CLEANING UP

We have defined several variables in the course of this section, some of which are no longer needed, so it is time to clean up:

```
> objects()

[1] "channel" "condition" "cooperation" "Guyer"
[5] "names" "sex" "v" "x"

> remove(channel, names, v, x)
> detach(package:RODBC)
```

We have retained the data frame Guyer and the vectors condition, cooperation, and sex for subsequent illustrations in this chapter. Because we are finished with the **RODBC** package, we have detached it.

2.1.5 GETTING DATA OUT OF R

We hope and expect that you will rarely have to get your data out of R to use with another program, but doing so is nevertheless quite straightforward. As in the case of reading data, there are many ways to proceed, but a particularly simple approach is to use the write.table or write.csv function to output a data frame to a plain-text file. The syntax for write.table is essentially the reverse of that for read.table. For example, the following command writes the Duncan data frame (from the attached car package) to a file:

```
> write.table(Duncan, "c:/temp/Duncan.txt")
```

By default, row labels and variable names are included in the file, data values are separated by blanks, and all character strings are in quotes, whether or not they contain blanks. This default behavior can be changed—see ?write.table.

The **foreign** package also includes some functions for exporting R data to a variety of file formats: Consult the documentation for the **foreign** package, help(package="foreign").

2.2 Working With Data Frames

It is perfectly possible in R to analyze data stored in vectors, but we generally prefer to begin with a data frame, typically read from a file via the read.table function or accessed from an R package. Almost all the examples in this *Companion* use data frames from the **car** package.

In many statistical packages, such as SPSS, a single data set is active at any given time; in other packages, such as SAS, individual statistical procedures typically draw their data from a single source, which by default in SAS is the

last data set created. This is not the case in R, where data may be used simultaneously from several sources, providing flexibility but with the possibility of interference and confusion.

There are essentially two ways to work with data in a data frame, both of which we will explain in this section:

- Attach the data frame to the search path via the attach command, making the variables in the data frame directly visible to the R interpreter.
- 2. Access the variables in the data frame as they are required without attaching the data frame.

New users of R generally prefer the first approach, probably because it is superficially similar to other statistical software in which commands reference an *active* data set. For reasons that we will explain, however, experienced users of R usually prefer *not* to attach data frames to the search path.

2.2.1 THE SEARCH PATH

When you type the name of a variable in a command, the R interpreter looks for an object of that name in the locations specified by the search path. We attach the Duncan data frame to the search path with the attach function and then use the search function to view the current path:

```
> attach(Duncan)
> search()

[1] ".GlobalEnv" "Duncan" "package:car"

[4] "package:survival" "package:splines" "package:leaps"

[7] "package:nnet" "package:MASS" "package:stats"

[10] "package:graphics" "package:grDevices" "package:utils"

[13] "package:datasets" "package:methods" "Autoloads"

[16] "package:base"
```

Now if we type the name of the variable prestige at the command prompt, R will look first in the global environment (.GlobalEnv), the region of memory in which R stores working data. If no variable named prestige is found in the global environment, then the data frame Duncan will be searched, because it was placed by the attach command in the second position on the search list. There is in fact no variable named prestige in the working data, but there is a variable by this name in the Duncan data frame, and so when we type prestige, we retrieve the prestige variable from Duncan, as we may readily verify:

```
> prestige
[1] 82 83 90 76 90 87 93 90 52 88 57 89 97 59 73 38 76 81 45 92
[21] 39 34 41 16 33 53 67 57 26 29 10 15 19 10 13 24 20 7 3 16
[41] 6 11 8 41 10
```

```
> Duncan$prestige
```

```
[1] 82 83 90 76 90 87 93 90 52 88 57 89 97 59 73 38 76 81 45 92 [21] 39 34 41 16 33 53 67 57 26 29 10 15 19 10 13 24 20 7 3 16 [41] 6 11 8 41 10
```

Typing Duncan\$prestige directly extracts the column named prestige from the Duncan data frame.⁶

Had prestige not been found in Duncan, then the sequence of attached packages would have been searched in the order shown, followed by a special list of objects (Autoloads) that are loaded automatically as needed (and which we will subsequently ignore), and finally the R base package. The packages in the search path shown above, beginning with the stats package, are part of the basic R system and are loaded by default when R starts up.

