

# R Data Import/Export

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Version 1.2.2 (2001-02-26)

R Development Core Team

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# Table of Contents

<b>Acknowledgements</b> .....	<b>1</b>
<b>1 Introduction</b> .....	<b>2</b>
1.1 Imports .....	2
1.2 Export to text files .....	3
1.3 XML .....	4
<b>2 Spreadsheet-like data</b> .....	<b>5</b>
2.1 Variations on <code>read.table</code> .....	5
2.2 Fixed-width-format files .....	6
2.3 Using <code>scan</code> directly .....	6
2.4 Re-shaping data .....	7
2.5 Flat contingency tables .....	8
<b>3 Importing from other statistical systems</b> ....	<b>9</b>
3.1 Minitab, SAS, SPSS, Stata .....	9
3.2 S-PLUS .....	9
3.3 Octave .....	9
<b>4 Relational databases</b> .....	<b>10</b>
4.1 Why use a database? .....	10
4.2 Overview of RDBMSs .....	10
4.2.1 SQL queries .....	11
4.2.2 Data types .....	12
4.3 R interface packages .....	12
4.3.1 Package <code>RPgSQL</code> .....	12
4.3.2 Package <code>RODBC</code> .....	14
4.3.3 Package <code>RMySQL</code> .....	16
4.3.4 Package <code>RmSQL</code> .....	17
<b>5 Binary files</b> .....	<b>18</b>
5.1 Package <code>Rstreams</code> .....	18
5.2 Binary data formats .....	19
<b>6 Connections</b> .....	<b>20</b>
6.1 Types of connections .....	20
6.2 Output to connections .....	21
6.3 Input from connections .....	21
6.3.1 Pushback .....	22
6.4 Listing and manipulating connections .....	22
6.5 Binary connections .....	23

<b>7</b>	<b>Network interfaces .....</b>	<b>24</b>
7.1	Reading from sockets .....	24
7.2	Using <code>download.file</code> .....	24
7.3	DCOM interface .....	24
7.4	CORBA interface .....	24
<b>Appendix A</b>	<b>References .....</b>	<b>26</b>
	<b>Function and variable index .....</b>	<b>27</b>
	<b>Concept index .....</b>	<b>29</b>

## Acknowledgements

The relational databases part of this manual is based in part on an earlier manual by Douglas Bates and Saikat DebRoy. The principal author of this manual was Brian Ripley.

Many volunteers have contributed to the packages used here. The principal authors of the packages mentioned are

<b>CORBA</b>	Duncan Temple Lang
<b>e1071</b>	Friedrich Leisch
<b>foreign</b>	Thomas Lumley, Saikat DebRoy and Douglas Bates
<b>hdf5</b>	Marcus Daniels
<b>Java</b>	John Chambers and Duncan Temple Lang
<b>netCDF</b>	Thomas Lumley
<b>RmSQL</b>	Torsten Hothorn
<b>RMySQL</b>	David James and Saikat DebRoy
<b>RODBC</b>	Michael Lapsley
<b>RSPerl</b>	Duncan Temple Lang
<b>RPgSQL</b>	Timothy Keitt
<b>RSPython</b>	Duncan Temple Lang
<b>Rstreams</b>	Brian Ripley and Duncan Murdoch

Brian Ripley is the author of the support for connections.

# 1 Introduction

Reading data into a statistical system for analysis and exporting the results to some other system for report writing can be frustrating tasks that can take far more time than the statistical analysis itself, even though most readers will find the latter far more appealing.

This manual describes the import and export facilities available either in R itself or via packages which are available from CRAN. Some of the packages described are still under development (and hence in the Devel area on CRAN) but they already provide useful functionality.

Unless otherwise stated, everything described in this manual is available for both Unix/Linux and Windows versions of R.

In general, statistical systems like R are not particularly well suited to manipulations of large-scale data. Some other systems are better than R at this, and part of the thrust of this manual is to suggest that rather than duplicating functionality in R we can make the other system do the work! (For example Therneau & Grambsch (2000) comment that they prefer to do data manipulation in SAS and then use **survival5** in S for the analysis.) Several recent packages allow functionality developed in languages such as **Java**, **perl** and **python** to be directly integrated with R code, making the use of facilities in these languages even more appropriate. (See the **Java**, **RSPerl** and **RSPython** packages.)

It is also worth remembering that R like S comes from the Unix tradition of small reusable tools, and it can be rewarding to use tools such as **awk** and **perl** to manipulate data before import or after export. The case study in Becker, Chambers & Wilks (1988, Chapter 9) is an example of this, where Unix tools were used to check and manipulate the data before input to S. R itself takes that approach, using **perl** to manipulate its databases of help files rather than R itself, and the function **read.fwf** used a call to a **perl** script until it was decided not to require **perl** at run-time. The traditional Unix tools are now much more widely available, including on Windows.

## 1.1 Imports

The easiest form of data to import into R is a simple text file, and this will often be acceptable for problems of small or medium scale. The primary function to import from a text file is **scan**, and this underlies most of the more convenient functions discussed in [Chapter 2 \[Spreadsheet-like data\]](#), [page 5](#).

However, all statistical consultants are familiar with being presented by a client with a floppy disc or CD-R of data in some proprietary binary format, for example ‘an Excel spreadsheet’ or ‘an SPSS file’. Often the simplest thing to do is to use the originating application to export the data as a text file (and statistical consultants will have copies of the commonest applications on their computers for that purpose). However, this is not always possible, and [Chapter 3 \[Importing from other statistical systems\]](#), [page 9](#) discusses what facilities are available to access such files directly from R.

In a few cases, data have been stored in a binary form for compactness and speed of access. One application of this that we have seen several times is imaging data, which is normally stored as a stream of bytes as represented in memory, possibly preceded by a header. Such data formats are discussed in [Chapter 5 \[Binary files\]](#), [page 18](#) and [Section 6.5 \[Binary connections\]](#), [page 23](#).

For much larger databases it is common to handle the data using a database management system (DBMS). There is once again the option of using the DBMS to extract a plain file, but for many such DBMSs the extraction operation can be done directly from an R package: See [Chapter 4 \[Relational databases\]](#), page 10. Importing data via network connections is discussed in [Chapter 7 \[Network interfaces\]](#), page 24.

## 1.2 Export to text files

Exporting results from R is usually a less contentious task, but there are still a number of pitfalls. There will be a target application in mind, and normally a text file will be the most convenient interchange vehicle. (If a binary file is required, see [Chapter 5 \[Binary files\]](#), page 18.)

Function `cat` underlies the functions for exporting data. It takes a `file` argument, and the `append` argument allows a text file to be written via successive calls to `cat`.

The commonest task is to write a matrix or data frame to file as a rectangular grid of numbers, possibly with row and column labels. This can be done by the functions `write.table` and `write`. Function `write` just writes out a matrix or vector in a specified number of columns (and transposes a matrix). Function `write.table` is more convenient, and writes out a data frame (or an object that can be coerced to a data frame) with row and column labels.

There are a number of issues that need to be considered in writing out a data frame to a text file.

