loop. We do not need to replace the for loop just because it is slow, but we need to replace it to avoid extra, unimportant bookkeeping code. The following examples will clarify the need to replace a for loop with its plyr counterpart.

The mean calculation of each variable within each species in the iris dataset using the for loop in base R can be coded as follows:

```
mean.iris.loop <- NULL
for(species in unique(iris$Species))
  iris_sub <- iris[iris$Species==species,]</pre>
  column means <- colMeans(iris sub[,-5])</pre>
  mean.iris.loop <- rbind(mean.iris.loop,column means)</pre>
rownames(mean.iris.loop) <- unique(iris$Species)</pre>
mean.iris.loop
            Sepal.Length Sepal.Width Petal.Length Petal.Width
                                                            0.246
                   5.006
                                 3.428
                                               1.462
setosa
versicolor
                   5.936
                                 2.770
                                               4.260
                                                            1.326
virginica
                   6.588
                                 2.974
                                               5.552
                                                            2.026
```

The same mean calculation, but this time using the plyr package, can be coded as follows:

```
library (plyr)
ddply(iris,~Species,function(x) colMeans(x[,-which(colnames(x)=="Species")]))
mean.iris.loop
```

	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
setosa	5.006	3.428	1.462	0.246
versicolor	5.936	2.770	4.260	1.326
virginica	6.588	2.974	5.552	2.026

Note that we can easily perform the same calculation with very little code using the plyr package compared to the for loop in base R.

#### Intuitive function names

To perform any kind of data processing, we need to know the type of input we have to provide and the expected format of output. In most R functions, it is difficult to understand from function names what type of input they accept and what the expected outputs are. Function names in plyr packages are much more intuitive and instructive about their input and output type. Each function is named according to the type of input it accepts and the type of output it produces. The first letter of the function name specifies the input and the second letter specifies the output type; a represents array, d represents data frame, 1 represents list, and \_ (underscore) represents the output discarded. For example, the function name adply() takes input as array and produces output as a data frame. The following table gives us a complete idea about function-naming conventions used in the plyr package:

Input	Output			
	Array	Data frame	List	Discarded
Array	aaply	adply	alply	a_ply
Data frame	daply	ddply	dlply	d_ply
List	laply	ldply	llply	l_ply

We can see that there are three types of input and four types of output. Users can easily get an idea of the types of input and output from the function names. Another interesting feature is that we do not need to learn all the 12 functions; rather, it is sufficient to learn the three types of input and four types of output.

Other than the function names in the table, there are some special cases of operating on arrays that correspond to the mapply() function in base R. In base R, mapply() can take multiple inputs as separate arguments, whereas a\*ply() takes only a single array argument. However, the separate argument in mapply() should be of the same length. The mapply() functions that are equivalent to plyr are maply(), mdply(), mlply(), and  $m_ply()$ .

Note that whenever a function name is written using a star symbol, such as a\*ply(), it indicates that the common input is an array. The output can be in any format: array, data frame, or list. Optionally, the output can be discarded.

To explain the intuitive nature of the input and output, we will now provide an example using the iris data that we used in an earlier example. This time, we will use the iris3 dataset; this is the same data, but stored in a three-dimensional array format. We will calculate the mean of each variable for each species as shown in the following code:

```
# class of iris3 dataset is array
class(iris3)
[1] "array"

# dimension of iris3 dataset
dim(iris3)
[1] 50 4 3
```

The following code snippet calculates the column mean for each species with the input as an array and the output as a data frame:

```
# Calculate column mean for each species and output will be data
frame
iris_mean <- adply(iris3,3,colMeans)</pre>
class(iris mean)
[1] "data.frame"
iris mean
         X1 Sepal L. Sepal W. Petal L. Petal W.
     Setosa
               5.006
                                1.462 0.246
                       3.428
             5.936
2 Versicolor
                       2.770
                                4.260
                                        1.326
3 Virginica
             6.588
                       2.974
                                5.552
                                        2.0266
```

The following code snippet calculates the column mean for each species with the input as an array and the output as an array too:

```
# again we will calculate the mean but this time output will be an
# array
iris_mean <- aaply(iris3,3,colMeans)
class(iris_mean)
[1] "matrix"</pre>
```

```
iris mean
```

```
Х1
             Sepal L. Sepal W. Petal L. Petal W.
                5.006
                         3.428
                                  1.462
 Setosa
                                           0.246
 Versicolor
                5.936
                         2.770
                                  4.260
                                           1.326
                6.588
                         2.974
                                  5.552
                                          2.026
 Virginica
# note that here the class is showing "matrix", since the output is a
# two dimensional array which represents matrix
# Now calculate mean again with output as list
iris mean <- alply(iris3,3,colMeans)</pre>
class(iris mean)
[1] "list"
iris mean
$'1'
Sepal L. Sepal W. Petal L. Petal W.
   5.006 3.428
                     1.462
                              0.246
$'2'
Sepal L. Sepal W. Petal L. Petal W.
   5.936 2.770
                   4.260
                              1.326
$131
Sepal L. Sepal W. Petal L. Petal W.
  6.588
            2.974
                   5.552
                              2.026
attr(,"split type")
[1] "array"
attr(, "split labels")
         X1
1
      Setosa
2 Versicolor
3 Virginica
```

#### Input and arguments

The functions in the plyr package accept various input objects: data frames, arrays, and lists. Each input object has its own rule to split the process. In this section, we will discuss input and arguments. The rules of splitting can be described shortly as follows:

- Arrays are sliced by dimension into lower dimensional pieces, and the
  corresponding common function is a\*ply(), where the array is the common
  input and the output can be one among an array, data frame, or list.
- Data frames are sliced and subset by a combination of variables from that dataset. The corresponding common function is d\*ply(), where the data frame is the common input and the output can be one among an array, data frame, or list.
- The elements of a list are processed separately, and the common function
  is 1\*ply(), where the common input is a list and the output can be an array,
  data frame, or list.