Suppose, now, that we attach the Prestige data frame to the search path. The default behavior of the attach function is to attach a data frame in the second position on the search path, after the global environment:

```
> attach(Prestige)
       The following object(s) are masked from Duncan :
        education income prestige type
       The following object(s) are masked from package:datasets:
        women
> search()
 [1] ".GlobalEnv"
                      "Prestige"
                                         "Duncan"
[4] "package:car"
                      "package:survival" "package:splines"
[7] "package:leaps"
                      [10] "package:stats"
                      "package:graphics" "package:grDevices"
                      "package:datasets" "package:methods"
[13] "package:utils"
[16] "Autoloads"
                      "package:base"
```

Consequently, the data frame Prestige is attached *before* the data frame Duncan; and if we now simply type prestige, then the prestige variable in Prestige will be located *before* the prestige variable in Duncan is encountered:

```
> prestige

[1] 68.8 69.1 63.4 56.8 73.5 77.6 72.6 78.1 73.1 68.8

[11] 62.0 60.0 53.8 62.2 74.9 55.1 82.3 58.1 58.3 72.8

[21] 84.6 59.6 66.1 87.2 66.7 68.4 64.7 34.9 72.1 69.3

...

[91] 38.9 36.2 29.9 42.9 26.5 66.1 48.9 35.9 25.1 26.1

[101] 42.2 35.2
```

⁶Information on indexing data frames is presented in Section 2.3.4.

The prestige variable in Duncan is still there—it is just being *shadowed* or *masked* (i.e., hidden) by prestige in Prestige, as the attach command warned us:

```
> Duncan$prestige
[1] 82 83 90 76 90 87 93 90 52 88 57 89 97 59 73 38 76 81 45 92
[21] 39 34 41 16 33 53 67 57 26 29 10 15 19 10 13 24 20 7 3 16
[41] 6 11 8 41 10
```

Because the variables in one data frame can shadow the variables in another, attaching more than one data frame at a time can lead to unanticipated problems and should generally be avoided. You can remove a data frame from the search path with the detach command:

```
> detach(Prestige)
> search()

[1] ".GlobalEnv" "Duncan" "package:car"

[4] "package:survival" "package:splines" "package:leaps"

[7] "package:nnet" "package:MASS" "package:stats"

[10] "package:graphics" "package:grDevices" "package:utils"

[13] "package:datasets" "package:methods" "Autoloads"

[16] "package:base"
```

Calling detach with no arguments detaches the second entry in the search path and, thus, produces the same effect as detach (Prestige).

Now that Prestige has been detached, prestige again refers to the variable by that name in the Duncan data frame:

```
> prestige
[1] 82 83 90 76 90 87 93 90 52 88 57 89 97 59 73 38 76 81 45 92
[21] 39 34 41 16 33 53 67 57 26 29 10 15 19 10 13 24 20 7 3 16
[41] 6 11 8 41 10
```

The working data are the first item in the search path, and so globally defined variables shadow variables with the same names anywhere else along the path. This is why we use an uppercase letter at the beginning of the name of a data frame. Had we, for example, named the data frame prestige rather than Prestige, then the variable prestige within the data frame would have been shadowed by the data frame itself. To access the variable would then require a potentially confusing expression, such as prestige\$prestige.

Our focus here is on manipulating data, but it is worth mentioning that R locates functions in the same way that it locates data. Consequently, functions earlier on the path can shadow functions of the same name later on the path.

In Section 1.1.3, we defined a function called myMean, avoiding the name mean so that the mean function in the **base** package would not be shadowed. The **base** function mean can calculate trimmed means as well as the ordinary arithmetic mean; for example,

```
> mean(prestige)
[1] 47.68889
> mean(prestige, trim=0.1)
[1] 47.2973
```

Specifying mean (prestige, trim=0.1) removes the top and bottom 10% of the data, calculating the mean of the middle 80% of observations. Trimmed means provide more efficient estimates of the center of a heavy-tailed distribution—for example, when outliers are present; in this example, however, trimming makes little difference.