### 1. Precision

These functions are based on `cat` not `print`, and the precision to which numbers are printed is governed by the current setting of `options(digits)`. It may be necessary to increase this to avoid losing precision. For more control, use `format` on a data frame, possibly column-by-column.

### 2. Header line

R prefers the header line to have no entry for the row names, so the file looks like

	dist	climb	time
Greenmantle	2.5	650	16.083
...			

Some other systems require a (possibly empty) entry for the row names, which is what `write.table` will provide if argument `col.names = NA` is specified. Excel is one such system.

### 3. Separator

A common field separator to use in the file is a comma, as that is unlikely to appear in any of the fields, in English-speaking countries. Such files are known as CSV (comma separated values) files. In some locales the comma is used as the decimal point (set this in `write.table` by `dec = ","`) and there CSV files use the semicolon as the field separator.

Using a semicolon or tab (`sep = "\t"`) are probably the safest options.

#### 4. Missing values

By default missing values are output as NA, but this may be changed by argument `na`. Note that NaNs are treated as NA by `write.table`, but not by `cat` nor `write`.

#### 5. Quoting strings

By default strings are quoted (including the row and column names). Argument `quote` controls quoting of character and factor variables.

Some care is needed if the strings contain embedded quotes. Three useful forms are

```
> df <- data.frame(a = I("a \" quote"))
> write.table(df)
"a"
"1" "a \" quote"
> write.table(df, qmethod = "double")
"a"
"1" "a "" quote"
> write.table(df, quote = FALSE, sep = ",")
a
1,a " quote
```

The second is the form of escape commonly used by spreadsheets.

It is possible to use `sink` to divert the standard R output to a file, and thereby capture the output of (possibly implicit) `print` statements. This is not usually the most efficient route, and the `options(width)` setting may need to be increased.

## 1.3 XML

When reading data from text files, it is the responsibility of the user to know and to specify the conventions used to create that file, e.g. the comment character, whether a header line is present, the value separator, the representation for missing values (and so on) described in [Section 1.2 \[Export to text files\]](#), [page 3](#). A markup language which can be used to describe not only content but also the structure of the content can make a file self-describing, so that one need not provide these details to the software reading the data.

The eXtensible Markup Language – more commonly know simply as XML – can be used to provide such structure, not only for standard datasets but also more complex data structures. XML is becoming extremely popular and is emerging as a standard for general data markup and exchange. It is being used by different communities to describe geographical data such as maps, graphical displays, mathematics and so on.

The **XML** package provides general facilities for reading and writing XML documents within both R and S-PLUS in the hope that we can easily make use of this technology as it evolves. Several people are exploring how we can use XML for, amongst other things, representing datasets to be shared across different applications; storing R and S-PLUS objects so they can be shared by both systems; representing plots via SVG (Scalable Vector Graphics, a dialect of XML); representing function documentation; generating “live” analyses/reports that contain text, data and code.

A description of the facilities of the **XML** package is outside the scope of this document: see the package’s Web page at <http://www.omegahat.org/RXML> for details and examples.



## 2 Spreadsheet-like data

In [Section 1.2 \[Export to text files\]](#), page 3 we saw a number of variations on the format of a spreadsheet-like text file, in which the data are presented in a rectangular grid, possibly with row and column labels. In this section we consider importing such files into R.

### 2.1 Variations on `read.table`

The function `read.table` is the most convenient way to read in a rectangular grid of data. Because of the many possibilities, there are several other functions that call `read.table` but change a group of default arguments.

Some of the issues to consider are:

#### 1. Header line

We recommend that you specify the `header` argument explicitly. Conventionally the header line has entries only for the columns and not for the row labels, so is one field shorter than the remaining lines. (If R sees this, it sets `header = TRUE`.) If presented with a file that has a (possibly empty) header field for the row labels, read it in by something like

```
read.table("file.dat", header = TRUE, row.names = 1)
```

Column names can be given explicitly via the `col.names`; explicit names override the header line (if present).

#### 2. Separator

Normally looking at the file will determine the field separator to be used, but with white-space separated files there may be a choice between `sep = ""` which uses any white space (spaces, tabs or newlines) as a separator, `sep = " "` and `sep = "\t"`. Note that the choice of separator affects the input of quoted strings.

#### 3. Quoting

By default character strings can be quoted by either `'` or `"`, and in each case all the characters up to a matching quote are taken as part of the character string. The set of valid quoting characters (which might be none) is controlled by the `quote` argument. For `sep = "\n"` the default is changed to `quote = ""`.

If no separator character is specified, quotes can be escaped within quoted strings by immediately preceding them by `\`, C-style.

If a separator character is specified, quotes can be escaped within quoted strings by doubling them as is conventional in spreadsheets. For example

```
'One string isn't two',"one more"
```

can be read by

```
read.table("testfile", sep = ",")
```

This does not work with the default separator.

#### 4. Missing values

By default the file is assumed to contain the character string `NA` to represent missing values, but this can be changed by the argument `na.strings`, which is a vector of one or more character representations of missing values.

Empty fields in numeric columns are also regarded as missing values.

### 5. Unfilled lines

It is quite common for a file exported from a spreadsheet to have all trailing empty fields (and their separators) omitted. To read such files set `fill = TRUE`.

### 6. White space in character fields

If a separator is specified, leading and trailing white space in character fields is regarded as part of the field. To strip the space, use argument `strip.white = TRUE`.

### 7. Blank lines

By default, `read.table` ignores empty lines. This can be changed by setting `blank.lines.skip = FALSE`, which will only be useful in conjunction with `fill = TRUE`, perhaps to indicate missing data in a regular layout.

Convenience functions `read.csv` and `read.delim` provide arguments to `read.table` appropriate for CSV and tab-delimited files exported from spreadsheets in English-speaking locales. The variations `read.csv2` and `read.delim2` are appropriate for use in countries where the comma is used for the decimal point.

If the options to `read.table` are specified incorrectly, the error message will usually be of the form

```
row.lens=
[1] 4 4 5 5 5 4 6 4 4 4 5 4 4 5 5
Error in read.table("file.dat", header = TRUE) :
  all rows must have the same length.
```

This may give enough information to find the problem, but the auxiliary function `count.fields` can be useful to investigate further.

## 2.2 Fixed-width-format files

Sometimes data files have no field delimiters but have fields in pre-specified columns. This was very common in the days of punched cards, and is still sometimes used to save file space.

Function `read.fwf` provides a simple way to read such files, specifying a vector of field widths. The function reads the file into memory as whole lines, splits the resulting character strings, writes out a temporary tab-separated file and then calls `read.table`. This is adequate for small files, but for anything more complicated we recommend using the facilities of a language like `perl` to pre-process the file.

## 2.3 Using scan directly

Both `read.table` and `read.fwf` use `scan` to read the file, and then process the results of `scan`. They are very convenient, but sometimes it is better to use `scan` directly.

One reason can be to reduce memory usage. Function `read.table` reads the whole data file into memory and processes it into a data frame, thereby creating another copy of the data. If the post-processing is not required (for example if most of the variables are to be discarded) it may be feasible to use `scan` but not to use `read.table`.

