





Data manipulation in R

A program to use when size matters

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1/50





Why use R?

- Its free
- Its available on most operating systems Windows, OS X, Linux
- There are huge numbers of packages available
- Its becoming the international standard for statistics

Todays workshop

- A common scenario
- A friend has emailed you her data in a spreadsheet
- Todays workshop is not about impressing with R code

Why not use a spreadsheet?

- Data manipulation in Excel is VERY risk and time consuming
- A range of software packages are available for Excel
- Large data sets can exceed the size limits of standard programs
- Spreadsheets don't have the inherent understanding of statistics that R has
- For example handling of NA's
- R is hot!

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James P. Howard. R Cookbook.

O'Reilly Media, Inc, 2011.

Phil Spector.

Data Manipulation with R. Use R series Springer, 2008

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Download it

- Open http://www.r-project.org
- Click CRAN (Under download on Top Left)
- Click http://cran.ms.unimelb.edu.au/ University of Melbourne

Windows

- Select Windows
- Select Base
- Download R (suggest latest version)

OS X

- Select Select OS X
- Select R-3.2.2.pkg (or the version that matches your OS version)

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5/50





2+5
[1] 7
Create a sequence of numbers
X = 2:10
Display basic statistical measures
summary(X)
Min. 1st Qu. Median Mean 3rd Qu. Max.
2 4 6 6 8 10
use q() to quit





low about RStudio

- https://www.rstudio.com/products/rstudio/download/
- Its also on your thumb drive

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To access the documentation type

help.start() help(summary) args(summary) example(sd) ??package

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To search R documentation

- RSiteSearch("key phrase")
- help(adf.test,package="tseries")
- To search for a tutorial for a package vignette(package="packagename")
- For an intro to vignettes see https://cran.r-project.org/web/packages/sos/vignettes/sos.pdf
- Examples on the web http://shiny.rstudio.com/gallery/

Custom Google search focused on R-specific websites

http://rseek.org

Coding Q&A site

http://stackoverflow.com http://stats.stakexchange.com

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Vectors

- Vectors $I \leftarrow c(1, 3, 4, 7, 11)$
- Refer to elements using array I[c(2,5)] 2nd and 5th elements of I

Data Frames

```
a <- c(35,23,24,65)
e <- c("Peter", "John", "Mark", NA)
f <- c(TRUE,TRUE,TRUE,FALSE)
team <- data.frame(a,e,f)
names(team) <- c("Age","Names","Passed") # variable names
str(team)
## 'data.frame': 4 obs. of 3 variables:
## $ Age : num 35 23 24 65
## $ Names: Factor w/ 3 levels "John","Mark",..: 3 1 2 NA
## $ Passed: logi TRUE TRUE TRUE FALSE</pre>
```



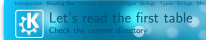


Research on how to work creatively based on case studies of successful R&D projects developed into Agile

- Keep the "manages" away
- Work sustainably
- People over process
- Iterative development

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10/5





Where are we

```
getwd()
setwd("/Users/pcru/SizeDoesMatter1")
dir() #This lists the files
ls() #This lists the variables
```

http://www.statmethods.net/input/contents.html

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To read a csv table as a table tr

 $\texttt{tab1} \leftarrow \texttt{as.matrix}(\texttt{read.csv}(\texttt{file} = \texttt{"filetable.csv"}, \texttt{sep} = \texttt{","}, \texttt{header} = \texttt{FALSE}))$

But our table is an excel file

- What about a package?
- http

//www.thertrader.com/2014/02/11/a-million-ways-to-connect-r-and-excel/

- Installing the R package xlsx
- CRAN mirror http://cran.csiro.au
- Change in preferences

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table1-read.xlsx2("1_R Wkshp_dummy data_OTU table.xlsx", sheetName =
"Sheet1",header=FALSE,rowNames=FALSE,transpose=TRUE,endRow=18)

