# Code\_Lib\_Import

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2025-01-26

# Getting the Data

Defining Workspace or directory

```
x <- getwd() # find working directory
## [1] "C:/Users/Owner/OneDrive/Documents/CompSci/CourseraDS/CourseraDataScience"
dir.create("testdir") # create a directory if doesn't exist, args: dir name, for nested recursive = tru
## Warning in dir.create("testdir"): 'testdir' already exists
setwd("testdir") # set working dir
file.create("mytest.R") # create file in wd
## [1] TRUE
file.exists("mytest.R") # check if file or directory exists in wd
## [1] TRUE
file.info("mytest.R") # file metadata, use $ operator to grab specific items
            size isdir mode
                                          mtime
## mytest.R 0 FALSE 666 2025-01-26 11:35:44 2025-01-26 11:35:44
                          atime exe
## mytest.R 2025-01-26 11:35:44 no
file.rename("mytest.R", "mytest2.R") # rename
## [1] TRUE
```

```
file.copy("mytest2.R","mytest3.R") # copy file
## [1] FALSE
file.remove("mytest2.R") # remove file
## [1] TRUE
file.path("mytest3.R") # relative path
## [1] "mytest3.R"
setwd(x) # could use relative setwd("../") to move up one, setwd("./data"), or absolute path directly u
dir() # output files in directory. Also list.files()
## [1] "API_auth_ex.R"
                                     "Code-Library.pdf"
## [3] "Code Library.Rmd"
                                     "Code_Lib_Import.Rmd"
## [5] "coded.R"
                                     "complete.R"
## [7] "corr.R"
                                     "Course-Notes.pdf"
## [9] "Course Notes.Rmd"
                                     "CourseraDataScience.Rproj"
## [11] "example.h5"
                                     "getdata_data_ss06pid.csv"
## [13] "getdata_wksst8110.for"
                                     "hw1_data.csv"
## [15] "pollutantmean.R"
                                     "Programming 2.3"
## [17] "quiz_data_3.1.2.xlsx"
                                     "quiz_data_3.1.csv"
## [19] "Rprof.out"
                                     "rprog_data_ProgAssignment3"
## [21] "specdata"
                                     "specdata.zip"
## [23] "testdir"
files_full <- list.files("specdata", full.names=TRUE) # pull all file names from a directory
ls() # prints the objects in work space
## [1] "files_full" "x"
rm(list=ls()) # clear workspace
rm(list=setdiff(ls(), "x")) # clear workspace except x
version #R info version
##
                  x86_64-w64-mingw32
## platform
## arch
                  x86_64
## os
                  mingw32
## crt
                  ucrt
## system
                  x86_64, mingw32
## status
## major
                  4.1
## minor
## year
                  2024
## month
                  06
```

```
## day
                  14
                  86737
## svn rev
## language
## version.string R version 4.4.1 (2024-06-14 ucrt)
## nickname
                 Race for Your Life
sessionInfo() #R info version, packages
## R version 4.4.1 (2024-06-14 ucrt)
## Platform: x86_64-w64-mingw32/x64
## Running under: Windows 11 x64 (build 22631)
## Matrix products: default
##
##
## locale:
## [1] LC_COLLATE=English_United States.utf8
## [2] LC_CTYPE=English_United States.utf8
## [3] LC_MONETARY=English_United States.utf8
## [4] LC_NUMERIC=C
## [5] LC_TIME=English_United States.utf8
##
## time zone: America/New York
## tzcode source: internal
## attached base packages:
## [1] stats
                graphics grDevices utils
                                               datasets methods
                                                                   base
## loaded via a namespace (and not attached):
## [1] compiler_4.4.1
                        fastmap_1.2.0
                                                              tools_4.4.1
                                            cli_3.6.3
## [5] htmltools_0.5.8.1 rstudioapi_0.17.1 yaml_2.3.10
                                                              rmarkdown_2.29
## [9] knitr_1.49
                          xfun_0.49
                                            digest_0.6.37
                                                              rlang_1.1.4
## [13] evaluate_1.0.3
source("coded.R") # load code into console
## [1] "Hello World"
args(ls()) # get arguments for a function
## NULL
help(ls) # access documentation on ls() function
## starting httpd help server ... done
?ls # same. for operator use ?`:`
# Interface to outside world
file(description = "hw1_data.csv") # open connection to standard, uncompressed file. Helps for partial
```

```
## description "hw1_data.csv"
## class
             "file"
## mode
              "r"
## text
              "text"
              "closed"
## opened
              "yes"
## can read
              "yes"
## can write
# gzfile() # connection to file w compression gzip
# bzfile() # connection to file w compression bzip2
jh <- url("http://www.jhsph.edu", "r") # connection to webpage
close(jh) # to end connection
# Download data from website
fileUrl <- "https://stg-arcgisazurecdataprod1.az.arcgis.com/exportfiles-1506-24014/Part1_Crime_Beta_596
if(!file.exists("./testdir/crimedata.csv")) download.file(fileUrl, destfile = "./testdir/crimedata.csv"
dateDownloaded <- date() # Keep track of date downloaded for file</pre>
CSV and XLSX
# Read table or csv data into R
x <- read.table("hw1_data.csv", header = TRUE, sep = ",") # reading tabular data from text files, retur
head(x)
    Ozone Solar.R Wind Temp Month Day
## 1
       41
             190 7.4 67
                                5
                                   1
## 2
       36
             118 8.0 72
                                   2
                                   3
             149 12.6 74
## 3
       12
                                5
## 4
       18
             313 11.5 62
                                5
                                   4
## 5
       NA
              NA 14.3 56
                               5 5
## 6
       28
              NA 14.9
                        66
                                5
x <- read.csv("hw1_data.csv") # Same but default separator is ", " and header = TRUE
# write.table(x)
# help read.table with colClasses with smaller sample
initial <- read.table("hw1_data.csv", header = TRUE, sep = ",", nrows = 100)</pre>
classes <- sapply(initial, class)</pre>
tabAll <- read.table("hw1_data.csv", header = TRUE, sep = ",", colClasses = classes)
\# Read Excel file into R
x <- "https://stg-arcgisazurecdataprod1.az.arcgis.com/exportfiles-1506-24014/Part1 Crime Beta 383480701
if(!file.exists("./testdir/crimexl.xlsx")) {
 download.file(x, destfile = "./testdir/crimexl.xlsx", method = "curl") # improve
 dateXl <- date()</pre>
```

