# Lab 4: Learning R IGF RNA-Seq Workshop 2017

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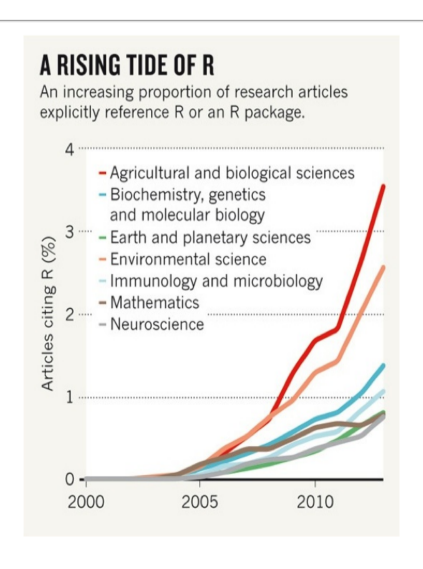
6/22/2017

# **Outline**

- Introduction of R
- Data structure (vector and data frame)
- Data importing and exporting
- Plotting
- String operations

# Why R?

- R is great at statistical computing and graphics
- R is free
- R has great community supports



# R example 1 (R for statistics)

#### Chi-square test

```
d <- c(12, 36, 24, 70)
dm <- matrix(d, nrow=2, byrow=T)
dm

[,1] [,2]
[1,] 12 36
[2,] 24 70

chisq.test(dm)</pre>
```

```
Pearson's Chi-squared test with Yates' continuity correction data: dm
X-squared = 7.8894e-31, df = 1, p-value = 1
```

# R example 2 (R for graphs)

ts="http://129.130.89.83/tmp/public/RNASeq/RNASeq2017/codes/xcard.R"
source(ts)