Depending on the input type, there are two or three main arguments for the common functions: a\*ply(), d\*ply(), and l\*ply(). The following are the main arguments for these common functions:

```
a*ply(.data, .margins, .fun, ..., .progress = "none")
d*ply(.data, .variables, .fun, ..., .progress = "none")
l*ply(.data, .fun, ..., .progress = "none")
```

The first argument, .data, is the input dataset that needs to be processed by splitting up, and the output will be combined from each split. The .margins or .variables argument specifies how the data should be split up into smaller pieces. The .fun argument specifies the processing task; this can be any function that is applicable to each split of the input. If we omit the .fun argument, the input data is just converted to the output structure specified by the function. If we want to monitor the progress of the processing task, the .progress argument should be specified. It will not show the progress status by default.

In the following example, we will see what will happen if we do not specify the <code>.fun</code> argument in any function of the <code>plyr</code> package. If we give the input as an array and want the output as a data frame, but we haven't given a <code>.fun</code> argument, the <code>adply()</code> function will just convert the array object into a data frame. Here is an example:

```
# converting 3 dimensional array to a 2 dimensional data frame
iris_dat <- adply(iris3, .margins=3)
class(iris_dat)</pre>
```

The .margins argument works in a similar manner to the apply function in base R. It does the following:

- Slices up a row by specifying .margins = 1
- Slices up a column by .margins = 2
- Slices up the individual cells by .margins = c(1,2)

The .margins argument works correspondingly for higher dimensions, with a combinatorial explosion in the number of possible ways to slice up the array.

#### Comparing default R and plyr

In this section, we will compare code side by side to solve the same problem using both default R and plyr. Reusing the iris3 data, we are now interested in producing five number summary statistics for each variable group by species. The five numbers will be minimum, mean, median, maximum, and standard deviation. The output will be a list of data frames.

To calculate the summaries for the five numbers, follow these steps:

- 1. Define a function that will calculate five number summary statistics for a given vector.
- 2. Produce the output of this function in a data frame object.
- 3. Apply this function in the iris3 dataset using a for loop.
- 4. Apply the same function using the apply() function of the plyr package.

An example that explains the calculation of the five number summary statistics is as follows:

```
# Function to calculate five number summary
fivenum.summary <- function(x)
{
   results <-data.frame(min=apply(x,2,min),
   mean=apply(x,2,mean),
   median=apply(x,2,median),
   max=apply(x,2,max),
   sd=apply(x,2,sd))
   return(results)
}</pre>
```

The following is how we calculate the summaries for the five numbers using a for loop with default R:

```
# initialize the output list object
all stats <- list()</pre>
# the for loop will run for each species
for(i in 1:dim(iris3)[3])
  sub data <- iris3[,,i]</pre>
  all stat species <- fivenum.summary(sub data)
  all stats[[i]] <- all stat species
# class of the output object
class(all stats)
[1] "list"
all stats
[[1]]
          min mean median max
Sepal L. 4.3 5.006 5.0 5.8 0.3524897

      Sepal W. 2.3 3.428
      3.4 4.4 0.3790644

      Petal L. 1.0 1.462
      1.5 1.9 0.1736640

Petal W. 0.1 0.246
                       0.2 0.6 0.1053856
[[2]]
          min mean median max
Sepal L. 4.9 5.936 5.90 7.0 0.5161711
Sepal W. 2.0 2.770 2.80 3.4 0.3137983
Petal L. 3.0 4.260 4.35 5.1 0.4699110
Petal W. 1.0 1.326 1.30 1.8 0.1977527
```

```
[[3]]

min mean median max sd

Sepal L. 4.9 6.588 6.50 7.9 0.6358796

Sepal W. 2.2 2.974 3.00 3.8 0.3224966

Petal L. 4.5 5.552 5.55 6.9 0.5518947

Petal W. 1.4 2.026 2.00 2.5 0.2746501
```

Let's calculate the same statistics, but this time using the alply() function from the plyr package:

```
all stats <- alply(iris3,3,fivenum.summary)</pre>
class(all stats)
[1] "list"
all_stats
$'1'
        min mean median max
Sepal L. 4.3 5.006
                    5.0 5.8 0.3524897
Sepal W. 2.3 3.428
                     3.4 4.4 0.3790644
Petal L. 1.0 1.462 1.5 1.9 0.1736640
Petal W. 0.1 0.246
                    0.2 0.6 0.1053856
$'2'
         min mean median max
                                    sd
Sepal L. 4.9 5.936 5.90 7.0 0.5161711
Sepal W. 2.0 2.770 2.80 3.4 0.3137983
Petal L. 3.0 4.260 4.35 5.1 0.4699110
Petal W. 1.0 1.326 1.30 1.8 0.1977527
$131
        min mean median max
                                    sd
Sepal L. 4.9 6.588 6.50 7.9 0.6358796
Sepal W. 2.2 2.974 3.00 3.8 0.3224966
Petal L. 4.5 5.552 5.55 6.9 0.5518947
Petal W. 1.4 2.026 2.00 2.5 0.2746501
attr(,"split_type")
[1] "array"
attr(, "split labels")
         Х1
1
     Setosa
2 Versicolor
3 Virginica
```

#### **Multiargument functions**

Sometimes, we have to deal with functions that take multiple arguments, and the values of each argument can come from a data frame, a list, or an array. The plyr package has intuitive and user friendly functions to work with multiargument functions. In this section, we will see an example of generating random numbers from normal distribution with various combinations of mean and standard deviation. The values of mean and standard deviation are stored in a data frame. Now, we will generate random numbers using default R functions, such as the for loop, and also using the mlply() function from the plyr package. The parameter combinations are given in the following table:

n	Mean	Standard deviation	
25	0	1	
50	2	1.5	
100	3.5	2	
200	2.5	5	
500	0.1	2	

With these parameter combinations, we will generate normal random numbers using default R and plyr as shown in the following code:

```
# define parameter set
parameter.dat <- data.frame(n=c(25,50,100,200,400), mean=c(0,2,3.5,2.5)
,0.1), sd=c(1,1)
.5, 2, 5, 2))
# displaying parameter set
parameter.dat
   n mean sd
 25 0.0 1.0
1
  50 2.0 1.5
3 100 3.5 2.0
4 200 2.5 5.0
5 400 0.1 2.0
# random normal variate generate using base R
# set seed to make the example reproducible
set.seed(12345)
```