Suppose that we define our own mean function, making no provision for trimming:

```
> mean <- function(x) {
+     warning("the mean function in the base package is shadowed")
+     sum(x)/length(x)
+ }</pre>
```

The first line in our mean function prints a warning message. The purpose of the warning is simply to verify that our function executes in place of the mean function in the **base** package. Had we *carelessly* shadowed the standard mean function, we would not have politely provided a warning:

```
> mean(prestige)
[1] 47.68889
Warning message:
In mean(prestige):the mean function in the base package is shadowed
```

The essential point here is that because our mean function resides in the global environment, it is encountered on the search path *before* the mean function in the **base** package. Shadowing the standard mean function is inconsequential as long as our function is equivalent; but if, for example, we try to calculate a trimmed mean, our function does not work:

```
> mean(prestige, trim=0.1)
Error in mean(prestige, trim = 0.1):unused argument(s)(trim = 0.1)
```

Shadowing standard R functions is a practice generally to be avoided. Suppose, for example, that a robust-regression function tries to calculate a trimmed mean, but fails because the standard mean function is shadowed by our redefined mean function. If we are not conscious of this problem, the resulting error message may prove cryptic.

We can, however, use the same name for a variable and a function, as long as the two do not reside in the working data. Consider the following example:

```
> mean <- mean(prestige) # uses then overwrites our mean function
> mean
[1] 47.68889
```

Specifying mean <- mean (prestige) causes our mean function to calculate the mean prestige and then stores the result in a variable called mean, which has the effect of destroying our mean function (and good riddance to it). The variable mean in the working data does not, however, shadow the function mean in the base package:

```
> mean(prestige, trim=0.1)
[1] 47.2973
```

Before proceeding, let us tidy up a bit:

```
> remove(mean)
> detach(Duncan)
```

2.2.2 AVOIDING attach

Here are some compelling reasons for *not* attaching data frames to the search path:

- Variables in attached data frames may mask other objects, and variables in attached data frames themselves may be masked by objects of the same name—for example, in the global environment.
- Attaching a data frame makes a copy of the data frame; the attached version is a snapshot of the data frame at the moment when it is attached. If changes are made to the data frame, these are not reflected in the attached version of the data. Consequently, after making such a change, it is necessary to detach and reattach the data frame. We find this procedure awkward, and inexperienced users of R may not remember to detach and reattach the data, leading to confusion about the current state of the attached data.
- We have observed that new users of R tend not to detach data frames after they are done with them. Often they will attach multiple versions of a data frame in the same session, which potentially results in confusion.

There are several strategies that we can use to avoid attaching a data frame:

• We can reference the variables in a data frame explicitly: for example,

```
> mean(Duncan$prestige)
[1] 47.68889
```

• Statistical-modeling functions in R usually include a data argument, which can be set to a data frame, conveniently specifying where the data for the model are to be found: for example,

• The with command can be used to evaluate an R expression in the environment of a data frame: for example,

We will use with frequently in the rest of this Companion.

2.2.3 MISSING DATA

Missing data are a regrettably common feature of real data sets. Two kinds of issues arise in handling missing data:

- There are relatively profound statistical issues concerning how best to use available information when missing data are encountered (see, e.g., Little and Rubin, 2002; and Schafer, 1997). We will ignore these issues here, except to remark that R is well designed to make use of sophisticated approaches to missing data.⁷
- There are intellectually trivial but often practically vexing mechanical issues concerning computing with missing data in R. These issues, which are the subject of the present section, arise partly because of the diverse data structures and kinds of functions available simultaneously to the R user. Similar issues arise in *all* statistical software, however, although they may sometimes be disguised.

As we have explained, on data input, missing values are typically encoded by the characters NA. The same characters are used to print missing information. Many functions in R know how to handle missing data, although sometimes they have to be explicitly told what to do.

To illustrate, let us examine the data set Freedman in the car package:

```
> head(Freedman) # first 6 rows
          population nonwhite density crime
               675 7.3 746 2602
Akron
                713
                        2.6
                               322 1388
Albany
                               NA 5018
Albuquerque
                NA
                        3.3
Allentown
                534
                        0.8
                               491 1182
                              1612 3341
Anaheim
               1261
                       1.4
Atlanta
               1330
                       22.8
                              770 2805
              # number of rows and columns
> dim(Freedman)
[1] 110
```

⁷Notable packages for handling missing data include amelia, mi, mice, and norm, which perform various versions of multiple imputation of missing data.