## Warning: package 'readxl' was built under R version 4.4.2

library(readxl) # xlsx, XLConnect other options

## A connection with

```
cameraData <- read_excel("./testdir/crimexl.xlsx", sheet = 1, col_types = "text", n_max = 100)</pre>
head(cameraData)
## # A tibble: 6 x 23
     RowID CCNumber CrimeDateTime CrimeCode Description Inside_Outside Weapon Post
     <chr> <chr>
                                   <chr>
                                             <chr>
                                                          <chr>
                                                                         <chr>
                    <chr>>
                                                                                <chr>>
                                             BURGLARY
                                                          <NA>
                                                                                835
## 1 1
           14I10336 41880.5
                                   5A
                                                                         NA
## 2 2
           14H13075 41880.989583~ 4E
                                             COMMON ASS~ <NA>
                                                                         NA
                                                                                515
           14H13066 41880.965277~ 6G
## 3 3
                                             LARCENY
                                                          < NA >
                                                                         NA
                                                                                114
## 4 4
           14I00593 41880.291666~ 6J
                                             LARCENY
                                                          <NA>
                                                                         NΑ
                                                                                732
## 5 5
           14H13360 41880.979166~ 6G
                                             LARCENY
                                                          <NA>
                                                                         NA
                                                                                213
## 6 6
           14H12804 41880.475694~ 4E
                                             COMMON ASS~ <NA>
                                                                         NA
                                                                                412
## # i 15 more variables: Gender <chr>, Age <chr>, Race <chr>, Ethnicity <chr>,
      Location <chr>, Old_District <chr>, New_District <chr>, Neighborhood <chr>,
       Latitude <chr>, Longitude <chr>, GeoLocation <chr>, PremiseType <chr>,
       Total_Incidents <chr>, x <chr>, y <chr>
## #
# write also works
```

# Read text file and R objects

# serialize()

```
lines <- readLines("coded.R") # reading lines of text file, return character vector

## Warning in readLines("coded.R"): incomplete final line found on 'coded.R'

## coded.R

## coded.R

# editable textual format retains metadata, helpful for version control, corruption fixable, memory cos dget("coded.R") # reading R objects deparsed into text files

## [1] "Hello World"

dput("coded.R") # takes R object, create R code to reconstruct object saving attributes, names

## "coded.R"

source("coded.R") # reading in R code files

## [1] "Hello World"

#dump() # multiple R objects

# load() # read in saved workspace read binary objects into R

# save()

# unservalize() # read single R objects in binary form
```