Loading the xlsx package

```
## Loading required package: xlsx
## Warning: package 'xlsx' was built under R version 3.1.3
## Loading required package: rJava
## Warning: package 'rJava' was built under R version 3.1.3
## Loading required package: methods
## Loading required package: xlsxjars
## Loading required package: xtable
```





Where from

- install command
- install.packages(pkgs)

Citing Packages

- Citing packages
- Getting the bibtex entry into endnote
- http://www.lib.uts.edu.au/question/5955/ how-can-i-import-bibliography-endnote-bibtex-latex-what-about-converting-other-way

```
 \begin{tabular}{ll} $\times$-citation() \\ xit-citation(package="RSQLite") \\ to Bibtex(x) \\ sessionInfo() \\ packages_in_use \leftarrow c(sessionInfo()\$basePkgs, names(sessionInfo()\$loadedOnly)) \\ the_citations_list \leftarrow lapply(X=packages_in_use, FUN=citation) \\ the_citations_list \leftarrow lapply(X=packa
```

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	X1	X2	Х3	X4	X5	X6	X7
1	Group	Contaminated					
2	Site	1			2		
3	Sample ID	10000	10001	10002	10003	10004	10005
4	Rep	1	2	3	1	2	3
5	phormidiaceae	24872	24872	5822	7538	7201	7538
6	streptococcaceae	11	7	14	8	10	8

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Transposing

We need to transpose the table and set the column names correctly

```
table1t=setNames(data.frame(t(table1[,-1])),table1[,1])
```

 $\label{lem:http://rgm3.lab.nig.ac.jp/RGM/R_rdfile?f=Ecdat/man/read.transpose.Rd&d=R_CC http://stackoverflow.com/questions/17288197/reading-a-csv-file-organized-horizontally$

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17/50





require(stringr)

A look at the stringer package

stri_c(str1,str2)

concatenates two string

str_len(str)

require(stringr)

Loading required package: stringr

```
table1t$Rep<-str_replace(table1t$Rep,"[rep]{3}?","\\1")
table1t$Rep<-str_replace(table1t$Rep,"A","1")
table1t$Rep<-str_replace(table1t$Rep,"B","2")
table1t$Rep<-str_replace(table1t$Rep,"C","3")
table1t$Rep<-as.factor(table1t$Rep)</pre>
```

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TDD - First do it the easy way first

```
ctridx<-which(table1t$Group=="Control")</pre>
table1t$Group[1:48] <- "Contaminated"
table1t$Group[(ctridx+1):48]<-"Control"
    ttt \leftarrow table1t\$Site
    \quad \quad \quad \text{for(i in } c(2:length(table1t\$Site)))
        temp \leftarrow as.character(table1t\$Site[i])
        \mathsf{tempb} {\leftarrow} \mathsf{as.character}(\mathsf{ttt}[\mathsf{i}{-}1])
        if (table1t $Site[i]==""")
             ttt[i] \leftarrow tempb
        if (! table1t $ Site [( i )]=="")
            ttt\,[\,i\,]\!\leftarrow\!temp
    table1t $Site ←ttt
## X3
## 1
## Levels: 1 2 3 4 FALSE TRUE
## X4
## Levels: 1 2 3 4 FALSE TRUE
## X5
```

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Levels: 1 2 3 4 FALSE TRUE

X6

2

Reading Tables Reading a table of other types



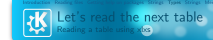
- http://www.statmethods.net/input/importingdata.html
- http://stackoverflow.com/questions/17288197/ reading-a-csv-file-organized-horizontally
- http://rgm3.lab.nig.ac.jp/RGM/R_rdfile?f=Ecdat/man/read. transpose.Rd&d=R_CC
- Input files from Stata

```
library(foreign)
mydata ← read.dta("c:/mydata.dta")
```

Levels: 1 2 3 4 FALSE TRUE

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setwd("/Users/pcru/SizeDoesMatter1")



Need coffee !!