These data, on 110 U.S. metropolitan areas, were originally from the 1970 Statistical Abstract of the United States and were used by Freedman (1975) as part of a wide-ranging study of the social and psychological effects of crowding. Freedman argues, by the way, that high density tends to intensify social interaction, and thus the effects of crowding are not simply negative. The variables in the data set are as follows:

- population: Total 1968 population, in thousands.
- nonwhite: Percent nonwhite population in 1960.
- density: Population per square mile in 1968.
- crime: Number of serious crimes per 100,000 residents in 1969.

Some of Freedman's data are missing—for example, the population and density for Albuquerque. Here are the first few values for density:

```
> head(Freedman$density, 20) # first 20 values
[1] 746 322 NA 491 1612 770 41 877 240 147 272 1831
[13] 1252 832 630 NA NA 328 308 1832
```

Suppose, now, that we try to calculate the median density; as we will see shortly, the density values are highly positively skewed, so using the *mean* as a measure of the center of the distribution would be a bad idea:

```
> median(Freedman$density)
[1] NA
```

R tells us that the median density is missing. This is the pedantically correct answer: Several of the density values are missing, and consequently we cannot in the absence of those values know the median, but this is probably not what we had in mind when we asked for the median density. By setting the na.rm (NA-remove) argument of median to TRUE, we instruct R to calculate the median of the remaining, nonmissing values:

```
> median(Freedman$density, na.rm=TRUE)
[1] 412
```

Several other R functions that calculate statistical summaries, such as mean, var (variance), sd (standard deviation), and quantile (quantiles), also have na.rm arguments, but not all R functions handle missing data in this manner.

Most plotting functions simply ignore missing data. For example, to construct a scatterplot of crime against density, including only the observations with valid data for both variables, we enter

```
> with(Freedman, {
+    plot(density, crime)
+    identify(density, crime, row.names(Freedman))
+ ))
```

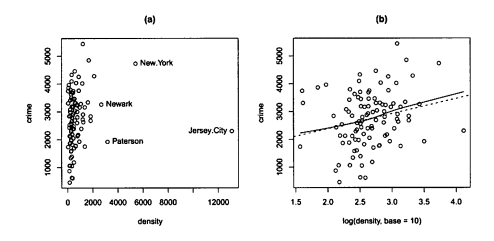


Figure 2.1 Scatterplot of crime by population density for Freedman's data. (a) Original density scale, with a few high-density cities identified interactively with the mouse, and (b) log-density scale, showing linear least-squares (broken) and lowess nonparametric-regression (solid) lines. Cases with one or both values missing are silently omitted from both graphs.

The resulting graph, including several observations identified with the mouse, appears in Figure 2.1a. Recall that we identify observations by pointing at them with the mouse and clicking the left mouse button; exit from identify by pressing the esc key in Mac OS X or, in Windows, by clicking the right mouse button and selecting *Stop*. It is apparent that density is highly positively skewed, making the plot very difficult to read. We would like to try plotting crime against the log of density but wonder whether the missing data will spoil the computation. The log function in R behaves sensibly, however: The result has a missing entry wherever—and only where—there was a missing entry in the argument:

```
> log(c(1, 10, NA, 100), base=10)
[1] 0 1 NA 2
```

Other functions that compute on vectors in an element-wise fashion—such as the arithmetic operators—behave similarly.

We, therefore, may proceed as follows, producing the graph in Figure 2.1b:9

```
> with(Freedman, plot(log(density, base=10), crime))
```

This graph is much easier to read, and it now appears that there is a weak, positive relationship between crime and density. We will address momentarily how to produce the lines in the plot.

Statistical-modeling functions in R have a special argument, na.action, which specifies how missing data are to be handled; na.action is set to

⁸Transformations, including the log transformation, are the subject of Section 3.4.