Function `scan` has many arguments, most of which we have already covered under `read.table`. The most crucial argument is `what`, which specifies a list of modes of variables

to be read from the file. If the list is named, the names are used for the components of the returned list. Modes can be numeric, character or complex, and are usually specified by an example, e.g. 0, "" or 0i. For example

```
cat("2 3 5 7", "11 13 17 19", file="ex.dat", sep="\n")
scan(file="ex.dat", what=list(x=0, y="", z=0), flush=TRUE)
```

returns a list with three components and discards the fourth column in the file.

There is a function `readLines` which will be more convenient if all you want is to read whole lines into R for further processing.

## 2.4 Re-shaping data

Sometimes spreadsheet data is in a compact format that gives the covariates for each subject followed by all the observations on that subject. R's modelling functions need observations in a single column. Consider the following sample of data from repeated MRI brain measurements

Status	Age	V1	V2	V3	V4
P	23646	45190	50333	55166	56271
CC	26174	35535	38227	37911	41184
CC	27723	25691	25712	26144	26398
CC	27193	30949	29693	29754	30772
CC	24370	50542	51966	54341	54273
CC	28359	58591	58803	59435	61292
CC	25136	45801	45389	47197	47126

There are two covariates and up to four measurements on each subject. The data were exported from Excel as a file 'mr.csv'.

We can use `stack` to help manipulate these data to give a single response.

```
zz <- read.csv("mr.csv", strip.white = TRUE)
zzz <- cbind(zz[gl(nrow(zz), 1, 4*nrow(zz)), 1:2], stack(zz[, 3:6]))
```

with result

	Status	Age	values	ind
X1	P	23646	45190	V1
X2	CC	26174	35535	V1
X3	CC	27723	25691	V1
X4	CC	27193	30949	V1
X5	CC	24370	50542	V1
X6	CC	28359	58591	V1
X7	CC	25136	45801	V1
X11	P	23646	50333	V2
...				

Function `unstack` goes in the opposite direction, and may be useful for exporting data.

Another way to do this is to use the (experimental) function `reshapeLong`, by

```
> reshapeLong(zz, V1:V4)
  Status Age reshape.i reshape.j reshape.v
1      P 23646         1        V1    45190
2      P 23646         1        V2    50333
```

3	P	23646	1	V3	55166
4	P	23646	1	V4	56271
5	CC	26174	2	V1	35535
6	CC	26174	2	V2	38227
7	CC	26174	2	V3	37911
8	CC	26174	2	V4	41184
...					

## 2.5 Flat contingency tables

Displaying higher-dimensional contingency tables in array form typically is rather inconvenient. In categorical data analysis, such information is often represented in the form of bordered two-dimensional arrays with leading rows and columns specifying the combination of factor levels corresponding to the cell counts. These rows and columns are typically “ragged” in the sense that labels are only displayed when they change, with the obvious convention that rows are read from top to bottom and columns are read from left to right. In R, such “flat” contingency tables can be created using `fTable`, which creates objects of class “`fTable`” with an appropriate print method.

As a simple example, consider the R standard data set `UCBAdmissions` which is a 3-dimensional contingency table resulting from classifying applicants to graduate school at UC Berkeley for the six largest departments in 1973 classified by admission and sex.

```
> data(UCBAdmissions)
> fTable(UCBAdmissions)
```

		Dept	A	B	C	D	E	F
Admit	Gender							
Admitted	Male		512	353	120	138	53	22
	Female		89	17	202	131	94	24
Rejected	Male		313	207	205	279	138	351
	Female		19	8	391	244	299	317

The printed representation is clearly more useful than displaying the data as a 3-dimensional array.

There is also a function `read.fTable` for reading in flat-like contingency tables from files. This has additional arguments for dealing with variants on how exactly the information on row and column variables names and levels is represented. The help page for `read.fTable` has some useful examples. The flat tables can be converted to standard contingency tables in array form using `as.table`.

Note that flat tables are characterized by their “ragged” display of row (and maybe also column) labels. If the full grid of levels of the row variables is given, one should instead use `read.table` to read in the data, and create the contingency table from this using `xtabs`.

## 3 Importing from other statistical systems

In this chapter we consider the problem of reading a binary data file written by another statistical system. This is often best avoided, but may be unavoidable if the originating system is not available.

### 3.1 Minitab, SAS, SPSS, Stata

Package **foreign** on CRAN provides import facilities for files produced by these four statistical systems, and for export to Stata.

Stata `.dta` files are a binary file format. Files from versions 5.0 and 6.0 of Stata can be read and written by functions `read.dta` and `write.dta`.

Function `read.mtp` imports a ‘Minitab Portable Worksheet’. This returns the components of the worksheet as an R list.

Function `read.xport` reads a file in SAS Transport (XPORT) format and return a list of data frames.

Function `read.spss` can read files created by the ‘save’ and ‘export’ commands in SPSS. It returns a list with one component for each variable in the saved data set.

### 3.2 S-PLUS

Package **Rstreams** on CRAN contains a function `readSfile` which can read binary objects produced by S-PLUS 3.x, 4.x or 2000 on Unix or Windows (and can read them on a different OS). This is able to read many but not all S objects: in particular it can read vectors, matrices and data frames and lists containing those.

Package **Rstreams** also contains a function `data.restore` to read S-PLUS data dumps (created by `data.dump`) with the same restrictions. It may be possible to read data dumps from S-PLUS 5.x and 6.x written with `data.dump(oldStyle=T)`.

### 3.3 Octave

Octave is a numerical linear algebra system, and function `read.octave` in package **e1071** can read the first vector or matrix from an Octave ASCII data file created using the Octave command `save -ascii`.

## 4 Relational databases

### 4.1 Why use a database?

There are limitations on the types of data that R handles well. Since all data being manipulated by R are resident in memory, and several copies of the data can be created during execution of a function, R is not well suited to extremely large data sets. Data objects that are more than a few (tens of) megabytes in size can cause R to run out of memory.

R does not easily support concurrent access to data. That is, if more than one user is accessing, and perhaps updating, the same data, the changes made by one user will not be visible to the others.

R does support persistence of data, in that you can save a data object or an entire worksheet from one session and restore it at the subsequent session, but the format of the stored data is specific to R and not easily manipulated by other systems.

Database management systems (DBMSs) and, in particular, relational DBMSs (RDBMSs) *are* designed to do all of these things well. Their strengths are

1. To provide fast access to selected parts of large databases.
2. Powerful ways to summarize and cross-tabulate columns in databases.
3. Store data in more organized ways than the rectangular grid model of spreadsheets and R data frames.
4. Concurrent access from multiple clients running on multiple hosts while enforcing security constraints on access to the data.
5. Ability to act as a server to a wide range of clients.

The sort of statistical applications for which DBMS might be used are to extract a 10% sample of the data, to cross-tabulate data to produce a multi-dimensional contingency table, and to extract data group by group from a database for separate analysis.

