

# 10. Interfacing R

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# Interfacing R

With Bioconductor, R can do a **huge** proportion of the analyses you'll want – but not everything

- Intensive (or anachronistic) C++, FORTRAN work, e.g. for pedigrees
- 'Speciality' analyses; some need different computing architecture
- Fancy interactive graphics

R can be used to 'manage' other software. Today we'll illustrate some favorite examples

# Starting other software

NB these commands are for Windows only; see help files for e.g. Unix versions

- shell() does the equivalent of a DOS-style command
- shell("notepad") starts the Notepad editor
- If the command takes arguments, put them in the same string; shell("notepad myfile.txt")

The system() and shell.exec() commands do much the same thing.

# Starting other software

Some more options for shell();

- wait; R 'hangs' until completion
- translate; makes forward and backslashes work properly
- intern; return the output as an R object

For other options see the system() help page, for example minimized=TRUE.

Paths for files can be a little messy; shell() starts in your working directory (find it using getwd()). For files outside of this, give the full pathway.

paste() is useful, if you need to do a lot of this sort of thing.

### Examples

Code for a really mundane job;

```
for(i in 1:100){
    infile <- paste("gene",i,"data.txt", sep="")
    outfile <- paste("gene",i,"phase.out", sep="")
    shell(paste("PHASE",infile,outfile))
}</pre>
```

... this will churn away for hours, although with no error-control.

```
Why did we use wait=TRUE here? (the default)
```

### Examples

- WinBUGS implements Bayesian analyse; it's not super-fast but is very flexible
- It needs special (& clever) architecture to achieve this
- WinBUGS' input, output, graphics are all rather clunky
- R is better; so R2WinBUGS calls WinBUGS for the difficult bits, and does all the 'translation' itself
- This is done with (repeated) use of system()

## Outline

Many programs already exist to do useful analyses. It is more convenient to call them from R than to rewrite them in R.

Sometimes this involves calling the C code directly, sometimes just involves using R to write input files for another program

Examples:

- Graphviz: drawing networks
- PMF: input files for ancient Fortran software
- Google Earth: displaying outliers in context.

GraphViz (http://www.graphviz.org) is a free program for drawing networks, written by AT&T researchers.

Its input format looks like

```
"15" [shape= box,regular=1 ,height= 0.5 ,width= 0.75 ,style=filled,color= grey ] ;
"16" [shape= circle ,height= 0.5 ,width= 0.75 ,style=filled,color= grey ] ;
"2x3" [shape=diamond,style=filled,label="",height=.1,width=.1] ;
"2" -> "2x3" [dir=none,weight=1] ;
"3" -> "2x3" [dir=none,weight=1] ;
"2x3" -> "1" [dir=none,weight=2] ;
"2x3" -> "4" [dir=none,weight=2] ;
"2x3" -> "5" [dir=none,weight=2] ;
"2x3" -> "6" [dir=none,weight=2] ;
```

The sem package uses GraphViz to display path diagrams for structural equation models and the gap package uses it to draw pedigrees.

In gap the pedtodot() function writes a GraphViz input file from a pedigree in GAS or LINKAGE format.

	pid	id	fid	mid	sex	aff	GABRB1	D4S1645
1	10081	1	2	3	2	2	7/7	7/10
2	10081	2	0	0	1	1	-/-	-/-
3	10081	3	0	0	2	2	7/9	3/10
4	10081	4	2	3	2	2	7/9	3/7
5	10081	5	2	3	2	1	7/7	7/10
6	10081	6	2	3	1	1	7/7	7/10
7	10081	7	2	3	2	1	7/7	7/10
8	10081	8	0	0	1	1	-/-	-/-
9	10081	9	8	4	1	1	7/9	3/10
10	10081	10	0	0	2	1	-/-	-/-
11	10081	11	2	10	2	1	7/7	7/7
12	10081	12	2	10	2	2	6/7	7/7
13	10081	13	0	0	1	1	-/-	-/-
14	10081	14	13	11	1	1	7/8	7/8
15	10081	15	0	0	1	1	-/-	-/-
16	10081	16	15	12	2	1	6/6	7/7