⁹An alternative would have been to plot crime against density using a log axis for density: plot(density, crime, log="x"). See Chapters 3 and 7 for general discussions of plotting data in R.

a function that takes a data frame as an argument and returns a similar data frame composed entirely of valid data (see Section 4.8.5). The default na-action is na.omit, which removes all observations with missing data on any variable in the computation. All the examples in this Companion use na.omit. An alternative, for example, would be to supply an na.action that imputes the missing values.

The prototypical statistical-modeling function in R is 1m, which is described extensively in Chapter 4. For example, to fit a linear regression of crime on the log of density, removing observations with missing data on either crime or density, enter the command

The 1m function returns a linear-model object; because the returned object was not saved in a variable, the interpreter simply printed a brief report of the regression. To plot the least-squares line on the scatterplot in Figure 2.1:

```
> abline(lm(crime ~ log(density, base=10), data=Freedman),
lty="dashed")
```

The linear-model object returned by 1m is passed to abline, which draws the regression line; specifying the line type 1ty="dashed" produces a broken line.

Some functions in R, especially the older ones, make no provision for missing data and simply fail if an argument has a missing entry. In these cases, we need somewhat tediously to handle the missing data ourselves. A relatively straightforward way to do so is to use the complete.cases function to test for missing data, and then to exclude the missing data from the calculation.

For example, to locate all observations with valid data for both crime and density, we enter:

```
> good <- with(Freedman, complete.cases(crime, density))
> head(good, 20) # first 20 values

[1] TRUE TRUE FALSE TRUE TRUE TRUE TRUE TRUE TRUE TRUE
[11] TRUE TRUE TRUE TRUE TRUE FALSE FALSE TRUE TRUE TRUE
```

We then use good to select the valid observations by indexing (a topic described in Section 2.3.4). For example, it is convenient to use the lowess function to add a nonparametric-regression smooth to our scatterplot (Figure 2.1b), but lowess makes no provision for missing data:¹⁰

¹⁰The lowess function is described in Section 3.2.1.

```
> with (Freedman,
+ lines(lowess(log(density[good], base=10), crime[good], f=1.0)))
```

By indexing the predictor density and response crime with the logical vector good, we extract only the observations that have valid data for both variables. The argument f to the lowess function specifies the span of the lowess smoother-that is, the fraction of the data included in each localregression fit; large spans (such as the value 1.0 employed here) produce smooth regression curves.

Suppose, as is frequently the case, that we analyze a data set with a complex pattern of missing data, fitting several statistical models to the data. If the models do not all use exactly the same variables, then it is likely that they will be fit to different subsets of nonmissing observations. Then if we compare the models with a likelihood ratio test, for example, the comparison will be invalid.11

To avoid this problem, we can first use na. omit to filter the data frame for missing data, including all the variables that we intend to use in our data analysis. For example, for Freedman's data, we may proceed as follows, assuming that we want subsequently to use all four variables in the data frame:

```
> head(Freedman.good) # first 6 rows
          population nonwhite density crime
               675 7.3
                713
                        2.6
```

> Freedman.good <- na.omit(Freedman)

746 2602 Akron 322 1388 Albany 491 1182 534 0.8 Allentown Anaheim 1261 1.4 1612 3341 1330 22.8 770 2805 Atlanta 331 7.0 41 3306 Bakersfield

> dim(Freedman.good) # number of rows and columns

[1] 100

A note of caution: Filtering for missing data on variables that we do not intend to use can result in discarding data unnecessarily. We have seen cases where students and researchers inadvertently and needlessly threw away most of their data by filtering an entire data set for missing values, even when they intended to use only a few variables in the data set.

Finally, a few words about testing for missing data in R: A common error is to assume that one can check for missing data using the == (equals) operator, as in

```
> NA == c(1, 2, NA, 4)
[1] NA NA NA NA
```

Testing equality of NA against any value in R returns NA as a result. After all, if the value is missing, how can we know whether it's equal to something else? The proper way to test for missing data is with the function is.na:

¹¹How statistical-modeling functions in R handle missing data is described in Section 4.8.5.

```
> is.na(c(1, 2, NA, 4))
[1] FALSE FALSE TRUE FALSE
```

For example, to count the number of missing values in the Freedman data frame:

```
> sum(is.na(Freedman))
[1] 20
```

This command relies on the automatic *coercion* of the logical values TRUE and FALSE to one and zero, respectively.