### 4.2 Overview of RDBMSs

Traditionally there have been large (and expensive) commercial RDBMSs (**Informix**; **Oracle**; **Sysbase**; IBM's DB/2; Microsoft SQL Server on Windows) and academic and small-system databases (such as MySQL, PostgreSQL, Microsoft Access, . . .), the former marked out by much greater emphasis on data security features. The line is blurring, with the Open Source PostgreSQL having more and more high-end features, and 'free' versions of Informix, Oracle and Sysbase being made available on Linux.

There are other commonly used data sources, including spreadsheets, non-relational databases and even text files (possibly compressed). Open Database Connectivity (ODBC) is a standard to use all of these data sources. It originated on Windows (see <http://www.microsoft.com/data/odbc/>) but is also implemented on Linux.

All of the packages described later in this chapter provide clients to client/server databases. The database can reside on the same machine or (more often) remotely. There is an ISO standard (in fact several: SQL-92 is ISO/IEC 9075, also known as ANSI

X3.135-1992) for an interface language called SQL (Structured Query Language, sometimes pronounced ‘sequel’: see Bowman *et al.* 1996) which these DBMSs support to varying degrees.

### 4.2.1 SQL queries

The more comprehensive R interfaces generate SQL behind the scenes for common operations, but direct use of SQL is needed for complex operations in all. Conventionally SQL is written in upper case, but many users will find it more convenient to use lower case in the R interface functions.

A relational DBMS stores data as a database of *tables* (or *relations*) which are rather similar to R data frames, in that they are made up of *columns* or *fields* of one type (numeric, character, date, currency, . . .) and *rows* or *records* containing the observations for one entity.

SQL ‘queries’ are quite general operations on a relational database. The classical query is a SELECT statement of the type

```
SELECT State, Murder FROM USArrests WHERE rape > 30 ORDER BY Murder
```

```
SELECT t.sch, c.meanses, t.sex, t.achieve
FROM student as t, school as c WHERE t.sch = c.id
```

```
SELECT sex, COUNT(*) FROM student GROUP BY sex
```

```
SELECT sch, AVG(sestat) FROM student GROUP BY sch LIMIT 10
```

The first of these selects two columns from the R data frame **USArrests** that has been copied across to a database table, subsets on a third column and asks the results be sorted. The second performs a database *join* on two tables **student** and **school** and returns four columns. The third and fourth queries do some cross-tabulation and return counts or averages. (The five aggregation functions are COUNT(\*), SUM, MAX, MIN and AVG, each applied to a single column.)

SELECT queries use FROM to select the table, WHERE to specify a condition for inclusion (or more than one condition separated by AND or OR), and ORDER BY to sort the result. Unlike data frames, rows in RDBMS tables are best thought of as unordered, and without an ORDER BY statement the ordering is indeterminate. You can sort (in lexicographical order) on more than one column by separating them by commas. Placing DESC after an ORDER BY puts the sort in descending order.

SELECT DISTINCT queries will only return one copy of each distinct row in the selected table.

The GROUP BY clause selects subgroups of the rows according to the criterion. If more than one column is specified (separated by commas) then multi-way cross-classifications can be summarized by one of the five aggregation functions. A HAVING clause allows the select to include or exclude groups depending on the aggregated value.

If the SELECT statement contains an ORDER BY statement that produces a unique ordering, a LIMIT clause can be added to select (by number) a contiguous block of output rows. This can be useful to retrieve rows a block at a time. (It may not be reliable unless the ordering is unique, as the LIMIT clause can be used to optimize the query.)

There are queries to create a table (`CREATE TABLE`, but usually one copies a data frame to the database in these interfaces), `INSERT` or `DELETE` or `UPDATE` data. A table is destroyed by a `DROP TABLE` ‘query’.

## 4.2.2 Data types

Data can be stored in a database in various data types. The range of data types is DBMS-specific, but the SQL standard defines many types, including the following that are widely implemented (often not by the SQL name).

`float(p)` Real number, with optional precision. Often called `real` or `double` or `double precision`.

`integer` 32-bit integer. Often called `int`.

`smallint` 16-bit integer

`character(n)`  
fixed-length character string. Often called `char`.

`character varying(n)`  
variable-length character string. Often called `varchar`.

`boolean` true or false. Sometimes called `bool`.

`date` calendar date

`time` time of day

`timestamp`  
date and time

There are variants on `time` and `timestamp`, with `timezone`.

The more comprehensive of the R interface packages hide the type conversion issues from the user.

## 4.3 R interface packages

There are four packages available on CRAN to help R communicate with DBMSs. They provide different levels of abstraction. Some provide means to copy whole data frames to and from databases. All have functions to select data within the database via SQL queries, and (except **RmSQL**) to retrieve the result as a whole as a data frame or in pieces (usually as groups of rows, but **RPgSQL** can retrieve columns). All except **RODBC** are (currently) tied to one DBMS.

### 4.3.1 Package RPgSQL

Package **RPgSQL** at <http://rpgsql.sourceforge.net/> and on CRAN provides an interface to **PostgreSQL**.

PostgreSQL is described by its developers as ‘the most advanced open source database server’ (Momjian, 2000). It would appear to be buildable for most Unix-alike OSes and Windows (under Cygwin or U/Win). PostgreSQL has most of the features of the commercial RDBMSs.



**RPgSQL** is the most mature and comprehensive of these RDBMS interfaces.

To make use of **RPgSQL**, first open a connection to a database using `db.connect`. (Currently only one connection can be open at a time.) Once a connection is open an R data frame can be copied to a PostgreSQL table by `db.write.table`, whereas `db.read.table` copies a PostgreSQL table to an R data frame.

**RPgSQL** has the interesting concept of a *proxy data frame*. A data frame proxy is an R object that inherits from the `"data.frame"` class, but contains no data. All accesses to the proxy data frame generate the appropriate SQL query and retrieve the resulting data from the database. A proxy data frame is set up by a call to `bind.db.proxy`. To remove the proxy, just remove the object which `bind.db.proxy` created.

A finer level of control is available via sending SQL queries to the PostgreSQL server via `db.execute`. This leaves a result in PostgreSQL's result cache, unless flushed by `clear = TRUE` (the default). Once a result is in the cache, `db.fetch.result` can be used to fetch the whole result as a data frame. Functions such as `db.result.columns` and `db.result.rows` will report the number of columns and rows in the cached table, and `db.read.column` will fetch a single column (as a vector). An individual cell in the result can be read by `db.result.get.value`. `db.clear.result` will clear the result cache.

One disadvantage is that PostgreSQL maps all table and column names to lower case, so for maximal flexibility, only use lower case in R names. Functions `sql.insert` and `sql.select` provide convenience wrappers for the INSERT and SELECT queries.