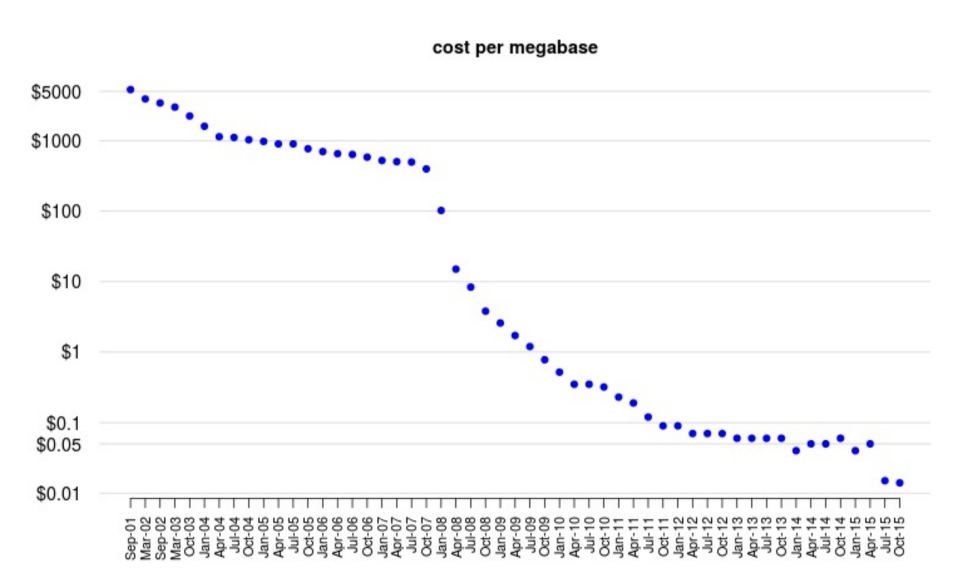


# R example 2 code

```
# Christmas tree
L <- matrix(
                0 , 0.1,
  c(0.03, 0,
   0.85, 0.00, 0.00, 0.85,
0.8, 0.00, 0.00, 0.8,
    0.2, -0.08, 0.15, 0.22,
   -0.2, 0.08, 0.15, 0.22,
0.25, -0.1, 0.12, 0.25,
-0.2, 0.1, 0.12, 0.2),
  nrow=4)
# ... and each row is a translation vector
B <- matrix(
  c(0, 0,
    0, 1.5,
    0, 1.5,
    0, 0.85,
    0, 0.85,
    0, 0.3,
    0, 0.4),
  nrow-2)
prob = c(0.02, 0.6, .08, 0.07, 0.07, 0.07, 0.07)
# Iterate the discrete stochastic map
N = 1e5 #5 # number of iterations
x = matrix(NA,nrow=2,ncol=N)
x[,1] = c(0,2) # initial point
k <- sample(1:7,N,prob,replace=TRUE) # values 1-7
 x[,i] = crossprod(matrix(L[,k[i]],nrow=2),x[,i-1]) + B[,k[i]] # iterate
# Plot the iteration history
#png('card.png')
par(bg='darkblue',mar=rep(0,4))
plot(x=x[1,],y=x[2,],
     col=grep('green',colors(),value=TRUE),
     axes=FALSE,
     cex=.1,
    xlab='',
ylab='' )#,pch='.')
bals <- sample(N,20)
cex=2,
       pch=19
text(x=-.7,y=8,
labels='Merry',
     adj=c(.5,.5),
     srt-45,
     vfont=c('script','plain'),
     cex=3,
     col='gold'
text(x=0.7,y=8,
     labels='Christmas',
     adj-c(.5,.5),
     srt=-45,
     vfont=c('script','plain'),
     cex=3,
     col='gold'
```

# R example 3 (R for graphs)

cs="http://129.130.89.83/tmp/public/RNASeq/RNASeq2017/codes/trend.R"
source(cs)



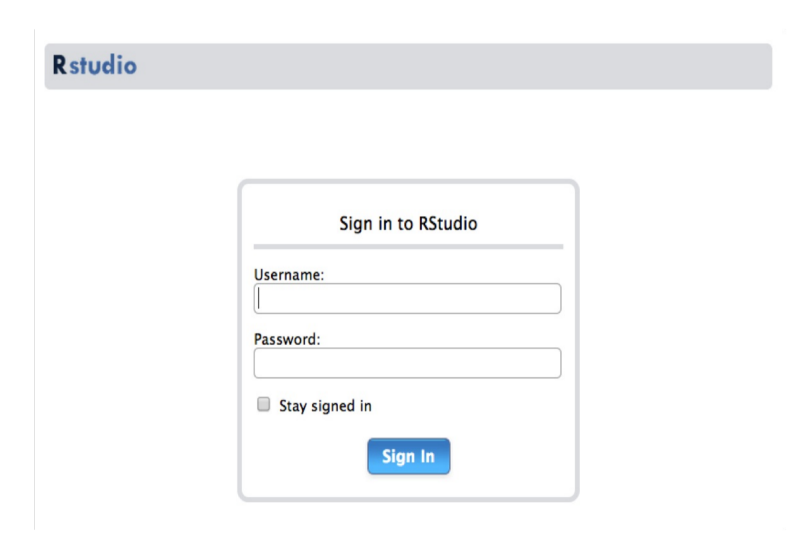
#### Where can we use R?

Rstudio is an open source integrated development environment (IDE) for R

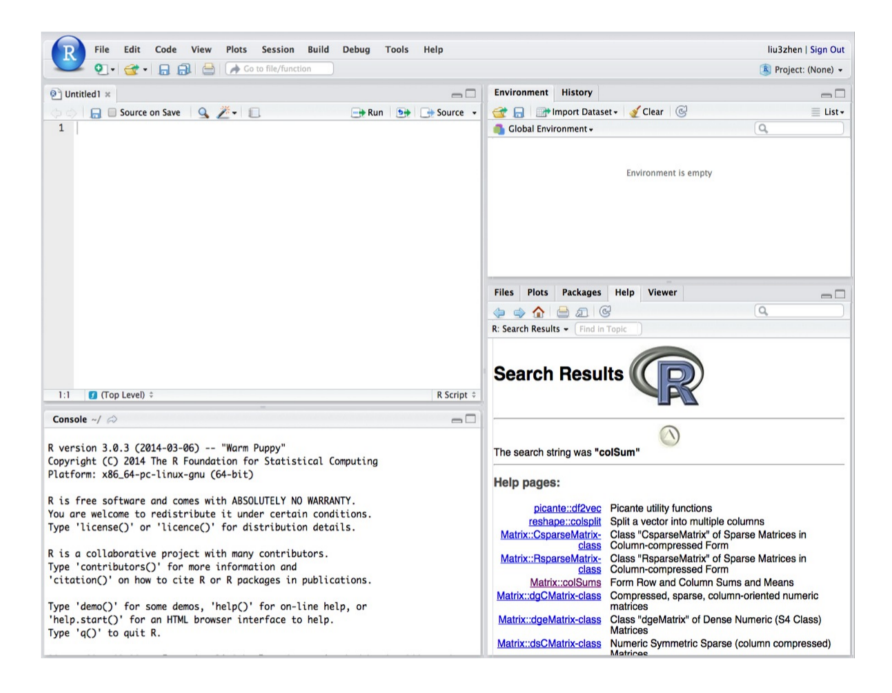
- On your own machine (Rstudio Desktop)
  - Download and install R
  - o Download and install Rstudio
- Use Rstudio at Beocat (Rstudio server)
  - o rstudio.beocat.cis.ksu.edu
  - Your KSU ID and password to login