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Oh NO

- All columns have been set to factors
- Dates have different formats

```
str(table2[,1:11])
## 'data.frame': 48 obs. of 11 variables:
                           : Factor w/ 2 levels "Contaminated",..: 1 1 1 1 1 1 1 1 1 1 ...
## $ Group
## $ Site
                           : Factor w/ 4 levels "1", "2", "3", "4": 1 1 1 2 2 2 1 1 1 2 ...
                           : Factor w/ 18 levels "10000", "10001", ...: 1 2 3 4 5 6 7 8 9 1 ...
## $ Sample.ID
                           : Factor w/ 9 levels "1","2","3","A",...: 1 2 3 1 2 3 7 8 9 7 ...
## $ Rep
                           : Factor w/ 2 levels "14-May-14", "N/A": 1 1 1 1 1 1 1 1 1 1 ...
## $ Spill.date
## $ Sample.collection.date: Factor w/ 4 levels "15.5.14","17/5/14",..: 1 1 1 1 1 2 2 2 2 2 ...
## $ labnum
                           : Factor w/ 36 levels "2000", "2001", ...: 1 2 3 4 5 6 7 8 9 19 ...
                           : Factor w/ 39 levels "10", "105", "108", ... 27 30 28 26 25 27 12 15 13 7
## $ phosphate..ppb.
## $ ammonia..ppb.
                           : Factor w/ 41 levels "10","103","1042",...: 10 14 15 6 7 4 31 34 32 28 .
## $ chlorophyll..ug.L.
                          : Factor w/ 38 levels "1", "10", "11", ...: 20 23 21 25 17 18 16 14 15 12 ...
                           : Factor w/ 31 levels "100", "120", "31", ...: 5 4 3 7 6 5 8 7 9 11 ...
## $ DO....
```

#dir()
table2<-read.xlsx2("2_R Wkshp_dummy data_Env Data_incl2outliersMK.xlsx", sheetName ="Sheet2</pre>

	Group	Site	Sample.ID	Rep	Spill.date	Sample.collection.date
1	Contaminated	1	10000	1	14-May-14	15.5.14
2	Contaminated	1	10001	2	14-May-14	15.5.14
3	Contaminated	1	10002	3	14-May-14	15.5.14
4	Contaminated	2	10003	1	14-May-14	15.5.14
5	Contaminated	2	10004	2	14-May-14	15.5.14
6	Contaminated	2	10005	3	14-May-14	15.5.14

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Site

Break it down

##

First read a few rows only

Group

table2<-read.xlsx2("2_R Wkshp_dummy data_Env Data_incl2outliersMK.xlsx", sheetName = "Sheet
sapply(table2,mode)</pre>

Spill.date

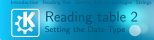
Sample.ID

```
"character"
                     "numeric"
                                   "numeric" "character"
                                                              "character"
       rowNames as.Data.frame
       "logical"
                     "logical"
sapply(table2,class)
                                                              Spill.date
          Group
                          Site
                                   Sample.ID
                                                       Rep
     "character"
                     "numeric"
                                   "numeric"
                                               "character"
                                                              "character"
##
       rowNames as.Data.frame
      "logical"
                     "logical"
```

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colClasses

- The variable colClasses can be used to specify the row types.
- We need to set **stringsAsFactor=FALSE** or all columns with be loaded as factors
- The dates are in a non-standard format so we need to read them as chars first

table2b<-read.xlsx2("2_R Wkshp_dummy data_Env Data_incl2outliersMK.xlsx", sheetName = "Sheet2",heade:
sapply(table2,class)</pre>

```
## Group Site Sample.ID Rep Spill.date
## "character" "numeric" "numeric" "character"
## rowNames as.Data.frame
## "logical" "logical"
```

table2f<-table2 table2f\$Spill.date<-as.Date(table2f\$Spill.date,"%d-%b-%y") table2f\$Sample.collection.date<-as.Date(table2f\$Sample.collection.date,"%d.%m.%y") ## Error in as.Date.default(table2f\$Sample.collection.date, "%d.%m.%y"): do not know how to convert 'table2f\$Sample.collection.date' to class "Date" asses=c(" #sapply(table2f,mode) sapply(table2f,class) ## Spill.date Group Site Sample.ID "character" "numeric" "numeric" "Date "character" rowNames as.Data.frame "logical" "logical"