2.2.4 NUMERIC VARIABLES AND FACTORS

If we construct R data frames by reading data from text files using read. - table or from numeric and character vectors using data. frame, then our data frames will consist of two kinds of data: numeric variables and factors. Both read.table and data.frame by default translate character data and logical data into factors.

Before proceeding, let us clean up a bit:

```
> objects()

[1] "condition" "cooperation" "Freedman.good"

[4] "good" "Guyer" "sex"

> remove(good, Freedman.good)
```

Near the beginning of this chapter, we entered data from Fox and Guyer's (1978) experiment on anonymity and cooperation into the global variables cooperation, condition, and sex.¹² The latter two variables are character vectors, as we verify for condition:

```
> condition
```

```
[1] "public" "public" "public" "public" "public" [6] "public" "public" "public" "public" "public" "public" [11] "anonymous" "a
```

We can confirm that this is a vector of character values using the *predicate function* is.character, which tests whether its argument is of mode "character":

```
> is.character(condition)
```

[1] TRUE

¹²Variables created by assignment at the command prompt are global variables defined in the working data.

After entering the data, we defined the data frame Guyer, which also contains variables named cooperation, condition, and sex. We will remove the global variables and will work instead with the data frame:

```
> remove(cooperation, condition, sex)
```

Look at the variable condition in the Guyer data frame:

```
> Guyer$condition
```

```
public
 [1] public
              public
                       public
                                           public
                                                    public
                                public
[7] public
              public
                       public
                                           anonymous anonymous
[13] anonymous anonymous anonymous anonymous anonymous
[19] anonymous anonymous
Levels: anonymous public
> is.character(Guyer$condition)
[1] FALSE
> is.factor(Guyer$condition)
[1] TRUE
```

As we explained, condition in the data frame is a factor rather than a character vector. A factor is a representation of a categorical variable; factors are stored more economically than character vectors, and the manner in which they are stored saves information about the levels (category set) of a factor. When a factor is printed, its values are not quoted, as are the values of a character vector, and the levels of the factor are listed.

Many functions in R, including statistical-modeling functions such as 1m, know how to deal with factors. For example, when the generic summary function is called with a data frame as its argument, it prints various statistics for a numeric variable but simply counts the number of observations in each level of a factor:

> summary(Guyer)

Factors have unordered levels. An extension, called *ordered factors*, is discussed, along with factors, in the context of linear models in Section 4.6.

2.2.5 MODIFYING AND TRANSFORMING DATA

Data modification in R usually occurs naturally and unremarkably. When we wanted to plot crime against the log of density in Freedman's data, for

example, we simply specified log (density, base=10).¹³ Similarly, in regressing crime on the log of density, we just used log (density, base=10) on the right-hand side of the linear-model formula.

Creating new variables that are functions of other variables is straightforward. For example, the variable cooperation in the Guyer data set counts the number of cooperative choices out of a total of 120 choices. To create a new variable with the percentage of cooperative choices in each group:

```
> perc.coop <- 100*Guyer$cooperation/120
```

The new variable perc. coop resides in the working data, not in the Guyer data frame. It is generally advantageous to add new variables such as this to the data frame from which they originate: Keeping related variables together in a data frame avoids confusion, for example.

```
> Guyer$perc.coop <- 100*Guyer$cooperation/120
> head(Guyer) # first 6 rows
 cooperation condition sex perc.coop
1
    49 public male 40.83333
2
        64 public male 53.33333
3
       37 public male 30.83333
4
       52 public male 43.33333
       68 public male 56.66667
5
6
        54
              public female 45.00000
```

A similar procedure may be used to *modify* an existing variable in a data frame. The following command, for example, replaces the original cooperation variable in Guyer with the logit (log-odds) of cooperation:

The transform function can be used to create and modify several variables in a data frame at once. For example, if we have a data frame called Data with variables named a, b, and c, then the command

```
> Data <- transform(Data, c=-c, asq=a^2, a.over.b=a/b)</pre>
```

replaces Data by a new data frame in which the variables a and b are included unchanged, c is replaced by -c, and two new variables are added—asq, with the squares of a, and a.over.b, with the ratios of a to b.