# Rstudio at Beocat

rstudio.beocat.cis.ksu.edu



#### Rstudio interface



# Getting started, R commands

**Expression**: evaluated, printed, and the value lost

```
2 + 4

[1] 6

68 * 0.15

[1] 10.2
```

**Assignment** assign values to a variable evaluated, the value passed to a variable but NOT printed

assignment operator: <- or =</pre>

```
y <- 2
y = 2
info <- "hello world"
cat(info)</pre>
```

hello world

#### Notes

• Comments (#): Notes to scripts, starting with a hashtag ('#'), everything to the end of the line is a comment.

y <- 2 + 4 # an example of the assignment

• Variable names are case senstive

```
y <- 2
Y <- 3
y
```

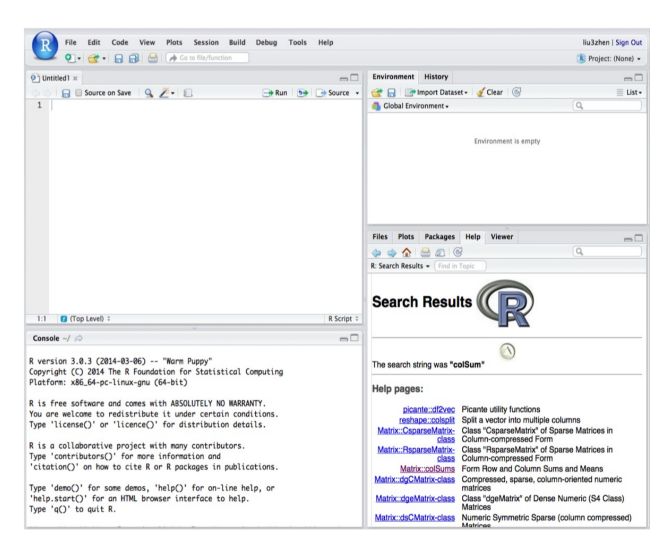
[1] 2

Υ

[1] 3

# **Executing commands**

- PC window: control + return (enter)
- Apple MAC: command + return (enter)



# vector: multiple elements

A vector is a single entity consisting of an ordered collection of numbers, characters, logical quantities, etc.

concatenate command: c()

- Numeric vector c(10.4, 5.6, 3.1, 6.4, 21.7)
- Logical vector c(TRUE, FALSE, TRUE, TRUE)
- Character vector c("a", "b", "c")
- Missing value (NA, not available)
   c("a", "b", "c", NA)

# vector manipulation (I)

```
# Numeric vector
x \leftarrow c(10.4, 5.6, 3.1, 6.4, 21.7)
sum(x)
[1] 47.2
2*x
[1] 20.8 11.2 6.2 12.8 43.4
### extract 2nd elements
x[2]
[1] 5.6
```

# vector manipulation (II)

# Logical vector

lv <- c(TRUE, FALSE, TRUE, TRUE) !lv lv == FALSE</pre>

# vector manipulation (III)

```
# Character vector
cv <- c("a", "b", "c")
cv2 <- paste(cv, 1:3, sep="")
cv2

[1] "a1" "b2" "c3"

# Missing value
mvv <- c("a", "b", "c", NA)
is.na(mvv)</pre>
```

[1] FALSE FALSE TRUE

# vector manipulation (IV)

Vectors must have their values with the same mode, either numeric, character, logical, or other types.

#### conversion to other modes

```
z <- 0:9
is.numeric(z)</pre>
```

#### [1] TRUE

```
digits <- as.character(z) # convert to character
d <- as.numeric(digits) # convert to numeric</pre>
```

# vector manipulation (V)

• Select a subset of a vector

```
x <- c(4, 5, 7, 3, 9)
x[c(2, 3)]

[1] 5 7

x[x>6]

[1] 7 9
```

[1] 5 7 3

• Modify a vector

```
x[3] <- 23.1
c(x, 10.9)
```

[1] 4.0 5.0 23.1 3.0 9.0 10.9

# Question 1

Can a vector contain different types of elements?

```
c(1, "a")
c(1, TRUE)
c(TRUE, "a")
c(1, "a", TRUE)
```