First the code prints nodes for each individual, with sex and affectedness information

```
for (s in 1:n) cat(paste("\"", id.j[s], "\" [shape=",
    sep = ""), shape.j[s], ",height=", height, ",width=",
    width, ",style=filled,color=", shade.j[s], "] ;\n")
```

#### giving output like

"16" [shape= circle ,height= 0.5 ,width= 0.75 ,style=filled,color= grey ] ;

It then works out all the matings and creates small nodes for each mating and lines connecting the parents to these nodes

giving output like

```
"2x3" [shape=diamond,style=filled,label="",height=.1,width=.1] ;
"2" -> "2x3" [dir=none,weight=1] ;
"3" -> "2x3" [dir=none,weight=1] ;
```

and then connects children to parents.



[Bioconductor also has GraphViz more integrated with R in the RGraphViz package]

### **PMF:** factor analysis

PMF is a program for constrained factor analysis in analytic chemistry. It is controlled by an ugly text input file:

```
pmfini<-c(" ##PMF2 .ini file for: Simulations from R",</pre>
" ## Monitor code M: if M>1, PMF2 writes output every Mth step",
" ## For finding errors, use M<1 to output debug information",
" ##
          М
                  PMF2 version number",
11
          1
                     4.2",
" ## Dimensions: Rows, Columns, Factors. Number of \"Repeats\"",
11
                  0nt.0
                             @ns@
                                         @sources@
                                                           1".
" ##
     \FPEAK\" (>0.0 for large values and zeroes on F side)",
11
       @FPEAK@".
" ## Mode(T:robust, F:non-robust) Outlier-distance
                                                            (T=True F=False)",
11
               @isrobust@
                                                   CoutlierC".
" ## Codes C1 C2 C3 for X_std-dev, Errormodel EM=[-10 ... -14]",
11
          0.0100
                    0.0000
                              0.0100
                                         -12",
" ## G Background fit: Components Pullup_strength",
11
                              0
                                      0.0000".
                                      Initially skipped",
" ## Pseudorandom numbers:
                             Seed
11
                              @seed@
                                             0",
" ## Iteration control table for 3 levels of limit repulsion \"lims\"",
```

The @value@ are places where we want to substitute in a value.

### **PMF:** factor analysis

R code for the substitutions looks like

```
temp<-gsub("@FPEAK@",formatC(fpeak,digits=4,format="f"),pmfini)
temp<-gsub("@isrobust@",isrobust,temp)
seed<-as.character(as.integer(seed))
temp<-gsub("@seed@", seed, temp)</pre>
```

We can write data files needed by PMF, and then write the control file, then call PMF with the system() function. After PMF finishes we read in the results.

```
write.table(cX,file=xfile, quote=FALSE, col.names=FALSE, row.names=FALSE)
write.table(cU,file=efile, quote=FALSE, col.names=FALSE, row.names=FALSE)
if (!debug)
    on.exit(unlink(c(xfile,efile,inifile)))
writeLines(temp,inifile)
```

```
sysval<-system(paste(pmf,inifile), intern=TRUE,invisible=!debug)</pre>
```

ffactor<-read.table(outfiles\$f,row.names=sourcenames,col.names=species)
gfactor<-read.table(outfiles\$g,row.names=times,col.names=sourcenames)</pre>

From the user's viewpoint it looks as though everything was done in R.

# SVG+tooltips

SVG (Scalable Vector Graphics) is a non-bitmap graphics format for the web.

The RSvgDevice and RSVGTipsDevice packages allow R output to SVG format.

We can use this to create graphs with links and tooltips. For example, a funnelplot showing associations between a large number of SNPs and VTE.

Point at a dot to see the SNP it represents, and click to go to information about the gene.

# SVG+tooltips

Google Earth is controlled by KML files specifying locations. KML is another plain text format.

and then send it to Google Earth with the shell.exec(filename) function, which opens a file using whatever is the appropriate program.

# Google Earth

The identify() function lets the user select a point on a scatterplot.



In this example the points are locations where air pollution was measured, and we can call Google Earth to look at the location.