We can explore these functions in a simple example. The database 'testdb' had already been set up, and as PostgreSQL was running on a standalone machine no further authentication was required to connect.

```
> library(RPgSQL)
> db.connect(dbname="testdb")    # add authentication as needed
Connected to database "testdb" on ""
> data(USArrests)
> usarrests <- USArrests
> names(usarrests) <- tolower(names(USArrests))
> db.write.table(USArrests, write.row.names = TRUE)
> db.write.table(usarrests, write.row.names = TRUE)
> rm(USArrests, usarrests)
## db.ls lists tables in the database.
> db.ls()
[1] "USArrests" "usarrests"
> db.read.table("USArrests")
              Murder Assault UrbanPop Rape
Alabama         13.2      236      58 21.2
Alaska          10.0      263      48 44.5
...
## set up a proxy data frame. Remember USArrests has been removed
> bind.db.proxy("USArrests")
## USArrests is now a proxy, so all accesses are to the database
> USArrests[, "Rape"]
      Rape
1  21.2
2  44.5
```

```

...
> rm(USArrests) # remove proxy
> db.execute("SELECT rpgsql_row_names, murder FROM usarrests",
             "WHERE rape > 30 ORDER BY murder", clear=FALSE)
> db.fetch.result()
      murder
Colorado    7.9
Arizona     8.1
California   9.0
Alaska      10.0
New Mexico  11.4
Michigan     12.1
Nevada       12.2
Florida     15.4
> db.rm("USArrests", "usarrests") # use ask=FALSE to skip confirmation
Destroy table USArrests? y
Destroy table usarrests? y
> db.ls()
character(0)
> db.disconnect()

```

Notice how the row names are mapped if `write.row.names = TRUE` to a field `rpgsql_row_names` in the database table and transparently restored provided we preserve that field in the query.

**RPgSQL** provides means to extend its mapping between R classes within a data frame and PostgreSQL types.

### 4.3.2 Package RODBC

Package **RODBC** on CRAN provides an interface to database sources supporting an ODBC interface. This is very widely available, and allows the same R code to access different database systems. **RODBC** runs on both Linux and Windows, and many database systems provide support ODBC, including most of those on Windows (such as Microsoft Access), and MySQL, Oracle and PostgreSQL on Unix/Linux.

You will need an ODBC Driver Manager such as unixODBC (<http://www.unixODBC.org>) or iODBC (<http://www.iODBC.org>) on Unix/Linux, and an installed driver for your database system. The FreeODBC project (<http://www.jepstone.net/FreeODBC/>) is a repository of information related to ODBC.

Two groups of interface functions are provided. The `odbc*` group provide a low-level interface to the basic ODBC functions: see the help page (`?RODBC`) for details. The `sql*` group provide an interface between R data frames and SQL tables.

Up to 16 simultaneous connections are possible. A connection is opened by a call to `odbcConnect` which returns a handle used for subsequent access to the database. A connection is closed by `odbcClose`. Details of the tables on a connection can be found using `sqlTables`.

Function `sqlSave` copies an R data frame to a table in the database, and `sqlFetch` copies a table in the database to an R data frame.

An SQL query can be sent to the database by a call to `sqlQuery`. This returns the result in an R data frame. (`sqlCopy` sends a query to the database and saves the result as a table in the database.) A finer level of control is attained by first calling `odbcQuery` and then `sqlGetResults` to fetch the results. The latter can be used within a loop to retrieve a limited number of rows at a time. `sqlGetResults` by default returns a data frame, but the raw results can also be obtained as a character matrix.

Here is an example using PostgreSQL, for which the ODBC driver maps column and data frame names to lower case. We use a database `testdb` we created earlier, and had the DSN (data source name) set up in ‘`~/odbc.ini`’ under `unixODBC`. Exactly the same code worked using `MyODBC` to access a MySQL database under Linux or Windows NT (where MySQL also maps names to lowercase). Under Windows, DSNs are set up in the ODBC applet in the Control Panel.

```
> library(RODBC)
## tell it to map names to l/case
> channel <- odbcConnect("testdb", uid="ripley", case="tolower")
## load a data frame into the database
> data(USArrests)
> sqlSave(channel, USArrests, rownames="state")
> rm(USArrests)
## list the tables in the database
> sqlTables(channel)
  TABLE_QUALIFIER TABLE_OWNER TABLE_NAME TABLE_TYPE REMARKS
1                NA           NA  usarrests      TABLE      NA
## list it
> sqlFetch(channel, "USArrests", rownames = TRUE)
               murder assault urbanpop rape
Alabama        13.2      236      58 21.2
Alaska         10.0      263      48 44.5
...
## an SQL query, originally on one line
> sqlQuery(channel, "select state, murder from USArrests
  where rape > 30 order by murder")
      state murder
1 Colorado    7.9
2 Arizona     8.1
3 California   9.0
4 Alaska     10.0
5 New Mexico  11.4
6 Michigan    12.1
7 Nevada      12.2
8 Florida     15.4
## remove the table
> sqlDrop(channel, "usarrests")
## close the connection
> odbcClose(channel)
```

As a simple example of using ODBC with a Excel spreadsheet, suppose that an DSN for spreadsheet ‘`bdr.xls`’ has been set up in the Control Panel. Then we can read from the spreadsheet by

```

> library(RODBC)
> channel <- odbcConnect("bdr.xls")
## list the spreadsheets
> sqlTables(channel)
  TABLE_CAT TABLE_SCHEM      TABLE_NAME  TABLE_TYPE REMARKS
1 C:\\bdr      NA          Sheet1$  SYSTEM TABLE      NA
2 C:\\bdr      NA          Sheet2$  SYSTEM TABLE      NA
3 C:\\bdr      NA          Sheet3$  SYSTEM TABLE      NA
4 C:\\bdr      NA Sheet1$Print_Area      TABLE      NA
## retrieve the contents of sheet 1
> sh1 <- sqlQuery(channel, "select * from [Sheet1$]")

```

Notice that the specification of the table is different from the name returned by `sqlTables`: this precludes the use of `sqlFetch`.

### 4.3.3 Package RMySQL

Package **RMySQL** in the Devel area on CRAN provides an interface to the MySQL database system (see <http://www.mysql.com> and Dubois, 2000.). This is part of a project to provide a common API for access from R to relational DBMSs using SQL. Currently this provides lower-level facilities than **RPgSQL** or **RODBC**.

MySQL exists on Unix/Linux and Windows: as from version 3.23 (Jan 2001) it is released under GPL. MySQL is a ‘light and lean’ database. (It preserves the case of names where the operating file system is case-sensitive, so not on Windows.) Package **RMySQL** has been used on both Linux and Windows.

A call to the function `MySQL` returns a database connection manager object, and then a call to `dbConnect` opens a database connection which can subsequently be closed by a call to the generic function `close`.

SQL queries can be sent by either `dbExec` or `dbExecStatement`. `dbExec` sends the query and retrieves the results as a list. (It will fail unless the results are from a SELECT-like statement.) `dbExecStatement` sends the query and returns an object of class "MySQLResultSet" which can be used to retrieve the results, and subsequently used to `close` the result.

Function `fetch` is used to retrieve some or all of the rows in the query result, as a list. The function `hasCompleted` indicates if all the rows have been fetched, and `getRowCount` returns the number of rows in the result.