# initialize blank list object to store the generated variable

```
dat <- list()</pre>
for(i in 1:nrow(parameter.dat))
dat[[i]] <- rnorm(n=parameter.dat[i,1],</pre>
              mean=parameter.dat[i,2],sd=parameter.dat[i,3])
}
# estimating mean from the newly generated data
estmean <- lapply(dat, mean)</pre>
estmean
[[1]]
[1] -0.001177287
[[2]]
[1] 2.417842
[[3]]
[1] 3.667193
[[4]]
[1] 2.999662
[[5]]
[1] 0.1765926
# Performing same task as above but this time use plyr package
dat plyr <- mlply(parameter.dat,rnorm)</pre>
estmean_plyr <- llply(dat_plyr,mean)</pre>
estmean_plyr
$'1'
[1] 0.4252469
$121
[1] 2.037528
$131
[1] 3.070231
$'4'
[1] 2.144276
$'5'
[1] 0.05399488
```

#### Summary

In this chapter, we discussed the importance of the split-apply-combine strategy. We understood what the split-apply-combine strategy is and why it is important in data manipulations. The split-apply-combine strategy can be implemented using base R, but requires a large amount of code and is not memory or time efficient. To overcome this limitation, we discussed the plyr package, in which group-wise data manipulation can be implemented efficiently. The functions within plyr are intuitive and instructive in terms of input and output type. A large variety of data processing can be done using only a few functions with common input and various outputs. For further reading, an interested user can look up (at the paper *The Split-Apply-Combine Strategy for Data Analysis*, by Wickham, which can be found at http://www.jstatsoft.org/v40/i01/paper). In the upcoming chapter, we will learn about reshaping a dataset, which is another important aspect of group-wise data manipulation.

# 4

## **Reshaping Datasets**

Reshaping data is a common and tedious task in real-life data manipulation and analysis. A dataset might come with different levels of grouping and we need to implement some reorientation to perform certain types of analyses. Datasets layout could be long or wide. In long-layout, multiple rows represent a single subject's record, whereas in wide-layout, a single row represents a single subject's record. Statistical analysis sometimes requires wide data and sometimes long data, and in such cases, we need to be able to fluently and fluidly reshape the data to meet the requirements of statistical analysis. Data reshaping is just a rearrangement of the form of the data – it does not change the content of the dataset. In this chapter, we will show you different layouts of the same dataset and see how they can be transferred from one layout to another. This chapter mainly highlights the melt and cast paradigm of reshaping datasets, which is implemented in the reshape contributed package. Later on, this same package is reimplemented with a new name, reshape2, which is much more time and memory efficient (the Reshaping Data with the reshape Package paper, by Wickham, which can be found at http://www.jstatsoft.org/v21/i12/paper). In this chapter, we will discuss the layout of a dataset and how we can change the layout using the new paradigm of reshaping datasets with melt and cast. To run the example of this chapter, readers need to install both the reshape and reshape2 packages.

#### The typical layout of a dataset

A single dataset can be rearranged in many different ways, but before going into rearrangement, let's look back at how we usually perceive a dataset. Whenever we think about any dataset, we think of a two-dimensional arrangement where a row represents a subject's (a subject could be a person and is typically the respondent in a survey) information for all the variables in a dataset and a column represents the information for each characteristic for all subjects. This means rows indicate records and columns indicate variables, characteristics, or attributes. This is the typical layout of a dataset. In this arrangement, one or more variables might play a role as an identifier and others are measured characteristics. For the purpose of reshaping, we could group the variables into two groups: identifier variables and measured variables.

- Identifier variables: These help to identify the subject from whom we took
  information on different characteristics. Typically, identifier variables are
  qualitative in nature and take a limited number of unique values. In database
  terms, an identifier is termed as the primary key, and this can be a single
  variable or a composite of multiple variables.
- Measured variables: These are those characteristics whose information
  we took from a subject of interest. These can be qualitative, quantitative,
  or a mix of both.

#### Long layout

In this layout, the dataset is arranged in such a way that a single subject's information is stored in multiple rows. We need a composite identification variable to identify a unique row. This type of layout is usually seen in a longitudinal dataset. The following is an example of this type of dataset:

sid	exmterm	math	literature	language
1	1	50	40	70
1	2	65	45	80
2	1	75	55	75
2	2	69	59	78

Notice that in the dataset we have repeated sid, but if we consider both sid and exmterm, each row can be identified uniquely. This layout is known as the long layout. The following is the R code to produce this data frame:

```
# Example of typical two dimensional data
# A demo dataset "students" with typical layout. This data
# contains two students' exam score of "math", "literature"
# and "language" in different term exam.
students <- data.frame(sid=c(1,1,2,2),
exmterm=c(1,2,1,2),
math=c(50,65,75,69),
literature=c(40,45,55,59),
language=c(70,80,75,78))
students
  sid exmterm math literature language
            1
1
                50
                           40
                                    70
            2 65
   1
                           45
                                    80
            1
3
    2
                75
                           55
                                    75
    2
            2
                69
                           59
                                    78
```

#### Wide layout

In this layout, each row represents all the information of a single subject. Usually, only one identification variable is enough to identify a unique subject, but a composite identification variable can be used. The main difference between wide and long layout is that the wide layout contains all the measured information in different columns. The following is the wide layout of the same data that we initially stored in long layout:

sid	math.1	literature.1	language.1	math.2	literature.2	language.2
1	50	40	70	65	45	80
2	<i>7</i> 5	55	75	69	59	78

Notice that in this layout, each row contains all the information corresponding to a single value of sid. This layout is known as the wide form. In a later section, we will see how we can convert long to wide and vice versa using R.