#### Just XML stuff

- XML (extensible markup language)
- http://en.wikipedia.org/wiki/XML
- store structured data in internet applications. From web scraping, API
- Components: markup (labels for structure) and content (text)
- Tags (labels): start end empty
- Elements: example of tag Hello, world
- Attributes: component of label
- XPath language: http://www.stat.berkeley.edu/~statcur/Workshop2/Presentations/XML.pdf
- /node (top level node), //node (node any level), node[@attr-name] node w attr name, node[@attr-name='bob']
- $\bullet \;\; \mathrm{XML} \;\; \mathrm{Short} \;\; \mathrm{Intro:} \;\; \mathrm{https://www.omegahat.net/RSXML/shortIntro.pdf}$
- XML Long Intro: https://www.omegahat.net/RSXML/Tour.pdf

```
# Reading file
library(XML)

## Warning: package 'XML' was built under R version 4.4.2

library(httr)
```

## Warning: package 'httr' was built under R version 4.4.2

```
fileUrl <- "https://www.w3schools.com/xml/simple.xml"
doc <- xmlTreeParse(rawToChar(GET(fileUrl)$content), useInternalNodes = TRUE) # read file into R to pars
rootNode <- xmlRoot(doc) # wrapper for structured element

# Access Data
xmlName(rootNode) # name of xml</pre>
```

## [1] "breakfast\_menu"

```
names(rootNode) # names of root node
           food
##
     food
                   food
                         food
                                 food
## "food" "food" "food" "food"
rootNode[[1]] # first food element, get first subelement of element using [[1]][[1]]
## <food>
##
     <name>Belgian Waffles</name>
##
     <price>$5.95</price>
     <description>Two of our famous Belgian Waffles with plenty of real maple syrup/description>
##
     <calories>650</calories>
## </food>
xmlSApply(rootNode,xmlValue) # give xml value of each root node
##
##
                                 "Belgian Waffles$5.95Two of our famous Belgian Waffles with plenty of
##
                      "Strawberry Belgian Waffles$7.95Light Belgian waffles covered with strawberries a
##
##
## "Berry-Berry Belgian Waffles$8.95Light Belgian waffles covered with an assortment of fresh berries a
##
                                                   "French Toast$4.50Thick slices made from our homemade
##
##
##
                           "Homestyle Breakfast$6.95Two eggs, bacon or sausage, toast, and our ever-pop
xpathSApply(rootNode, "//name", xmlValue) # get node xmlValue for each names
## [1] "Belgian Waffles"
                                     "Strawberry Belgian Waffles"
## [3] "Berry-Berry Belgian Waffles" "French Toast"
## [5] "Homestyle Breakfast"
xpathSApply(rootNode,"//price",xmlValue)
## [1] "$5.95" "$7.95" "$8.95" "$4.50" "$6.95"
# Pokemon
library(rvest)
## Warning: package 'rvest' was built under R version 4.4.2
fileUrlBase <- "https://pokemondb.net/pokedex/stats/gen"</pre>
generations <- 1:8
# Scrape Data
pokemonList <- lapply(generations,function(x){</pre>
 html <- read_html(pasteO(fileUrlBase,x)) # create URL</pre>
 tableNode <- html_node(html,xpath="//*[(@id = 'pokedex')]") # read HTML, extract node using xpath sel
 data <- html_table(html,header=TRUE)[[1]] # convert to data frame</pre>
```

```
colnames(data)[1] <- "ID"</pre>
  data$Generation <- x # add generation ID
  data # return data frame to list
})
thePokemon <- do.call(rbind,pokemonList) # one data frame
types <- strsplit(gsub('[a-z])',' ',thePokemon$Type, perl=T)," ") # split Type column
the Pokemon $Type1 <- unlist(lapply(types,function(x) x[1])) # Type1 column
thePokemon$Type2 <- unlist(lapply(types,function(x) ifelse(is.na(x[2])," ",x[2]))) # Type2 column
pokemonNames <- strsplit(gsub('[a-z]\\K(?=[A-Z])', '--', thePokemon$Name, perl=T),"--") # split Name
thePokemon$Name <- unlist(lapply(pokemonNames,function(x) x[1]))
thePokemon$Form <- unlist(lapply(pokemonNames,function(x) ifelse(is.na(x[2])," ",x[2])))
the Pokemon <- the Pokemon [,c(1,2,14,12,13,4,5,6,7,8,9,10,11)] # reorder and remove unneccesary
thePokemon[29, "Name"] <- "Nidoran" # cleaning gender
thePokemon[29,"Form"] <- "Female"</pre>
thePokemon[32,"Name"] <- "Nidoran"</pre>
thePokemon[32, "Form"] <- "Male"</pre>
```