¹³We did not have to create a new variable, say log.density <- log(density, 10), as one may be required to do in a typical statistical package such as SAS or SPSS.

Transforming numeric data is usually a straightforward operation—simply using mathematical operators and functions. Categorizing numeric data and recoding categorical variables are often more complicated matters. Several functions in R are employed to create factors from numeric data and to manipulate categorical data, but we will limit our discussion to three that we find particularly useful: (1) the standard R function cut, (2) the recode function in the car package, and (3) the standard ifelse function.

The cut function dissects the range of a numeric variable into class intervals, or bins. The first argument to the function is the variable to be binned; the second argument gives either the number of equal-width bins or a vector of cut points at which the division is to take place. For example, to divide the range of perc. coop into four equal-width bins, we specify

```
> Guyer$coop.4 <- cut(Guyer$perc.coop, 4)
> summary(Guyer$coop.4)

(22.5,33.3] (33.3,44.2] (44.2,55] (55,65.9]
6 7 5 2
```

R responds by creating a factor, the levels of which are named for the end points of the bins. In the example above, the first level includes all values with Guyer\$perc.coop greater than 22.5 (which is slightly smaller than the minimum value of Guyer\$perc.coop) and less than or equal to 33.3, the cut point between the first two levels. Because perc.coop is not uniformly distributed across its range, the several levels of coop. 4 contain different numbers of observations. The output from the summary function applied to a factor gives a one-dimensional table of the number of observations in each level of the factor.

Suppose, alternatively, that we want to bin perc.coop into three levels containing roughly equal numbers of observations¹⁴ and to name these levels "low", "med", and "high"; we may proceed as follows:

The quantile function is used to locate the cut points. Had we wished to divide perc.coop into four groups, for example, we would simply have specified different quantiles, c(0, .25, .5, .75, 1), and of course supplied four values for the labels argument.

¹⁴Roughly equal numbers of observations in the three bins are the best we can do because n = 20 is not evenly divisible by 3.

The recode function in the car package, which is more flexible than cut, can also be used to dissect a quantitative variable into class intervals: for example,

The recode function works as follows:

- The first argument is the variable to be recoded, here perc. coop.
- The second argument is a character string, enclosed in single or double quotes, containing the recode specifications.
- Recode specifications are of the form old.values=new.value. There may be several recode specifications, separated by semicolons.
- The old.values may be a single value, including NA; a range, of the form minimum:maximum, as in the example, where the special values 10 and hi have been used to stand in for the smallest and largest values of the variable; a vector of values, typically specified with the c (combine) function; or the special symbol else, which, if present, should appear last.
- An observation that fits into more than one recode specification is assigned the value of the first one encountered. For example, a group with perc.coop exactly equal to 50 would get the new value 1.
- Character data may appear both as old.values and as new.value.
 You must be careful with quotation marks, however: If single quotes are employed to enclose the recode specifications, then double quotes must be used for the values (and vice versa).
- When a factor is recoded, the old.values should be specified as character strings; the result is a factor, even if the new.values are numbers, unless the argument as.factor.result is set to FALSE.
- Character data may be recoded to numeric, and vice versa. To recode
 a character or numeric variable to a factor, set as.factor.result
 =TRUE.
- If an observation does not satisfy any of the recode specifications, then the old.value for that observation is carried over into the result.

To provide a richer context for additional illustrations of the use of recode, we turn our attention to the Womenlf data frame from the car package:

```
> set.seed(12345) # for reproducibility
> (sample.20 <- sort(sample(nrow(Womenlf), 20))) # 20 random obs.

[1] 1 9 39 43 44 84 96 98 100 115 119 131 185 186 190
[16] 199 230 231 233 252

> Womenlf[sample.20, ] # 20 randomly selected rows
```

```
partic hincome children
                            region
   not.work 15 present Ontario
9
   not.work
              15 present Ontario
39 not.work
              9 present Atlantic
43 parttime
               28 absent Ontario
44 not.work
               23 present Ontario
   fulltime
84
               17 present Ontario
96 not.work
               17 present Ontario
               15
98 fulltime
                   absent Ontario
               15 present Ontario
100 not.work
115 parttime
               13 present Prairie
               15 absent
                               BC
119 fulltime
131 parttime
               19 present Ontario
185 not.work
               13 absent Ontario
               15 present
                               BC
186 parttime
                               BC
190 not.work
               23 present
199 fulltime
               10
                   absent
                           Quebec
                          Quebec
230 parttime
               23 present
231 not.work
               7 present Quebec
                            Quebec
233 fulltime
               15
                    absent
                            Quebec
252 not.work
               23
                    absent
```