#### Data frame

A data frame may be regarded as a matrix (table) with columns possibly of differing modes

• Making data frames

```
df <- data.frame(name=c("Josh", "rose"), age=c(23, 35))
df

name age
1 Josh 23
2 rose 35</pre>
```

# Working with a data frame

```
name age
1 Josh 23
2 rose 35
```

#### Trying these commands:

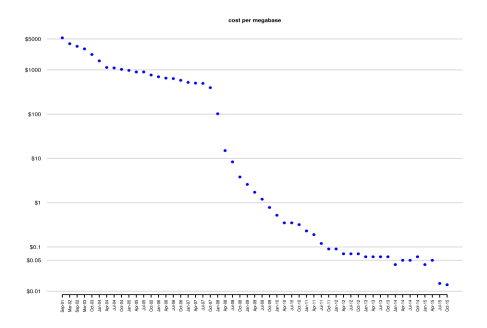
```
head(df, 1)
tail(df, 1)
str(df)
df[2, 1]
df[2, 2]
df[2]
df[, 2]
```

# Importing data

read.table(): to read a data frame (table)
read.delim, read.csv

```
cpm="http://129.130.89.83/tmp/public/RNASeq/RNASeq2017/data/cs.txt"
d <- read.delim(cpm)
head(d, 3)</pre>
```

```
Date Cost.per.Mb Cost.per.Genome
1 Sep-01 5292.39 95263072
2 Mar-02 3898.64 70175437
3 Sep-02 3413.80 61448422
```



# **Exporting data**

#### write.table() or write.csv()

To write a tab-delimited file

```
x <- data.frame(a = "pi", b = pi)
write.table(x, file="foo.txt", sep="\t", row.names=FALSE)</pre>
```

- file="foo.txt": foo.txt is the ouput file name
- sep="\t": separated by a tab (\t)
- row.names=FALSE: row names are not included in the output

### Problem

• Create a data frame

three columns: 1. Name 2. Major 3. Gender three rows (entries): your neighbors and you

- Write the data frame to an output file
- Read the file to R and add one more column (e.g., favorite color)

# Plotting: plot()

```
High-level plot: create a new plot plot(x, y, xlab, ylab, main, ...)
```

main

ylab

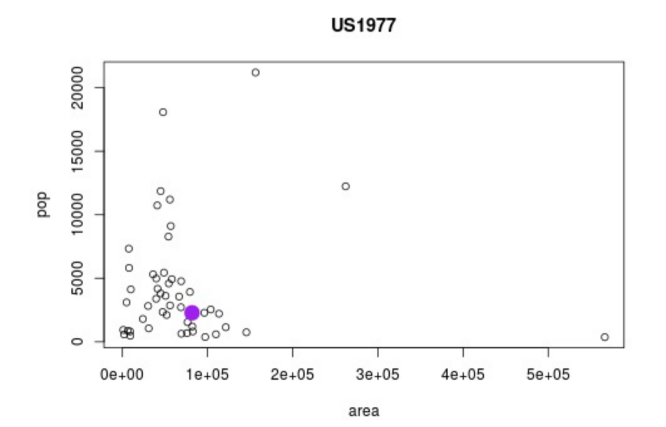
xlab

# Adding contents to a plot

Low-level plot: add to an existing plot

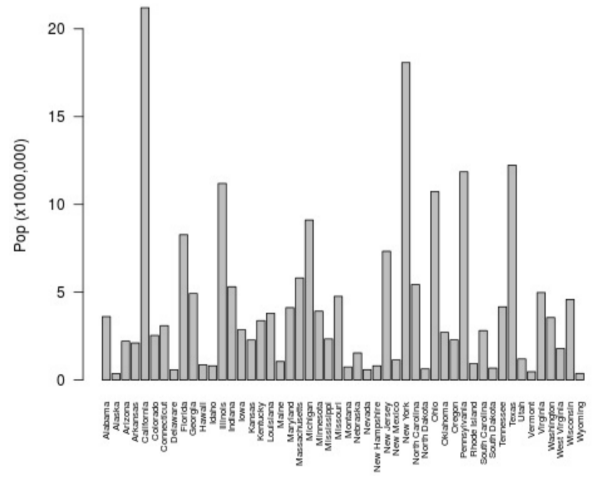
- add pointspoints()
- add lines lines()
- add text or legend text() legend()