#### Just JSON Stuff

- Javascript Object Notation- light weight data storage, common format from APIs, similar to XML but different syntax/format.
- Data stored as: number(double), strings(double quoted), boolean, array(ordered, comma sep, [] enclosed), objects(unordered, comma sep, {} enclosed, key:value pairs).
- http://en.wikipedia.org/wiki/JSON
- http://www.json.org/
- jsonlite vignette
- http://www.r-bloggers.com/new-package-jsonlite-a-smarter-json-encoderdecoder/

```
## Warning: package 'jsonlite' was built under R version 4.4.2

jsonData <- fromJSON("https://api.github.com/users/jtleek/repos") # can pass any format
names(jsonData) # names of data frame</pre>
```

```
[1] "id"
##
                                        "node_id"
##
  [3] "name"
                                       "full_name"
   [5] "private"
                                        "owner"
   [7] "html_url"
                                       "description"
##
  [9] "fork"
                                       "url"
## [11] "forks url"
                                       "kevs url"
## [13] "collaborators_url"
                                       "teams_url"
## [15] "hooks_url"
                                        "issue_events_url"
## [17] "events_url"
                                       "assignees_url"
## [19] "branches_url"
                                       "tags_url"
```

```
## [21] "blobs_url"
                                      "git_tags_url"
## [23] "git_refs_url"
                                      "trees_url"
## [25] "statuses_url"
                                      "languages_url"
## [27] "stargazers_url"
                                      "contributors_url"
## [29] "subscribers_url"
                                      "subscription_url"
## [31] "commits_url"
                                      "git_commits_url"
## [33] "comments_url"
                                      "issue_comment_url"
## [35] "contents_url"
                                      "compare_url"
## [37] "merges_url"
                                      "archive_url"
## [39] "downloads_url"
                                      "issues_url"
## [41] "pulls_url"
                                      "milestones_url"
## [43] "notifications_url"
                                      "labels_url"
## [45] "releases_url"
                                      "deployments_url"
## [47] "created_at"
                                      "updated_at"
## [49] "pushed_at"
                                       "git_url"
## [51] "ssh_url"
                                      "clone_url"
## [53] "svn_url"
                                      "homepage"
## [55] "size"
                                      "stargazers_count"
## [57] "watchers_count"
                                      "language"
## [59] "has_issues"
                                      "has_projects"
## [61] "has_downloads"
                                      "has_wiki"
## [63] "has_pages"
                                      "has_discussions"
## [65] "forks_count"
                                      "mirror_url"
## [67] "archived"
                                      "disabled"
## [69] "open_issues_count"
                                      "license"
## [71] "allow_forking"
                                      "is_template"
## [73] "web_commit_signoff_required" "topics"
## [75] "visibility"
                                      "forks"
## [77] "open_issues"
                                      "watchers"
## [79] "default_branch"
names(jsonData$owner) # names of column
                              "id"
                                                     "node_id"
## [1] "login"
## [4] "avatar url"
                                                     "url"
                              "gravatar_id"
## [7] "html_url"
                              "followers_url"
                                                    "following_url"
## [10] "gists_url"
                              "starred_url"
                                                     "subscriptions_url"
## [13] "organizations_url"
                              "repos_url"
                                                     "events_url"
## [16] "received_events_url" "type"
                                                    "user_view_type"
## [19] "site_admin"
jsonData$owner$login # who logged in
## [1] "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek"
## [9] "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek"
## [17] "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek"
## [25] "jtleek" "jtleek" "jtleek" "jtleek" "jtleek" "jtleek"
# write data frame to JSON
myjson <- toJSON(iris, pretty = TRUE)</pre>
cat(myjson)
```

```
## [
##
       "Sepal.Length": 5.1,
##
##
       "Sepal.Width": 3.5,
##
       "Petal.Length": 1.4,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
     },
##
     {
##
       "Sepal.Length": 4.9,
##
       "Sepal.Width": 3,
##
       "Petal.Length": 1.4,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
     },
##
##
       "Sepal.Length": 4.7,
       "Sepal.Width": 3.2,
##
       "Petal.Length": 1.3,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
##
##
##
       "Sepal.Length": 4.6,
##
       "Sepal.Width": 3.1,
##
       "Petal.Length": 1.5,
       "Petal.Width": 0.2,
##
##
       "Species": "setosa"
##
     },
##
     {
##
       "Sepal.Length": 5,
##
       "Sepal.Width": 3.6,
##
       "Petal.Length": 1.4,
##
       "Petal.Width": 0.2,
       "Species": "setosa"
##
##
     },
##
##
       "Sepal.Length": 5.4,
       "Sepal.Width": 3.9,
##
##
       "Petal.Length": 1.7,
##
       "Petal.Width": 0.4,
       "Species": "setosa"
##
##
     },
##
##
       "Sepal.Length": 4.6,
##
       "Sepal.Width": 3.4,
##
       "Petal.Length": 1.4,
##
       "Petal.Width": 0.3,
##
       "Species": "setosa"
##
     },
##
##
       "Sepal.Length": 5,
       "Sepal.Width": 3.4,
##
##
       "Petal.Length": 1.5,
```

```
"Petal.Width": 0.2,
##
##
       "Species": "setosa"
##
     },
##
     {
       "Sepal.Length": 4.4,
##
##
       "Sepal.Width": 2.9,
##
       "Petal.Length": 1.4,