There is no simple way to load a data frame into the database, so we have to work harder using direct SQL statements. The following will only work if the database is local: otherwise the file will need to be uploaded to the MySQL host.

```

> library(RMySQL)
## open a connection to a MySQL database
> con <- dbConnect(MySQL(), dbname = "test")
## list the tables in the database
> getTables(con)
## load a data frame into the database, deleting any existing copy
> data(USArrests)
## write the values to a file
> write.table(USArrests, "/tmp/arrests.dat", sep="\t",
              row.names = FALSE, col.names = FALSE)

```

```
## delete any existing table
> if(!is.na(match("arrests", getTables(con)[[1]])))
  dbExecStatement(con, "drop table arrests")
## character string here needs to be on one line
> dbExecStatement(con,
  "create table arrests (Murder float, Assault float,
  UrbanPop float, Rape float)")
> dbExecStatement(con,
  'load data infile "/tmp/arrests.dat" into table arrests')
> unlink("/tmp/arrests.dat")
## Select from the loaded database.
> as.data.frame(dbExec(con,
  "select * from arrests where Rape > 30 order by Murder"))
  Murder Assault UrbanPop Rape
1    7.9      204      78 38.7
2    8.1      294      80 31.0
3    9.0      276      91 40.6
4   10.0      263      48 44.5
5   11.4      285      70 32.1
6   12.1      255      74 35.1
7   12.2      252      81 46.0
8   15.4      335      80 31.9
> close(con)
```

#### 4.3.4 Package RmSQL

Package **RmSQL** on CRAN provides an interface to the Mini SQL database system (also known as mSQL, <http://www.hughes.com.au>, Yarger *et al.*, 1999). The package documentation describes mSQL as

Note that mSQL is NOT GPL licenced but free of charge for universities and noncommercial organisations.

**RmSQL** provides the most basic interface of those in this chapter, a wrapper to the C-API of mSQL with no additional functionality.

A database connection is opened by first selecting a host with `msqlConnect` and then a database by `msqlSelect`. The connection is closed by a call to `msqlClose`. Then an SQL query is sent by a call to `msqlQuery`, and the results stored by a call to `msqlStoreResult`. When a query is finished with, the result can be freed by `msqlFreeResult`.

Once the result of a query has been stored, the values can be retrieved row by row using `msqlFetchRow`. This fetches the rows in order unless the position is reset by a call to `msqlDataSeek`. A call to `msqlNumRows` gives the total number of rows in the result.

No example is given here as the basic interface makes any example lengthy, but there is one in the ‘Example’ directory of the package.

## 5 Binary files

### 5.1 Package Rstreams

Package **Rstreams** on CRAN provides a low-level interface to read from and write to binary files. It has been used to read in data from MRI experiments and Windows sound files, for example.

**Rstreams**' view of the file is as a stream of bytes. A file can be opened by the function `openstream` for either reading or writing. The return value is a number (in fact the file descriptor) that is used to reference the open file until it is closed by `closestream`.

Once a stream is open for reading, bytes can be transferred from it to an R object by one of the functions

```
readint(stream, n, size = 4, signed = TRUE, swapbytes = FALSE)
readfloat(stream, n, size = 8, swapbytes = FALSE)
readcomplex(stream, n, size = 8, swapbytes = FALSE)
readchar(stream, n = 1, len = NA, bufsize = 256)
```

These return an R object of an appropriate mode and storage mode. (Integers too large to be represented in storage mode "integer" will be read into a vector of storage mode "double".) Here `n` is the number of items to be read, but fewer will be read if there is insufficient data on the file.

The size of the data components on file need not be the same as that in the machine running R, and data written on a little-endian machine can be read on a big-endian machine or *vice versa* by setting `swapbytes = TRUE`.

Character data can be read by `readchar` either as fixed-length blocks of bytes (by specifying `len`) or as ASCII-zero-delimited strings. It is also possible to read lines of text separated by one of the standard end-of-line marks (LF, CRLF or CR) by

```
readlines(stream, n = 1, bufsize = 256, eol)
```

where by default the `eol` mark is detected automatically.

A file open for writing can be written to by any of

```
writeint(stream, data, size = 4, swapbytes = FALSE)
writefloat(stream, data, size = 8, swapbytes = FALSE)
writecomplex(stream, data, size = 8, swapbytes = FALSE)
writechar(stream, data, asciiz = FALSE)
```

where `data` is an R vector of an appropriate mode. There is no separator between character strings on the file unless `asciiz = TRUE` is specified.

Function `copystream` can copy a specified number of bytes from one open stream to another.

Function `truncate` truncates a stream at its current position, so all data after that point is lost.

## 5.2 Binary data formats

Packages **hdf5** (in the Devel area) and **netCDF** on CRAN provide experimental interfaces to NASA's HDF5 (Hierarchical Data Format, see <http://hdf.ncsa.uiuc.edu/HDF5/>) and to UCAR's netCDF data files (network Common Data Form, see <http://www.unidata.ucar.edu/packages/netcdf/>), respectively.

Both of these are systems to store scientific data in array-oriented way, including descriptions, labels, formats, units, . . . . HDF5 also allows *groups* of arrays, and the R interface writes maps lists to HDF5 groups, and can write numeric and character vectors and matrices.

The R interface can only read netCDF, not write it (yet).

## 6 Connections

*Connections* are used in R in the sense of Chambers (1998), a set of functions to replace the use of file names by a flexible interface to file-like objects. Connections were introduced to R in version 1.2.0, and the details are still experimental and subject to change. Their potential is not yet fully realised.

### 6.1 Types of connections

The most familiar type of connection will be a file, and file connections are created by function `file`. File connections can (if the OS will allow it for the particular file) be opened for reading or writing or appending, in text or binary mode. Currently (and unlike S), connections can only be opened for one of reading and writing/appending at a time.

Note that by default a connection is not opened when it is created. The rule is that a function using a connection should open a connection (needed) if the connection is not already open, and close a connection after use if it opened it. In brief, leave the connection in the state you found it in. There are generic functions `open` and `close` with methods to explicitly open and close connections.

Unix programmers are used to dealing with special files `stdin`, `stdout` and `stderr`. These exist as *terminal connections* in R. They may be normal files, but they might also refer to input from and output to a GUI console. (Even with the standard Unix R interface, `stdin` refers to the lines submitted from `readline` rather than a file.)

The three terminal connections are always open, and cannot be opened or closed. `stdout` and `stderr` are conventionally used for normal output and error messages respectively. They may normally go to the same place, but whereas normal output can be re-directed by a call to `sink`, error output is always sent to `stderr`. Note carefully the language used here: the connections cannot be re-directed, but output can be sent to other connections.

*Text connections* are another source of input. They allow R character vectors to be read as if the lines were being read from a text file. A text connection is created and opened by a call to `textConnection`, which copies the current contents of the character vector to an internal buffer at the time of creation.

Text connections can also be used to capture R output to a character vector. `textConnection` can be asked to create a new character object or append to an existing one, in both cases in the user's workspace. The connection is opened by the call to `textConnection`, and at all times the complete lines output to the connection are available in the R object. Closing the connection writes any remaining output to a final element of the character vector.