#### The new layout of a dataset

In R, the layout of a dataset is known differently than the typical layout we discussed in previous section. This new layout of a dataset consists of only the identification variables and a value per variable. The identification variable identifies a subject along with which measured variable the value represents, which is the long layout in this paradigm. In this new paradigm, each row represents one observation of one variable. Interestingly, the typical long and wide layouts are both known as wide layout in this new paradigm. In the new paradigm, long data is also known as molten data and the process of producing molten data is known as melting from the wide layout. The difference between this new layout of the data and the typical layout is that it now contains only the ID variable and a new column, value, which represents the value of that observation. The following is an example of molten data that comes from the typical long layout:

sid	exmterm	variable	value
1	1	math	50
1	2	math	65
2	1	math	75
2	2	math	69
1	1	literature	40
1	2	literature	45
2	1	literature	55
2	2	literature	59
1	1	language	70
1	2	language	80
2	1	language	75
2	2	language	78

In this dataset, we see that each row contains all the information of one student, which is known as the wide data. The following is the R code to generate this molten data:

```
# Example of molten data
library(reshape)
molten_students <- melt.data.frame
  (students,id.vars=c("sid","exmterm"))</pre>
```

The melt.data.frame function converts the wide data to long (molten) form, and the new layout will contain only the identification variables along with two other columns named variable and value. In the new layout, each row contains the observation of a single variable, which is also known as the long form. The variable column represents the identification information along with what is being measured and the value column contains the measurement itself.

## Reshaping the dataset from the typical layout

In this section, we will see how we can convert a typical long layout to a typical wide layout and vice versa. To perform this conversion, we will use the built-in reshape() function, which takes several arguments, but we will use the following arguments:

- data: This argument specifies the dataset that we want to change the layout of.
- direction: This argument specifies whether the data is long or wide. Note that here long and wide indicates the typical layout.
- idvar: This argument specifies the identification variable. It could be a single variable or multiple variables.
- timevar: This argument specifies how many times the values of idvar repeats for each subject.

The following example converts the long layout to a wide layout of the students' data that was created earlier:

After reshaping the data, we see that the rows contain each student's exam record. Now we will change the layout from wide to long using the same function:

```
# Now again reshape to long format
long students <- reshape
(wide students,direction="long",idvar="id")
long students
sid exmterm math.1 literature.1 language.1
                                         70
  2
          1
                 75
                              55
                                         75
          2
  1
                 65
                              45
                                         80
           2.
                 69
                              59
                                         78
```

The limitation of this default reshape function is that it can only deal with the long and wide structures. In reality, data might contain multiple nested levels. To deal with complex data structures, the reshape function is not useful; we could use the reshape package instead.

# Reshaping the dataset with the reshape package

As we have seen, there are two different paradigms for defining the layout of a dataset. To change the layout of a dataset following the steps of a new paradigm, we need to use the reshape packages, where all the functions are implemented following the new layout. The main idea of the reshape package is melting a dataset and then casting it to a suitable layout. In the section *The new layout of a dataset*, we have mentioned melting a dataset and what it looks like. Just to recall, in molten data, each row represents a single observation of a single variable in the dataset. Also, it contains only the identifier variables and a value variable to represent what is being measured. In this section, we will discuss melting with more examples and casting with molten datasets.

#### Melting data

In R, melting is a generic operation and can be applied to various data types, including data frames, arrays, and matrices. Though melting can be applied to different R objects, the most common use is melting a data frame. To perform melting operations using the melt function, we need to know what the identification variables are and what the measured variables in the original input dataset are.

If we do not specify the identification variables and measured variables, by default any factor variables are assumed as the ID variables and any numeric variables are assumed as measured variables. To avoid this ambiguous operation, it would be good to specify it explicitly. If we specify only one type of variable, either identification or measured, the function assumes the remaining variable is of the other category. For example, if we specify only the ID variables, the remaining variables will be considered as measured variables and vice versa. The following example will clarify these points:

```
# original data
students
  sid exmterm math literature language
1
    1
             1
                 50
                             40
                                       70
2
    1
             2
                 65
                             45
                                       80
3
    2
             1
                 75
                             55
                                       75
             2
4
    2
                 69
                             59
                                       78
# Melting by specifying both id and measured variables
melt(students,id=c("sid","exmterm"),
   measured=c("math","literature","language"))
                variable value
 sid exmterm
     1
              1
                       math
1
                                50
2
     1
              2
                       math
                                65
3
     2
              1
                       math
                                75
4
                       math
                                69
5
     1
              1 literature
                               40
6
     1
              2 literature
                               45
7
     2
              1 literature
                                55
8
     2
              2 literature
                                59
9
     1
              1
                  language
                                70
10
     1
              2
                  language
                                80
                  language
11
     2
              1
                                75
              2
12
     2
                  language
                                78
# Melting by specifying only id variables
melt(students,id=c("sid","exmterm"))
```

	sid	exmterm	variable	value
1	1	1	math	50
2	1	2	math	65
3	2	1	math	75
4	2	2	math	69
5	1	1	literature	40
6	1	2	literature	45
7	2	1	literature	55
8	2	2	literature	59
9	1	1	language	70
10	1	2	language	80
11	2	1	language	75
12	2	2	language	78

In the melting process, the melt function does not assume the ID or measured variables; there could be any number of variables in any order, which gives flexibility to deal with the complex dataset. One important thing to note is that whenever we use the melt function, all the measured variables should be of the same type; that is, the measured variables should be either *numeric*, *factor*, *character*, or *date*.