       "Petal.Width": 0.2,
##
##
       "Species": "setosa"
##
     },
##
     {
##
       "Sepal.Length": 4.9,
##
       "Sepal.Width": 3.1,
##
       "Petal.Length": 1.5,
##
       "Petal.Width": 0.1,
       "Species": "setosa"
##
##
     },
##
       "Sepal.Length": 5.4,
##
       "Sepal.Width": 3.7,
##
##
       "Petal.Length": 1.5,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
     },
##
     {
##
       "Sepal.Length": 4.8,
##
       "Sepal.Width": 3.4,
##
       "Petal.Length": 1.6,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
     },
##
     {
##
       "Sepal.Length": 4.8,
##
       "Sepal.Width": 3,
       "Petal.Length": 1.4,
##
##
       "Petal.Width": 0.1,
##
       "Species": "setosa"
##
     },
##
##
       "Sepal.Length": 4.3,
##
       "Sepal.Width": 3,
##
       "Petal.Length": 1.1,
##
       "Petal.Width": 0.1,
##
       "Species": "setosa"
##
     },
##
##
       "Sepal.Length": 5.8,
##
       "Sepal.Width": 4,
##
       "Petal.Length": 1.2,
##
       "Petal.Width": 0.2,
       "Species": "setosa"
##
##
     },
##
     {
##
       "Sepal.Length": 5.7,
```

```
"Sepal.Width": 4.4,
##
##
       "Petal.Length": 1.5,
       "Petal.Width": 0.4,
##
##
       "Species": "setosa"
##
     },
##
       "Sepal.Length": 5.4,
##
##
       "Sepal.Width": 3.9,
##
       "Petal.Length": 1.3,
##
       "Petal.Width": 0.4,
       "Species": "setosa"
##
##
     },
##
##
       "Sepal.Length": 5.1,
##
       "Sepal.Width": 3.5,
       "Petal.Length": 1.4,
##
##
       "Petal.Width": 0.3,
       "Species": "setosa"
##
##
     },
##
       "Sepal.Length": 5.7,
##
##
       "Sepal.Width": 3.8,
##
       "Petal.Length": 1.7,
       "Petal.Width": 0.3,
##
##
       "Species": "setosa"
##
     },
##
       "Sepal.Length": 5.1,
##
       "Sepal.Width": 3.8,
##
##
       "Petal.Length": 1.5,
       "Petal.Width": 0.3,
##
##
       "Species": "setosa"
##
     },
##
       "Sepal.Length": 5.4,
##
       "Sepal.Width": 3.4,
##
       "Petal.Length": 1.7,
##
##
       "Petal.Width": 0.2,
       "Species": "setosa"
##
##
     },
##
##
       "Sepal.Length": 5.1,
       "Sepal.Width": 3.7,
##
##
       "Petal.Length": 1.5,
       "Petal.Width": 0.4,
##
       "Species": "setosa"
##
##
     },
     {
##
##
       "Sepal.Length": 4.6,
##
       "Sepal.Width": 3.6,
       "Petal.Length": 1,
##
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
     },
```

```
##
##
       "Sepal.Length": 5.1,
       "Sepal.Width": 3.3,
##
##
       "Petal.Length": 1.7,
       "Petal.Width": 0.5,
##
##
       "Species": "setosa"
##
     },
##
##
       "Sepal.Length": 4.8,
##
       "Sepal.Width": 3.4,
##
       "Petal.Length": 1.9,
##
       "Petal.Width": 0.2,
##
       "Species": "setosa"
##
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##
##
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##
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##
##
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##
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##
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##
     {
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##
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##
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##
       "Petal.Width": 1.5,
##
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##
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     {
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##
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##
##
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##
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##
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##
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##
##
       "Petal.Width": 2,
##
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##
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##
##
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##
       "Sepal.Width": 2.8,
##
       "Petal.Length": 6.7,
##
       "Petal.Width": 2,
##
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##
     },
##
     {
##
       "Sepal.Length": 6.3,
```

```
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##
##
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       "Petal.Width": 1.8,
##
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##
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##
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##
##
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##
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##
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##
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##
##
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##
       "Sepal.Width": 3.2,
##
       "Petal.Length": 6,
##
       "Petal.Width": 1.8,
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##
##
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##
##
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##
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##
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##
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##
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##
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##
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       "Petal.Width": 1.8,
##
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##
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##
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##
       "Sepal.Width": 2.8,
##
##
       "Petal.Length": 5.6,
##
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##
##
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##
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       "Sepal.Width": 3,
##
##
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       "Petal.Width": 1.6,
##
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     {
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##
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##
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##
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```