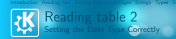
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25/50



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things up Setting the re





- The as.Data method can take a format string as the second variable
- The format strings are described in help on strptime
- But Spill.data has **two formats**
- We can use the if else function to combine them

table2bf<-table2b
table2bf\$Spill.date<-as.Date(table2bf\$Spill.date,"%d-%b-%y")
cdate1<-as.Date(table2bf\$Sample.collection.date,"%d.%m.%y")
cdate2<-as.Date(table2bf\$Sample.collection.date,"%d/%m/%y")
table2bf\$Sample.collection.date<-as.Date(ifelse(!is.na(cdate1),as.Date(cdate1),as.Date(cdate2)), origtable2bf\$Group<-as.factor(table2bf\$Group)
table2bf\$Rep<-as.factor(table2bf\$Rep)
dated<-table2bf\$Sample.collection.date-table2bf\$Spill.date</pre>

Count the NAs

na_count <-sapply(table2bf, function(y) sum(length(which(is.na(y)))))
na_count</pre>

##	Group	Site	Sample.ID
##	0	0	0
##	Rep	Spill.date	Sample.collection.date
##	0	24	0
##	labnum	phosphateppb.	ammoniappb.
##	0	0	0
##	chlorophyllug.L.	DO	rowNames
##	0	0	0
##	as.Data.frame		
##	0		

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```
Reading table 2

Just fix the Rep column using the stringer package again
```

```
require(stringr)
table2bf$Rep<-str\_replace(table2bf$Rep,"[rep]{3}?","\\1")
table2bf$Rep<-str_replace(table2bf$Rep,"A","1")</pre>
table2bf$Rep<-str_replace(table2bf$Rep,"B","2")</pre>
table2bf$Rep<-str_replace(table2bf$Rep, "C", "3")
table2bf$Rep<-as.factor(table2bf$Rep)</pre>
str(table2bf)
## 'data.frame': 48 obs. of 13 variables:
## $ Group
                           : Factor w/ 2 levels "Contaminated",..: 1 1 1 1 1 1 1 1 1 1 ...
## $ Site
                           : num 1 1 1 2 2 2 1 1 1 2 ...
                           : num 10000 10001 10002 10003 10004 ...
## $ Sample.ID
                           : Factor w/ 3 levels "1", "2", "3": 1 2 3 1 2 3 1 2 3 1 ...
## $ Rep
   $ Spill.date
                           : Date, format: "2014-05-14" "2014-05-14" ...
##
   $ Sample.collection.date: Date, format: "2014-05-15" "2014-05-15" ...
                           : num 2000 2001 2002 2003 2004 ...
   $ labnum
   $ phosphate..ppb.
                           : num 3020 3253 3169 2999 2879 ...
   $ ammonia..ppb.
                           : num 13880 14598 14676 10984 11657 ...
## $ chlorophyll..ug.L.
                           : num 302 323 315 352 289 296 254 248 250 220 ...
## $ DO....
                           : num 34 33 31 38 36 34 40 38 41 45 ...
## $ rowNames
                           : logi FALSE FALSE FALSE FALSE FALSE ...
                           : logi FALSE FALSE FALSE FALSE FALSE ...
## $ as.Data.frame
```

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Provided

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The inbuilt command merge

- R has a command merge
- To begin, start looking at the first 9 lines of the tables and merge them
- Need to use Group, Site, Sample.ID because otherwise it's not uniques