*Pipes* are a special form of file that connects to another process, and pipe connections are created by the function `pipe` (currently implemented on Unix and `Rterm` only). Opening a pipe connection for writing (it makes no sense to append to a pipe) runs an OS command, and connects its standard input to whatever R then writes to that connection. Conversely, opening a pipe connection for input runs an OS command and makes its standard output available for R input from that connection.

More types of connections are planned, including reading from and writing to sockets and URLs.



## 6.2 Output to connections

We have described functions `cat`, `write`, `write.table` and `sink` as writing to a file, possibly appending to a file if argument `append = TRUE`, and this is what they did prior to R version 1.2.0.

The current behaviour is equivalent, but what actually happens is that when the `file` argument is a character string, a file connection is opened (for writing or appending) and closed again at the end of the function call. If we want to repeatedly write to the same file, it is more efficient to explicitly declare and open the connection, and pass the connection object to each call to an output function. This also makes it possible to write to pipes, which was implemented earlier in a limited way via the syntax `file = "|cmd"` (which can still be used).

There is a function `writeLines` to write complete text lines to a connection.

Some simple examples are

```
zz <- file("ex.data", "w") # open an output file connection
cat("TITLE extra line", "2 3 5 7", "", "11 13 17",
    file = zz, sep = "\n")
cat("One more line\n", file = zz)
close(zz)

## convert decimal point to comma in output, using a pipe (Unix)
zz <- pipe(paste("sed s/\\. /, / >", "outfile"), "w")
cat(format(round(rnorm(100), 4)), sep = "\n", file = zz)
close(zz)
## now look at the output file:
file.show(outfile, delete.file = TRUE)

## capture R output: use examples from help(lm)
zz <- textConnection("ex.lm.out", "w")
sink(zz)
example(lm, prompt.echo = "> ")
sink()
close(zz)
## now 'ex.lm.out' contains the output for futher processing.
## Look at it by, e.g.,
cat(ex.lm.out, sep = "\n")
```

## 6.3 Input from connections

The basic functions to read from connections are `scan` and `readLines`. These take a character string argument and open a file connection for the duration of the function call, but explicitly opening a file connection allows a file to be read sequentially in different formats.

Other functions that call `scan` can also make use of connections. In particular `read.table` can use connections, but as it needs to read the data twice it is inefficient on a non-seekable connection. (It reads the datafile into an R character vector and operates on that via a text connection.)

Some simple examples are

```
## read in file created in last examples
readLines("ex.data")
unlink("ex.data")

## read listing of current directory (Unix)
readLines(pipe("ls -l"))

# remove trailing commas from an input file.
# Suppose we are given a file 'data' containing
450, 390, 467, 654, 30, 542, 334, 432, 421,
357, 497, 493, 550, 549, 467, 575, 578, 342,
446, 547, 534, 495, 979, 479
# Then read this by
scan(pipe("sed -e s/,,$// data"), sep=",")
```

### 6.3.1 Pushback

C programmers may be familiar with the `ungetc` function to push back a character onto a text input stream. R connections have the same idea in a more powerful way, in that an (essentially) arbitrary number of lines of text can be pushed back onto a connection via a call to `pushBack`.

Pushbacks operate as a stack, so a read request first uses each line from the most recently pushbacked text, then those from earlier pushbacks and finally reads from the connection itself. Once a pushbacked line is read completely, it is cleared. The number of pending lines pushed back can be found via a call to `pushBackLength`.

A simple example will show the idea.

```
> zz <- textConnection(LETTERS)
> readLines(zz, 2)
[1] "A" "B"
> scan(zz, "", 4)
Read 4 items
[1] "C" "D" "E" "F"
> pushBack(c("aa", "bb"), zz)
> scan(zz, "", 4)
Read 4 items
[1] "aa" "bb" "G" "H"
> close(zz)
```

Pushback is only available for connections opened for input in text mode.

## 6.4 Listing and manipulating connections

A summary of all the connections currently opened by the user can be found by `showConnections()`, and a summary of all connections, including closed and terminal connections, by `showConnections(all = TRUE)`

The generic function `seek` can be used to read and (on some connections) reset the current position for reading or writing. Unfortunately it depends on OS facilities which may be unreliable (e.g. with text files under Windows).

## 6.5 Binary connections

R version 1.2.1 adds functions `readBin` and `writeBin` to read and write from binary connections. A connection is opened in binary mode by appending "b" to the mode specification, that is using mode "rb" for reading, and mode "wb" or "ab" (where appropriate) for writing. The functions have arguments

```
readBin(con, what, n = 1, size = NA, endian = .Platform$endian)
writeBin(object, con, size = NA, endian = .Platform$endian)
```

In each case `con` is a connection which will be opened if necessary for the duration of the call, and if a character string is given it is assumed to specify a file name.

It is slightly simpler to describe writing, so we will do that first. `object` should be an atomic vector object, that is a vector of mode `numeric`, `integer`, `logical`, `character` or `complex`, without attributes. By default this is written to the file as a stream of bytes exactly as it is represented in memory.

`readBin` reads a stream of bytes from the file and interprets them as a vector of mode given by `what`. This can be either an object of the appropriate mode (e.g. `what=integer()`) or a character string describing the mode (one of the five given in the previous paragraph or "double" or "int"). `size` specifies the maximum number of vector elements to read from the connection: if fewer are available a shorter vector will be returned.

The remaining two arguments are used to write or read data for interchange with another program or another platform. By default binary data is transferred directly from memory to the connection or *vice versa*. This will not suffice if the file is to be transferred to a machine with a different architecture, but between almost all R platforms the only change needed is that of byte-order. Intel ix86-based machines, Compaq Alpha and Vaxen are *little-endian*, whereas Sun Sparc, mc680x0 series, IBM R6000, SGI and most others are *big-endian*. (Network byte-order (as used by XDR, eXternal Data Representation) is big-endian.) To transfer to or from other programs we may need to do more, for example to read 16-bit integers or write single-precision real numbers. This can be done using the `size` argument, which (usually) allows sizes 1, 2, 4, 8 for integers and logicals, and sizes 4, 8 and perhaps 12 or 16 for reals. Transferring at different sizes can lose precision, and should not be attempted for vectors containing NA's.

Character strings are read and written in C format, that is as a string of bytes terminated by a zero byte. Package **Rstreams** provides more flexibility, for files.

## 7 Network interfaces

Some limited facilities are available to exchange data at a lower level across network connections.

### 7.1 Reading from sockets

Base R comes with some facilities to communicate *via* BSD sockets on systems that support them (including the common Linux, Unix and Windows ports of R). One potential problem with using sockets is that these facilities are often blocked for security reasons or to force the use of Web caches, so these functions may be more useful on an intranet than externally.

The low-level interface is given by functions `make.socket`, `read.socket`, `write.socket` and `close.socket`. The function `httpclient` builds on these to read (directly) from an HTTP server.