#### Missing values in molten data

There could be two types of missing values in practice: one is **sampling zero** (that is, no response) and the other is **structural missing**. The sampling zeros are explicitly coded and represented in the dataset, but the structural missing depends on the structure of the dataset. The structural missing is implicit in the dataset represented by the absence of a certain combination of the ID variable. If we change the structure of a dataset from nested to crossed, the implicit missing no longer exists in the data, rather it explicitly appears in the new structure and care should be taken to deal with that data. The following simple example taken from the *Reshaping Data with the reshape Package* paper, by Wickham, which can be found at http://www.jstatsoft.org/v21/i12/paper, clearly explains implicit and explicit missing in two different data structures.

Consider a dataset with two ID variables, sex (male or female) and pregnant (yes or no). When the variables are nested, the missing value "pregnant male" is represented by its absence in the dataset, as shown in the following table. However, in a crossed view, we need to add the explicit missing value as there will now be a cell that must be filled with a value.

Sex	Pregnant	Value
Male	No	10
Female	No	14
Female	Yes	4

The cross view of this table can be represented as follows:

Sex	Pregnant	Not Pregnant
Male		10
Female	4	14

To deal with the implicit missing value, it is good to use na.rm=TRUE with the melt function to remove the structural missing. If we do not specify na.rm=TRUE during melting, we have to specify this during data analysis.

#### **Casting molten data**

Once we have molten data, we can rearrange it in any layout using the cast function from the reshape package. There are two main arguments required to cast molten data:

- data: This is the molten data that we want to reshape.
- formula: This is the casting formula to determine the layout of the output data; for example, which variable should go into columns and which should go into rows. If we do not specify a formula, the cast will return the classic data frame.

There are other argument options for performing certain types of operations, if required. The basic casting formula is col\_var\_1+col\_var\_2 ~ row\_var\_1+row\_var\_2, which describes the variables to appear in columns and rows. The following example shows how the cast function works:

# Melting students data

```
molten students <- melt(students,id.vars=c("sid","exmterm"))</pre>
molten students
  sid exmterm variable value
1
    1
           1
                  math
                          50
   1
          2
2
                   math
                          65
          1
2
3
   2
                         75
                  math
                 math
4
                         69
           1 literature
                         40
```

6	1	2	literature	45
7	2	1	literature	55
8	2	2	literature	59
9	1	1	language	70
10	1	2	language	80
11	2	1	language	75
12	2	2	language	78

Now use the cast function to return to the original data structure by specifying both row and column variables, as follows:

```
cast(molten_students,sid+exmterm~variable)
  sid exmterm math literature language
1
    1
            1
                50
                            40
2
    1
            2
                65
                            45
                                     80
3
            1
                75
                            55
                                     75
4
    2
            2
                69
                            59
                                     78
```

The following is the same operation, but specifying only row variables:

```
cast(molten students,...~variable)
  sid exmterm math literature language
1
                 50
                             40
2
    1
             2.
                 65
                             45
                                       80
3
                 75
                             55
            1
                                       75
4
    2
                 69
                             59
                                       78
```

We now rearrange the data in a way that sid is now a separate column for each student, as follows:

We again rearrange the data in a way that exmterm is now a separate column for each term, as follows:

```
cast (molten_students,...~exmterm)
  sid variable 1 2
1 1    math 50 65
2 1 literature 40 45
3 1 language 70 80
4 2    math 75 69
5 2 literature 55 59
6 2 language 75 78
```



Note that the column names of the last two examples are not valid column names because they contain numbers. It is a limitation of R. R cannot automatically label row or column names unambiguously, so we have to be careful about column names during analysis.

#### The reshape2 package

Though the reshape package has various functions to perform various tasks that cannot be done using built-in R functions, this package is slow. To make this more time and memory efficient, Wickham reimplemented this package and developed another package, reshape2. The reason behind the development of the new reshape2 package is to keep the functionality of the original reshape package so that users do not get confused. Some important new features of the reshape2 package are as follows:

- It is much better than the original reshape package in terms of memory and time efficiency
- It uses several functions instead of only the cast function
- The multidimensional marginal total can be calculated

#### **Summary**

This chapter introduced the theoretical framework for reshaping a dataset. The limitations of conventional approaches were pointed out and the new paradigm of data layout was highlighted. In the new paradigm, only two functions allow users to rearrange datasets into various layouts as required. This chapter also discussed structural missing and sampling zero, and how to deal with those missing during the melting process. For faster and large data rearrangement, readers were redirected to the reshape2 package. In the next chapter, we will discuss how R can be connected with databases and handle large-scale data.

# 5 R and Databases

We noticed earlier that a dataset can be stored in any format using different software as well as relational databases. Usually, large-scale datasets are stored in database software. In data mining and statistical learning, we need to process large-scale datasets. One of the major problems in R is memory usage. R is RAM intensive, and for that reason, the size of a dataset should be much smaller than its RAM. Also, one of the major drawbacks of R is its inability to deal with large datasets.

This chapter introduces how to deal with large datasets that are bigger than the computer's memory and dealing with a dataset by interacting with database software. In the first few sections, we describe how to interact with database software with **Open Database Connectivity (ODBC)** and import datasets. This chapter will present an example of memory issues and then describe ODBC using an example of MS Excel and MS Access, dealing with large datasets with specialized contributed R packages. This chapter ends with an introduction to data manipulation using SQL through the sqldf package.