```
merge(x, y, by = intersect(names(x), names(y)),
    by.x = by, by.y = by, all = FALSE, all.x = all, all.y = all,
    sort = TRUE, suffixes = c(".x",".y"),
    incomparables = NULL, ...)

tablc<-tablelt[1:9,]
tab2c<-table2b[1:9,]
m1<-merge(tablc,tab2c,by.x="Sample ID",by.y="Sample.ID")
m2<-merge(tablelt,table2bf,by.x=c("Group","Site","Sample ID"),by.y=c("Group","Site","Sample ID"),by.y=c("Group","Site","Sample ID","Rep"),by.y=c("Group","Site","Site","Sample ID","Rep"),by.y=c("Group","Site","Site","Site","Sample ID","Rep"),by.y=c("Group","Site","Site","Site","Sample ID","Rep"),by.y=c("Group","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","Site","S
```

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Follow up data from contaminated site

```
table3←read.xlsx2("3_Follow up data from contaminated site_MK.xlsx", sheetName = "Sheet1",header=TRUE,row rep("character",2),rep("numeric",18))) table3f←table3 table3f$Spill.date←as.Date(table3f$Spill.date,"%d.%m.%y") table3f$Sample.collection.date←as.Date(table3f$Sample.collection.date,"%d.%m.%y") sapply(table3f,mode) sapply(table3f,class)
```

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Joining table 3 to are merged tables

- We need to be careful to match everything
- Install the plyr package This has lots of useful functions for renaming var etc
- This means we need columns for corynebacteriaceae and porphyromondaceae
- Should these values be NA or 0?
- We will do one of each.
- Generally we would use NA but in this case 0 is better as its likely the rows were missing as non were detected

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```
require (plyr)
 Sample . ID←rep (20000,3)
 table3fi←cbind(table3f,Sample.ID)
#how many columns I can't count
 ncol(table3fi)
ncol(m3)
#now get the cols all right table3fii←table3fi[c(1,2,24,3,4:23)]
m3k-m3[c(1:4,19:20,5:18,21:26)]
ms:=ms[c[1:4,19:20,3:10,21:20]]
setdiff(names(m3i),names(table3fii))
m3i:=rename(m3i,c("Sample ID"="Sample.ID"))
corynebacteriaceae=rep(0,nrow(table3fii))
corynebacteriaceae←rep(Na,now(table3fii))
porphyromondaceae←rep(Na,nrow(table3fii))
table3fiii←cbind(table3fii, corynebacteriaceae, porphyromondaceae)
 setdiff(names(m3ii),names(table3fiii))
 \texttt{m3ii}\left[\,,c\left(7\!:\!24\right)\right] \,\leftarrow\, \texttt{sapply}\left(\,\texttt{m3ii}\left[\,,c\left(7\!:\!24\right)\right]\,,\texttt{as.numeric}\,\right)
 m3ii[,c(1:4)] \leftarrow sapply(m3ii[,c(1:4)],as.character)
#m3ii[,c("Site")] 

sapply (m3ii[,c("Site")], as.character)
table3fiii [,c(1:4)] \leftarrow sapply(table3fiii [,c(1:4)], as.character) table3fiii [,c(7:24)] \leftarrow sapply(table3fiii [,c(7:24)], as.numeric)
table4←rbind (m3ii, table3fiii
table4[,1] ← sapply(table4[,1], as.factor)
```

Loading required package: plyr

```
## [1] 24
## [1] 27
## [1] "Sample ID" "corynebacteriaceae"
```

[1] "Sample ID" "corynebacteriaceae" "porphyromondaceae"

character(0)

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reshape2

- vignette(reshape) doesn't work
- try http://had.co.nz/reshape/
- and http://seananderson.ca/2013/10/19/reshape.html

A small example for melt

- Suppose we what a box plot to see if there are outliers
- We will use ggplot2 box plot
- The box plot needs data in long format.
- To use this first **melt** the data
- We need to specify the unique key, the variable name and the value name
- The key is not unique.
- Then plot it