### 7.2 Using `download.file`

There are functions `read.table.url` and `scan.url` that appear to work like `read.table` and `scan` but access a Web resource at a specified URL. Appearances are deceptive, however, as these functions (and `url.show` and `source.url`) first download the Web resource to a local file and then apply the standard functions. Various methods can be used to download the file, including the programs `wget` and `lynx` and the direct use of BSD sockets.

### 7.3 DCOM interface

DCOM is a Windows protocol for communicating between different programs, possibly on different machines. Thomas Baier's `StatConnector` program available from CRAN under Software->Other->Non-standard provides an interface to the proxy DLL which ships with the Windows version of R and makes an DCOM server. This can be used to pass simple objects (vectors and matrices) to and from R and to submit commands to R.

The program comes with a Visual Basic demonstration, and there is an Excel plug-in by Erich Neuwirth available in the same area on CRAN. This interface is in the other direction to most of those considered here in that it is another application (Excel, or written in Visual Basic) that is the client and R is the server.

### 7.4 CORBA interface

CORBA (Common Object Request Broker Architecture) is similar to DCOM, allowing applications to call methods, or operations, in server objects running in other applications, potentially programmed in different languages and running on different machines. There is a **CORBA** package available from the Omegahat Project (at <http://www.omegahat.org/RSCORBA/>), currently for Unix but a Windows version looks to be possible.

This package allows R commands to be used to locate available CORBA servers, query the methods they provide, and dynamically invoke methods on these objects. R values given

as arguments in these calls are exported in the call and made available to that operation invocation. Primitive data types (vectors and lists) are exported by default, while more complex objects are exported by reference. Examples of using this include communicating with the Gnumeric (<http://www.gnumeric.org>) spreadsheet, and also interacting with the data visualization system `ggobi`.

One can also create CORBA servers within R, allowing other applications to call these methods. For example, one might offer access to a particular dataset or to some of R's modelling software. This is done dynamically by combining R data objects and functions. This allows one to explicitly export data and functionality from R.

One can also use the **CORBA** package to achieve distributed, parallel computing in R. One R session acts as a manager and dispatches tasks to different servers running in other R worker sessions. This uses the ability to invoke asynchronous or background CORBA calls in R. More information is available from the Omegahat Project, at <http://www.omegahat.org/RSCORBA/>.

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## Function and variable index

### B

`bind.db.proxy` ..... 13

### C

`cat` ..... 3, 21  
`close` ..... 16, 20  
`close.socket` ..... 24  
`closestream` ..... 18  
`copystream` ..... 18  
`count.fields` ..... 6

### D

`data.restore` ..... 9  
`db.clear.result` ..... 13  
`db.connect` ..... 13  
`db.execute` ..... 13  
`db.fetch.result` ..... 13  
`db.read.column` ..... 13  
`db.read.table` ..... 13  
`db.result.columns` ..... 13  
`db.result.get.value` ..... 13  
`db.result.rows` ..... 13  
`db.write.table` ..... 13  
`dbConnect` ..... 16  
`dbExec` ..... 16  
`dbExecStatement` ..... 16

### F

`fetch` ..... 16  
`file` ..... 20  
`format` ..... 3  
`ftable` ..... 8

### H

`hdf5` ..... 19  
`httpClient` ..... 24

### M

`make.socket` ..... 24  
`mysqlClose` ..... 17  
`mysqlConnect` ..... 17  
`mysqlDataSeek` ..... 17  
`mysqlFetchRow` ..... 17  
`mysqlFreeResult` ..... 17  
`mysqlNumRows` ..... 17  
`mysqlQuery` ..... 17  
`mysqlSelect` ..... 17

`mysqlStoreResult` ..... 17  
`MySQL` ..... 16

### N

`netCDF` ..... 19

### O

`odbcClose` ..... 14  
`odbcConnect` ..... 14  
`odbcFetchRows` ..... 14  
`odbcQuery` ..... 14  
`open` ..... 20  
`openstream` ..... 18

### P

`pipe` ..... 20  
`pushBack` ..... 22  
`pushBackLength` ..... 22

### R

`read.csv` ..... 6  
`read.csv2` ..... 6  
`read.delim` ..... 6  
`read.delim2` ..... 6  
`read.dta` ..... 9  
`read.ftable` ..... 8  
`read.fwf` ..... 6  
`read.mtp` ..... 9  
`read.octave` ..... 9  
`read.socket` ..... 24  
`read.spss` ..... 9  
`read.table` ..... 5, 21  
`read.table.url` ..... 24  
`read.xport` ..... 9  
`readBin` ..... 23  
`readchar` ..... 18  
`readcomplex` ..... 18  
`readfloat` ..... 18  
`readint` ..... 18  
`readlines` ..... 18  
`readLines` ..... 7, 21  
`readSfile` ..... 9  
`reshapeLong` ..... 7

**S**

scan .....	2, 6, 21
scan.url .....	24
seek .....	22
showConnections .....	22
sink .....	4, 21
sql.insert .....	13
sql.select .....	13
sqlCopy .....	14
sqlFetch .....	14
sqlGetResults .....	14
sqlQuery .....	14
sqlSave .....	14
sqlTables .....	14
stack .....	7
stderr .....	20
stdin .....	20
stdout .....	20

**T**

textConnection .....	20
truncate .....	18

**U**

unstack .....	7
---------------	---

**W**

write .....	3, 21
write.dta .....	9
write.socket .....	24
write.table .....	3, 21
writeBin .....	23
writchar .....	18
writcomplex .....	18
writefloat .....	18
wrint .....	18
writelines .....	21



# Concept index

## A

AWK ..... 2

## B

Binary files ..... 18, 23

## C

comma separated values ..... 3  
Connections ..... 20, 21, 22  
CORBA ..... 24  
CSV files ..... 3, 6

## D

DBMS ..... 10  
DCOM ..... 24

## E

Exporting to a text file ..... 3

## F

File connections ..... 20  
Fixed-width-format files ..... 6  
Flat contingency tables ..... 8

## H

Hierarchical Data Format ..... 19

## I

Importing from other statistical systems ..... 9

## L

lynx ..... 24

## M

Mini SQL database system ..... 17  
Minitab ..... 9  
Missing values ..... 4, 5  
MySQL database system ..... 15, 16

## N

network Common Data Form ..... 19

## O

Octave ..... 9  
ODBC ..... 10, 14  
Open Database Connectivity ..... 10, 14

## P

perl ..... 2, 6  
Pipe connections ..... 20  
PostgreSQL database system ..... 12, 15  
proxy data frame ..... 13  
Pushback on a connection ..... 22

## Q

Quoting strings ..... 4, 5

## R

Re-shaping data ..... 7  
Relational databases ..... 10

## S

S-PLUS ..... 9  
SAS ..... 9  
Spreadsheet-like data ..... 5  
SPSS ..... 9  
SQL queries ..... 11  
Stata ..... 9

## T

Terminal connections ..... 20  
Text connections ..... 20

## U

Unix tools ..... 2

## W

wget ..... 24

## X

XML ..... 4