```
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##
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##
##
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##
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##
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##
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##
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##
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##
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##
       "Petal.Width": 2.3,
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##
     },
##
##
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##
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##
##
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##
     },
##
##
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##
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##
       "Petal.Length": 5.5,
##
       "Petal.Width": 1.8,
##
       "Species": "virginica"
##
     },
##
##
       "Sepal.Length": 6,
##
       "Sepal.Width": 3,
##
       "Petal.Length": 4.8,
##
       "Petal.Width": 1.8,
```

```
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##
##
     },
##
##
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##
##
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##
##
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##
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##
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##
##
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##
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##
     },
##
##
       "Sepal.Length": 5.8,
##
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##
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##
       "Petal.Width": 1.9,
##
       "Species": "virginica"
##
     },
##
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##
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##
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       "Petal.Width": 2.3,
##
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##
##
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##
##
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##
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##
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##
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##
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##
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##
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##
       "Petal.Width": 2.3,
       "Species": "virginica"
##
##
     },
##
##
       "Sepal.Length": 6.3,
##
       "Sepal.Width": 2.5,
```

```
##
       "Petal.Length": 5,
##
       "Petal.Width": 1.9,
       "Species": "virginica"
##
##
     },
##
##
       "Sepal.Length": 6.5,
##
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       "Petal.Length": 5.2,
##
##
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##
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##
     },
##
##
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##
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##
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##
       "Petal.Width": 2.3,
##
        "Species": "virginica"
##
     },
##
       "Sepal.Length": 5.9,
##
##
       "Sepal.Width": 3,
##
       "Petal.Length": 5.1,
       "Petal.Width": 1.8,
##
##
        "Species": "virginica"
##
     }
## ]
# upload again
iris2 <- fromJSON(myjson)</pre>
head(iris2)
```

```
Sepal.Length Sepal.Width Petal.Length Petal.Width Species
##
## 1
              5.1
                           3.5
                                         1.4
                                                      0.2
                                                           setosa
## 2
              4.9
                           3.0
                                         1.4
                                                      0.2
                                                          setosa
## 3
              4.7
                           3.2
                                         1.3
                                                     0.2 setosa
## 4
              4.6
                           3.1
                                         1.5
                                                      0.2
                                                          setosa
## 5
              5.0
                           3.6
                                         1.4
                                                      0.2 setosa
## 6
              5.4
                           3.9
                                         1.7
                                                      0.4 setosa
```

## mySQL Stuff

- Free, widely-used open source database, esp in internet-based applications.
- Data structured in databases, inside is tables(dataset), inside is fields(column), row is record. IDs link databases.
- http://en.wikipedia.org/wiki/MySQL and http://www.mysql.com/
- Install MySQL: https://dev.mysql.com/doc/refman/5.7/en/installing.html
- http://cran.r-project.org/web/packages/RMySQL/RMySQL.pdf

- Commands: http://www.pantz.org/software/mysql/mysqlcommands.html and http://www.r-bloggers.com/mysql-and-r
- Be careful with SQL commands, DO NOT PUSH UNLESS REQUIRED!

#### library (RMySQL)