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The code

```
\label{eq:matable4} $$ \mathtt{matable4} = \mathtt{mat
```

require(reshape2)

Loading required package: reshape2

matable4<-melt(table4[,c(1:4,7:25)],variable.name = "microbe",value.name = "abundance", id=c("Group",</pre>

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Removing Outliers

- Outliers are defined 1.5 times the interquartile range above the upper quartile
- Assume that rows 12 and 14 in phosphate are errors as the 9 is typed twice
- Still issues with ammonia to explore

```
phosphate<-table4[,"phosphate..ppb."]
upper.limit <- quantile(phosphate) [4] + 1.5*IQR(phosphate)
lower.limit <- quantile(phosphate) [2] - 1.5*IQR(phosphate)
#table4[phosphate> upper.limit,c("Site","phosphate..ppb.")]
```

	Site	phosphateppb.
1	1	3020.00
2	1	3253.00
3	1	3169.00
12	1	9982.00
14	1	9982.00
16	1	1542.00

table4[12,"phosphate..ppb."]<-982
table4[14,"phosphate..ppb."]<-982</pre>

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Using ggplot

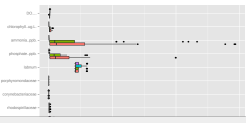
- As we have keys we need to specify the x and y
- Let's make the sites different colors
- The variable names are long so flip it with coord_flip()
- Looks like we have outliers...hmm

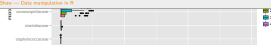
require(ggplot2)

Loading required package: ggplot2

ggplot(matable4,aes(x=microbe,y=abundance,fill=Site)) + geom_boxplot() + coord_flip()

Warning: Removed 24 rows containing non-finite values
(stat_boxplot).



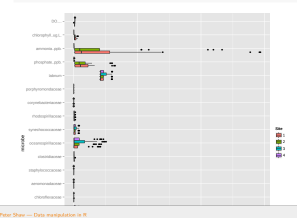




Look again ggplot

matable4<-melt(table4[,c(1:4,7:25)],variable.name = "microbe",value.name = "abundance", id= ggplot(matable4,aes(x=microbe,y=abundance,fill=Site)) + geom_boxplot() + coord_flip()

Warning: Removed 24 rows containing non-finite values
(stat_boxplot).







RSQLite

- Suppose merge is not enough? I know about SQL and want to do joins
- Install RSQLite
- We also need to install DBI

Loading required package: RSQLite

```
db <- dbConnect(SQLite(), dbname="Test.sqlite")</pre>
#qetConfiq()fstaged.queries
# sqldf(attach "Test1.sqlite" as new)
dbBegin(db)
## [1] TRUE
dbWriteTable(db,"table1",table1t,overwrite=TRUE)
## [1] TRUE
dbReadTable(db, "table1")
##
                    Group Site Sample.ID
                                             Rep phormidiaceae
## X2
             Contaminated
                                     10000
## X3
             Contaminated
                                     10001
                                               2
                                                         24872
## X4
             Contaminated
                                     10002
                                               3
                                                          5822
             contaminated
                                     10004
                                                           7ZUI
## X7
             Contaminated
                                     10005
                                               3
                                                          7538
## X8
             Contaminated
                                     10006
                                                          8467
                                     10007
```

Another important component of TDD is refactoring and unit tests

- Refactoring http://refactoring.com/
- http://www.r-bloggers.com/
 my-experience-of-learning-r-from-basic-graphs-to-performance-tuning/
- TDD in R http: //www.slideserve.com/andrew/test-driven-development-in-r
- Version Control tortiseSVN http://tortoisesvn.net/
- GitHub https://github.com/