The first are two examples demonstrating memory problems in R:

 The following example explains the memory limitation of a computer system. R stores everything in RAM, and a typical personal computer consists of limited RAM (depending on the computer's operating system, that is, 32-bit or 64-bit).

```
# Trying to create a vector of zero with length 2^32-1.
# Note that the RAM of the computer on we are generating
# this example is 8 GB with 64-bit Windows-7
# Professional edition. Processor core i5.

x <- rep(0, 2^31-1)
Error: cannot allocate vector of size 16.0 Gb</pre>
```

```
In addition: Warning messages:
1: Reached total allocation of 8078Mb: see help(memory.size)
2: Reached total allocation of 8078Mb: see help(memory.size)
3: Reached total allocation of 8078Mb: see help(memory.size)
4: Reached total allocation of 8078Mb: see help(memory.size)
```

• The preceding example clarifies that R cannot allocate a vector that has size larger than the RAM. Now we will see another example that is related to the maximally addressable range of different types of numbers. The maximum addressable range for integers is 2<sup>31</sup>-1.

```
# Maximum addressable range of inter vector
as.integer(2^31-1)
[1] 2147483647

# If we try to assign a vector of length greater than
# maximum addressable length then that will produce NA
as.integer(2^31)
[1] NA
Warning message:
NAs introduced by coercion
```

#### R and different databases

Before going on to discuss large-scale data handling using R, we will discuss how R can interact with database software through ODBC. There are two principal ways to connect to a database: the first uses the ODBC facility available on many computers and the second uses the DBI package of R along with a specialized package for the particular database needed to be accessed. If there is a specialized package available for a database, we may find that the corresponding DBI-based package gives better performance than the ODBC approach. On the other hand, if a database does not have a specialized package to access, using ODBC may be the only option.

#### R and Excel

An Excel file can be imported into R using ODBC. We will now create an ODBC connection with an MS Excel file with the connection string xlopen.

To create an ODBC connection string with an MS Excel file, we need to open the control panel of the operating system and then open **Administrative Tools** and then choose ODBC. A dialog box will now appear. Click on the **Add...** button and select an appropriate ODBC driver and then locate the desired file and give a data source name. In our case, the data source name is xlopen. The name of the Excel file can be anything, and in our case the file name is xlopen. The following R code will import the corresponding Excel file into the R environment:

```
# calling ODBC library into R
library(RODBC)

# creating connection with the database using odbc package.
# We created the connection following the steps outlined in the
# preceding paragraph

xldb<- odbcConnect("xlopen")

# In the odbcConnect() function the minimum argument required
# is the ODBC connection string.

# Now the connection created, using that connection we will import data
xldata<- sqlFetch(xldb, "CSVanscombe")

# Note here that "CSVanscombe" is the Excel worksheet name.</pre>
```

We can use other packages to import an Excel file, but at the same time R has the facility to import data using the ODBC approach. To use the ODBC approach on an Excel file, we firstly need to create the connection string using the system administrator.

#### R and MS Access

To import data from the MS Access database, the procedure is the same as with Excel. First, we need to create a connection string from the system administrator and then connect with the database from R using the RODBC package.

Let us consider the Access database containing three different tables: coveragepage, questionnaire1, and questionnaire2. The connection string to access this database is accessdata. The following command can be used to import all the three tables as separate data frames in R:

```
# calling odbc library
library(RODBC)

# connecting with database
access_con<- odbcConnect("accessdata")

# import separate table as separate R data frame
coverage_page<- sqlFetch(access_con, "coverpage")
ques1 <- sqlFetch(access_con, "questionnaire1")
ques2 <- sqlFetch(access con, "questionnaire2")</pre>
```

Using MS Excel and MS Access, we can deal with fairly large datasets, but sometimes it so happens that the dataset is too large and handling with Excel or Access is difficult. Also, Excel cannot deal with relational databases. To overcome this limitation, R has another functionality, which we will discuss in the following sections.

#### Relational databases in R

In this section, we will try to provide a concise overview of different packages in R for handling massive data and illustrate some of them.

A popular approach to dealing with bigger datasets is the use of SQL, a different programming language. It might not be difficult for someone to learn another programming language, but as we are dealing with and talking about using R, the community of R users try to develop specialized packages to deal with large datasets. Those contributed packages successfully create interfaces between R and different database software packages that use relational database management systems (RDBMS), such as MySQL (RMySQL), PostgreSQL (RPgSQL), and Oracle (ROracle). To get the full benefit of these specialized packages, we have to install third-party software, and one of the most popular packages is RMySQL. This package allows us to make connections between R and the MySQL server.

MySQL, which can deal with a mid-size, multi-platform RDBMS is a popular software in the open source community. Some of its advantages include high-performance, being open source, and being free for non-commercial use. In order to install this package properly, we need to download both the MySQL server and RMySQL.

There are several R packages available that allow direct interactions with large datasets within R, such as filehash, ff, and bigmemory. The idea is to avoid loading the whole dataset into memory.

#### The filehash package

The filehash package, which is used for solving large-data problems, was contributed by Roger Peng (The Interacting with Data using the filehash Package for R paper, available at http://cran.r-project.org/web/packages/filehash/vignettes/filehash.pdf). The idea behind the development of this package was to avoid loading the dataset into a computer's virtual memory. We must rather dump the large dataset into the hard drive and then assign an environment name for the dumped objects. Once a dataset is dumped into the hard drive, we can access the data using the assigned environment. In this way, we can deal with larger datasets and avoid the use of the computer's virtual memory and allow faster data manipulation. We will now discuss the basic steps of using this package through some examples.

Firstly, create a database that can be accessed later on. To create a database, we have to use the dbCreate function, which needs to be initialized (via dbInit) in order to be accessed, as shown in the following code. The dbInit function returns an S4 object that inherits from the filehash class.

```
library(filehash)
dbCreate("exampledb")
filehash db<- dbInit("exampledb")</pre>
```

The primary interface of filehash databases consists of the functions dbFetch, dbInsert, dbExists, dbList, and dbDelete. All of these functions are generic in nature and specific methods exist for the database that work in the backend. The first argument that is taken by the functions within this package is an object of the filehash class. To insert some data into the database, we can simply call dbInsert. We retrieve those data values with dbFetch, as shown in the following code:

```
dbInsert(filehash_db, "xx", rnorm(50))
value<- dbFetch(filehash_db, "xx")
summary(value)</pre>
```

The dbList function lists all of the keys that are available in the database, the dbExists function tests to see if a given key is in the database, and the dbDelete function deletes a key-value pair from the database, as shown in the following code:

```
dbInsert(filehash_db, "y", 4709)
dbDelete(filehash_db, "xx")
dbList(filehash_db)
dbExists(filehash_db, "xx")
```

There is another very useful command, dbLoad(), that works in a similar way to the attach() function. Using the filehash package, the objects are attached but stored on the local hard disk. We may also assess the objects in the filehash database using the usual standard R subset and accessor functions such as \$, [[, and [, as shown in the following code:

```
filehash_db$x<- runif(100)
summary(filehash_db$x)
summary(filehash_db[["x"]])
filehash_db$y<- rnorm(100, 2)
dbList(filehash db)</pre>
```

After initializing a database using the default DB1 format, it opens a file connection for reading and writing to the database file on the disk. This file connection will remain open until the database is closed via dbDisconnect or the database object in R is removed. There is a limit on the number of file connections that can be open at the same time, so to protect any database from unexpected results, we need to make sure the file connections are closed properly.