The sample function is used to pick a random sample of 20 rows in the data frame, selecting 20 random numbers without replacement from one to the number of rows in Womenlf; the numbers are placed in ascending order by the sort function.¹⁵

We use the set.seed function to specify the seed for R's pseudo-random number generator, ensuring that if we repeat the sample command, we will obtain the same sequence of pseudo-random numbers. Otherwise, the seed of the random-number generator will be selected unpredictably based on the system clock when the first random number is generated in an R session. Setting the random seed to a known value before a random simulation makes the result of the simulation reproducible. In serious work, we generally prefer to start with a known but randomly selected seed, as follows:

```
> (seed <- sample(2^31 - 1, 1))
[1] 974373618
> set.seed(seed)
```

The number $2^{31} - 1$ is the largest integer representable as a 32-bit binary number on most of the computer systems on which R runs (see Section 2.6.2).

The data in Women1f originate from a social survey of the Canadian population conducted in 1977 and pertain to married women between the ages of 21 and 30, with the variables defined as follows:

• partic: Labor-force participation, parttime, fulltime, or not.work (not working outside the home).

¹⁵ If the objective here were simply to sample 20 rows from Women1f, then we could more simply use the some function in the car package, some (Women1f, 20), but we will reuse this sample to check on the results of our recodes.

- hincome: Husband's income, in \$1,000s (actually, family income minus wife's income).
- children: Presence of children in the household: present or absent.
- region: Atlantic, Quebec, Ontario, Prairie (the Prairie provinces), or BC (British Columbia).

Now consider the following recodes:

```
> # recode in two ways:
> Womenlf$working <- recode(Womenlf$partic,
     ' c("parttime", "fulltime")="yes"; "not.work"="no" ')
> Womenlf$working.alt <- recode(Womenlf$partic,
     ' c("parttime", "fulltime")="yes"; else="no" ')
> Womenlf$working[sample.20] # 20 sampled observations
 [1] no no no yes no yes no yes no yes no yes no
[16] yes yes no yes no
Levels: no yes
> with(Womenlf, all(working == working.alt)) # check
[1] TRUE
> Womenlf$fulltime <- recode(Womenlf$partic,
+ ' "fulltime"="yes"; "parttime"="no"; "not.work"=NA ')
> Womenlf$fulltime[sample.20] # 20 sampled observations
[1] <NA> <NA> no <NA> yes <NA> no yes no
[13] <NA> no <NA> yes no <NA> yes <NA>
Levels: no yes
> Womenlf$region.4 <- recode(Womenlf$region,
+ 'c("Prairie", "BC")="West" ')
> Womenlf$region.4[sample.20] # 20 sampled observations
 [1] Ontario Ontario Atlantic Ontario Ontario Ontario
 [7] Ontario Ontario Ontario West West Ontario
[13] Ontario West West Quebec Quebec Quebec
[19] Quebec Quebec
Levels: Atlantic Ontario Quebec West
```

In all these examples, factors (either partic or region) are recoded, and consequently, recode returns factors as results:

- The first two examples yield identical results, with the second example illustrating the use of else. To verify that all the values in working and working alt are the same, we use the all function along with the element-wise comparison operator == (equals).
- In the third example, the factor fulltime is created, indicating whether
 a woman who works outside the home works full-time or part-time;
 fulltime is NA (missing) for women who do not work outside the
 home.
- The fourth and final example illustrates how values that are *not* recoded (here Atlantic, Quebec, and Ontario in the factor region) are simply carried over to the result.

The standard R ifelse command (discussed further in Section 8.3.1) can also be used to recode data. For example,