RSQLite

- Some links to RSQL ideas
- http://stackoverflow.com/questions/12307685/ join-more-than-2-tables-in-r-using-rsqlite
- https://support.rstudio.com/hc/en-us/articles/ 201057987-Quick-list-of-useful-R-packages
- https://cran.rstudio.com/web/packages/dplyr/vignettes/introduction.html

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Dropping Row and Columns with too many NAs

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In tidy data:

- Each variable forms a column.
- Each observation forms a row.
- Each type of observational unit forms a table.
- https://cran.r-project.org/web/packages/tidyr/vignettes/tidy-data.html
- http://pj.freefaculty.org/R/Rtips.html#toc-Subsection-1.11

Spit out the dates and numbers

dates4←table4 [, c (5,6)] abundance←table4 [, c (7:25)]

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45/5





sapply

- Also known as centring the data
- Ecological percentage of the sum of the variables
- We an use sweep to centre the data
- options(digits = 1) Just to make things pretty

sweepOutContinu=sweep(abundance,2.apply(abundance,2.min,na.rm=TRUE))
afterSweepContinu=sweep(sweepOutContinu,2.apply(sweepOutContinu,2.max,na.rm=TRUE),"/")
tableS=-cbind(table4[,c(1:6)], afterSweepContinu,days)
options(digits=1)
sweep(abundance, 2, colSums(abundance), FUN="/")
scale(abundance, center=FALSE, scale=colSums(abundance))





Calculating the number of days

We can just subtract as. Date fields

```
dates4<-table4[,c(5,6)]
abundance<-table4[,c(7:25)]
days<-dates4[,2]-dates4[,1]</pre>
```

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R has nice graphs

- A graphical output
- http://rcharts.io/gallery/
- R Graph gallery currently down try http://rgraphgallery.blogspot.com/
- A reference on where to go R thumbnails
- ggplot2 (scatter plot of 2 var and then 3 plots)
- To create a correlation heat map

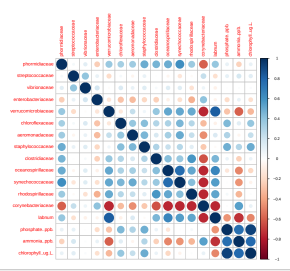
```
library(corrplot)
abuncor—cor(t5lessThan20col[.c(6:22)])
require(corrplot)
corrplot(abuncor, method = "circle")
```

[1] 23

Loading required package: corrplot

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49/50



LaTeX Beamer

http://latex-beamer.sourceforge.net/

Sharelatex Site

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https://www.sharelatex.com

A Data Cleaning Mooc

https://www.sharelatex.com



Help is on the way

- Parameterized Complexity Research Unit (PCRU) PhD students
- PhD student in Bioinformatics from Central South Uni

Your feedback on some ideas

- Using Sweave or Knitr
- Advanced Data Cleaning
- Network Centric data analysis

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Output of sessionInfo

sessionInfo()

```
## R version 3.1.2 (2014-10-31)
## Platform: x86_64-apple-darwin13.4.0 (64-bit)
## locale:
## [1] C
## attached base packages:
## [1] methods stats
                         graphics grDevices utils
                                                       datasets base
## other attached packages:
## [1] corrplot_0.73 RSQLite_1.0.0 DBI_0.3.1
                                                   ggplot2_1.0.0
## [5] reshape2_1.4.1 plyr_1.8.1 stringr_0.6.2 xtable_1.7-4
## [9] xlsx_0.5.7 xlsxjars_0.6.1 rJava_0.9-7
##
## loaded via a namespace (and not attached):
## [1] MASS_7.3-39
                        Rcpp_0.11.5
                                        colorspace_1.2-6 digest_0.6.8
```

grid_3.1.2

munsell_0.4.2

gtable_0.1.2

proto_0.3-10

[13] scales_0.2.4 tools_3.1.2

[5] evaluate_0.7.2 formatR_1.2

[9] highr_0.5

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##!ne_critations_trst <= tappitg(A=packages_tn_ase, row-critation)

labeling_0.3