Just like save.image in base R, there are some utilities included in the filehash package and two of them are dumpObjects and dumpImage. The dumpObjects utility saves an object into the filehash database so that it can be accessed in the future if required. It does not save objects into R itself, which allows faster processing. Similarly, dumpImage saves the entire workspace to a filehash database. The dumpList function takes a list and creates a filehash database with values from the list. The list must have a non-empty name for every element in order for dumpList to succeed. The dumpDF utility creates a filehash database from a data frame where each column of the data frame is an element in the database. Essentially, dumpDF converts the data frame to a list and then calls dumpList. The following example shows how we can use dumpDF:

```
dumpDF(read.table("anscombe.txt", header=T), dbName="massivedata")
massive environment<- db2env(db="massivedata")</pre>
```

The first element of dumpDF() is a data object. R will read the data within dumpDF(), so its memory does not have a copy of it. Space saved! So now our pretended large dataset, anscombe.txt, can be accessed through the env01 environment. To access it, we use with(). Suppose we want to perform a linear regression of y on x and access the data using the variable names. In such cases, if you assign an object name for the read.table command, the memory will have a copy of the data, which is not desirable. Using the with() function, we can fit a model or compute summary statistics as usual as follows:

```
fit<- with(massive_environment, lm(Y1~X1))
with(massive_environment, summary(Y1))
with(massive environment, Y1[1] <- 99))</pre>
```

#### The ff package

As we have seen in the example in the introductory section of this chapter, R can only address objects that fit within the memory limits of its RAM and the maximally addressable range of 2<sup>31</sup>-1 bytes. To overcome this limitation, Adler and Glaser, in 2010, developed the ff package. This package extends the R system and stores data in the form of native binary flat files in persistent storage such as hard disks, CDs, or DVDs rather than in the RAM. This package enables users to work on several large datasets simultaneously. It also allows the allocation of vectors or arrays that are larger than the RAM. The package comprises of two parts: one is the low-level layer written in C++ and the other is the high-level layer in R. This package is designed for convenient access to large datasets.

As users will only deal with the high-level layer, the following are the tasks we do in this layer:

- Opening/creating flat files: There are two basic functions, ff and ffm, to deal with opening and creating flat files. If we specify the length argument or the dim argument, a new file is created, otherwise R will open an existing file.
- **I/O operations**: These operations are controlled by the [] (for reading) and the [] <- (for writing) operators.
- Generic functions and methods for the ff and ffm objects: Methods for dim and length are provided and the sample function is converted to a generic function.

The primary argument for the functions ff and ffm require a filename in the file argument to specify the flat file. Whenever length (for ff) or dim (for ffm) is specified, as shown in the following code, a new flat file is created, otherwise an existing file is opened:

```
# A flat file with a length 10 is created
library(ff)
file1 <- ff(filename="file1", length=10, vmode="double")</pre>
str(file1)
list()
 - attr(*, "physical")=Class 'ff pointer' <externalptr>
  ..- attr(*, "vmode") = chr "double"
  ..- attr(*, "maxlength") = int 10
  ..- attr(*, "pattern") = chr "/"
  ..- attr(*, "filename") = chr "D:/Book on R/Writing/outline/data ch2/
file1"
  ..- attr(*, "pagesize") = int 65536
  ..- attr(*, "finalizer") = chr "close"
  ..- attr(*, "finonexit") = logi TRUE
  ... attr(*, "readonly") = logi FALSE
  ..- attr(*, "caching") = chr "mmnoflush"
 - attr(*, "virtual") = list()
  ..- attr(*, "Length") = int 10
  ... attr(*, "Symmetric") = logi FALSE
 - attr(*, "class") = chr [1:2] "ff vector" "ff"
```

The entries of file1 can be modified with the []<- operator. For example, the first 10 entries of the rivers dataset that contains the length of the 141 rivers in North America can be stored in an ff object, as shown in the following code:

```
# calling rivers data
data(rivers)
file1[1:10] <- rivers[1:10]

# Note that here file1 is an ff object whereas
# file1[...] returns default R vector
str(file1)</pre>
```

If required, we can perform sampling on the ff objects as follows:

```
# set seed to reproduce the example
set.seed(1337)
sample(file1,5,replace=FALSE)
[1] 735 392 524 450 600
```

Flat file objects are referenced when forming R objects using external pointers. In order to clear the references, the garbage collector, gc(), can be used as follows:

```
gc()
```

Calling gc() clears the reference to the file, but does not delete the file from the hard drive. Since the data is still present, the flat file can be opened again at a later stage.