# Scatter plot



# **Boxplot**

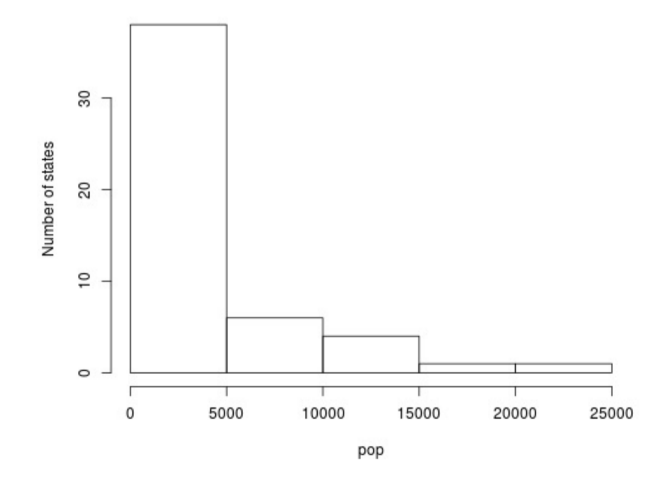




# Histogram

hist(pop, ylab="Number of states", main="US 1977 Population")





# String operations - nchar

**nchar()** nchar the sizes of the corresponding elements of a vector.

```
cvec <- c("google", "hello", "the", "world")
nchar(cvec)</pre>
```

[1] 6 5 3 5

# String operations - grep

**grep()** grep searches for matches to argument pattern within each element of a character vector

```
cvec
[1] "google" "hello" "the" "world"
grep("o", cvec)
[1] 1 2 4
```

# String operations – sub and gsub

**sub()** and **gsub()** sub and gsub perform replacement of the first and all matches respectively.

```
cvec
[1] "google" "hello" "the" "world"

sub("o", "O", cvec)

[1] "gOogle" "hello" "the" "wOrld"

gsub("o", "O", cvec)

[1] "gOOgle" "hello" "the" "wOrld"
```

# Package installation

Prepare for Lab 5: RNA-Seq analysis

```
# DESeq2
source("http://bioconductor.org/biocLite.R")
biocLite("DESeq2")

# GOSeq
biocLite("goseq")

# GO.db
biocLite("GO.db")
```

# Getting help

#### Usage of commands

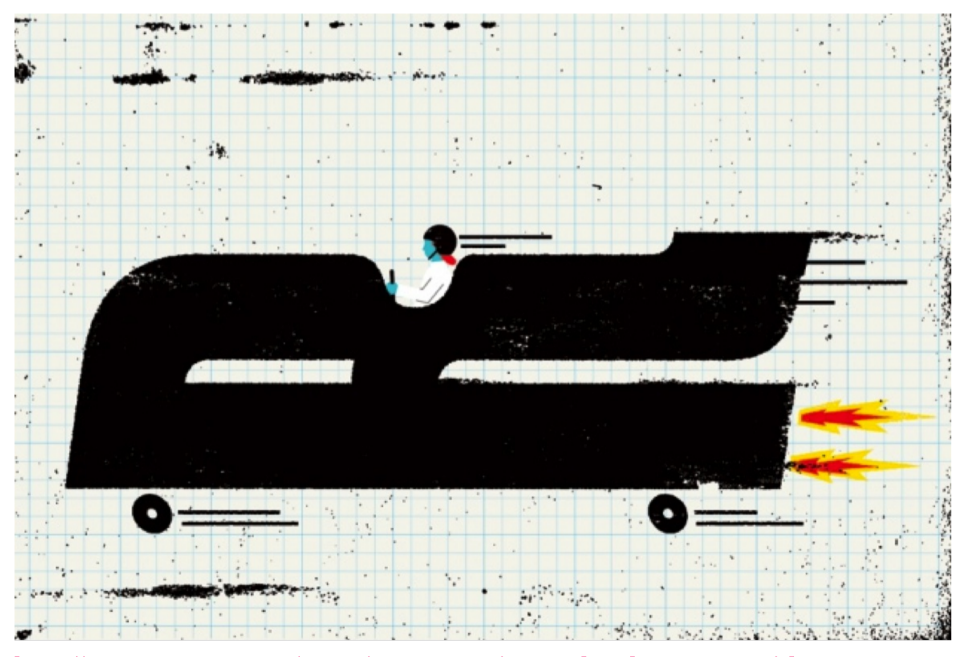
- help(nchar)
- ?nchar
- ??colsum

#### R Reference Card

- stack overflow
- google

Learning R at swirlstats

# Adventure with R



http://www.nature.com/news/programming-tools-adventures-with-r-1.16609

#### **Contact information**

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