#### R and sqldf

The sqldf package is an R package that allows users to run SQL statements within R. SQL is the popular programming language for manipulating data from relational databases, and the sqldf package creates an opportunity to work directly with SQL statements on an R data frame. With this package, the user can do the following tasks easily:

- Write alternate syntax for data frame manipulation, particularly for purposes
  of faster processing, since using sqldf (with SQLite as the underlying
  database) is often faster compared to performing the same manipulations
  in built-in R functions
- Read portions of large files into R without reading the entire file

The user need not perform the following tasks once they use sqldf because these are automatically done:

- Database setup
- Writing the create table statement, which defines each table
- Importing and exporting to and from the database
- The coercing of the returned columns to the appropriate class in common cases

#### Data manipulation using sqldf

We can perform any type of data manipulation to an R data frame either in memory or during import. The following example shows the selection of a portion of the iris dataset using the sqldf package:

```
# Selecting the rows from iris dataset where sepal length > 2.5
# and store that in subiris data frame
library(sqldf)
subiris<- sqldf("select * from iris where Sepal Width> 3")
head(subiris)
   Sepal Length Sepal Width Petal Length Petal Width Species
1
           5.1
                        3.5
                                      1.4
                                                   0.2
                                                        setosa
2
           4.7
                        3.2
                                      1.3
                                                  0.2
                                                        setosa
                        3.1
                                      1.5
3
           4.6
                                                  0.2
                                                        setosa
           5.0
                        3.6
                                      1.4
                                                  0.2
4
                                                        setosa
5
           5.4
                        3.9
                                      1.7
                                                  0.4
                                                        setosa
           4.6
                        3.4
                                      1.4
                                                  0.3
                                                        setosa
nrow(subiris)
[1] 67
```

We can also select a smaller number of columns while filtering out some of the rows with a specified condition. The following example selects only sepal length, petal length, and species; however, this time, rows are filtered by values for petal length greater than 1.4:

```
subiris2<-
sqldf("select Sepal Length, Petal Length, Species from iris where Petal
Length> 1.4")
nrow(subiris2)
[1] 126
head(subiris2)
  Sepal_Length
                  Petal Length
                                    Species
1
            4.6
                        1.5
                                    setosa
2
                        1.7
            5.4
                                    setosa
3
            5.0
                        1.5
                                    setosa
                        1.5
4
            4.9
                                    setosa
5
            5.4
                        1.5
                                    setosa
            4.8
                        1.6
6
                                    setosa
```

If the dataset is too large and cannot entirely be read into the R environment, we can import a portion of that dataset using sqldf. The following example shows how we can import a portion of a csv file using the sqldf functionality. We will use the read.csv.sql() function to perform this task. This is an interface to sqldf that works like read.csv in R, except that it also provides a sql= argument. Not all of the other arguments of read.csv are supported.

In the following example, we will import the iris.csv file. We will import only sepal width and petal width along with the species information where petal width is greater than 0.4:

```
iriscsv<-read.csv.sql("iris.csv",sql="select</pre>
Sepal Width, Petal Width, Species from file where Petal Width>0.4")
head(iriscsv)
     Sepal Width Petal Width
                                  Species
          3.3
                      0.5
1
                                  "setosa"
          3.5
                      0.6
2
                                 "setosa"
          3.2
                      1.4
                              "versicolor"
3
4
          3.2
                      1.5
                              "versicolor"
         3.1
5
                     1.5
                              "versicolor"
```

"versicolor"

An important thing to note is that in the original iris.csv file, the variable names were dot separated, but when we pass a SQL statement, we need to use an underscore as the variable name, otherwise it will output an error as follows:

1.3

2.3

6

```
iriscsv<-read.csv.sql("iris.csv",sql="select Sepal.Width,Petal.
Width,Species from file where Petal.Width>0.4")

Error in sqliteExecStatement(con, statement, bind.data) :
   RS-DBI driver: (error in statement: no such column: Sepal.Width)
```

We sometimes need to draw a random sample from a dataset but the original data file might be too large. In the following example, we will show how we can draw a random sample size of 10 from the iris data that is stored in the iris.csv file:

```
iris_sample<-
read.csv.sql("iris.csv",sql="select * from file order by random(*)
limit 10")
iris sample</pre>
```

	Sepal_Length Sep	al_Width P	etal_Length Pet	al_Widt	h Species
1	6.5	3.0	5.2	2.0	"virginica"
2	5.0	3.5	1.3	0.3	"setosa"
3	6.0	2.2	4.0	1.0	"versicolor"
4	6.9	3.1	5.4	2.1	"virginica"
5	6.2	2.8	4.8	1.8	"virginica"
6	5.1	3.8	1.9	0.4	"setosa"
7	5.8	2.6	4.0	1.2	"versicolor"
8	5.9	3.2	4.8	1.8	"versicolor"
9	6.4	2.9	4.3	1.3	"versicolor"
10	6.4	3.1	5.5	1.8	"virginica"

We can perform group-wise processing and aggregation using sqldf, which is a faster alternative to the aggregate function. For example, if we want to calculate the mean of each variable in the iris data for each species, the following is the code:

The base R counterpart for performing the same operation is as follows:

```
aggregate(iris[,-5],list(iris$Species),mean)
```

	Group.1	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
1	setosa	5.006	3.428	1.462	0.246
2	versicolor	5.936	2.770	4.260	1.326
3	virginica	6.588	2.974	5.552	2.026

Though both functions give us the same results, for larger datasets, sqldf is much faster than base R.

#### Summary

At the beginning of this chapter, we showed you how we can deal with an MS Excel file as a database and how an MS Access database table can be imported into R. One of the major problems in R is that its memory is bound by the system virtual memory, and that is why the data should be smaller in size than the memory of a dataset to be able to work with it. But in reality, datasets are often larger than the virtual memory and sometimes the length of the array or vector exceeds the maximum addressable range. To overcome these two limitations, R can be utilized with relational databases. Contributed R packages exist to help in dealing with such large datasets, and they have been highlighted in this chapter, particularly filehash and ff. We also discussed sqldf for faster data manipulation.

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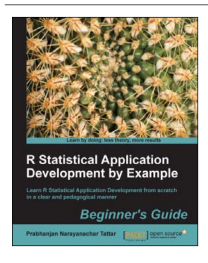


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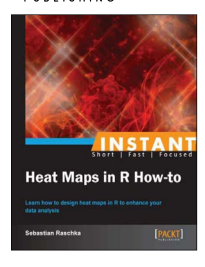
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