```
## Warning: package 'RMySQL' was built under R version 4.4.2

## Loading required package: DBI

## Warning: package 'DBI' was built under R version 4.4.2

ucscDb <- dbConnect(MySQL(), user="genome",host="genome-mysql.cse.ucsc.edu") # open a connection to the result <- dbGetQuery(ucscDb, "show databases;"); dbDisconnect(ucscDb); # run the mySQL command "show databases"</pre>
```

#### ## [1] TRUE

#### result # print available databases

##		Database
##	1	acaChl1
##	2	ailMel1
##	3	allMis1
##	4	allSin1
##	5	amaVit1
##	6	anaPla1
##	7	ancCey1
##	8	angJap1
##	9	anoCar1
##	10	anoCar2
##	11	anoGam1
##	12	anoGam3
##	13	apaSpi1
##	14	apaVit1
##	15	apiMel1
		-
##	16	apiMel2
##	17	apiMel2 apiMel2_justpushed
## ##	17 18	apiMel2 apiMel2_justpushed aplCal1
## ## ##	17 18 19	apiMel2 apiMel2_justpushed aplCal1 aptFor1
## ## ## ##	17 18 19 20	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1
## ## ## ## ##	17 18 19 20 21	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2
## ## ## ## ##	17 18 19 20 21 22	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1
## ## ## ## ##	17 18 19 20 21 22 23	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1
## ## ## ## ## ##	17 18 19 20 21 22 23 24	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1
## ## ## ## ## ## ##	17 18 19 20 21 22 23 24 25	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1 balPav1
## ## ## ## ## ## ##	17 18 19 20 21 22 23 24 25 26	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1 balPav1 bisBis1
## ## ## ## ## ## ## ##	17 18 19 20 21 22 23 24 25 26 27	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1 balPav1 bisBis1 bosTau2
## ## ## ## ## ## ## ## ## ## ## ## ##	17 18 19 20 21 22 23 24 25 26 27 28	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1 balPav1 bisBis1 bosTau2 bosTau3
## ## ## ## ## ## ## ##	17 18 19 20 21 22 23 24 25 26 27	apiMel2 apiMel2_justpushed aplCal1 aptFor1 aptMan1 aquChr2 araMac1 ascSuu1 balAcu1 balPav1 bisBis1 bosTau2

## 31	bosTau6
## 32	bosTau7
## 33	bosTau8
## 34	bosTau9
## 35	bosTauMd3
## 36	braFlo1
## 37	bruMal2
## 38	bucRhi1
## 39	burXyl1
## 40	caeAng2
## 41	caeJap1
## 42	caeJap4
## 43	caePb1
## 44	caePb2
## 45	caePb3
## 46	caeRem2
## 47	caeRem3
## 48	caeRem4
## 49	caeSp111
## 50	caeSp51
## 51	calAnn1
## 52	calJac1
## 53	calJac3
## 54	calJac4
## 55	calMil1
## 56	canFam1
## 57	canFam2
## 58	canFam3
## 59	canFam4
## 60	canFam5
## 61	canFam6
## 62	capCar1
## 63	carCri1
## 64	cavPor3
## 65	cb1
## 66	cb3
## 67	cb4
## 68	ce10
## 69	ce11
## 70	ce2
## 71	ce4
## 72	ce6
## 73	cerSim1
## 74	chaVoc2
## 75	cheMyd1
## 76	chlSab2
## 70 ## 77	chlUnd1
	choHof1
	chrPic1
## 80	chrPic2
## 81	ci1
## 82	ci2
## 83	ci3
## 84	colLiv1

##	85	colStr1
##	86	corBra1
##	87	corCor1
##	88	cotJap2
##	89	criGri1
##	90	criGriChoV1
##	91	criGriChoV2
##	92	cucCan1
##	93	danRer1
##	94	danRer10
##	95	danRer11
##	96	danRer2
##	97	danRer3
##	98	danRer4
##	99	danRer5
##	100	danRer6
##	101	danRer7
##	102	dasNov3
##	103	dipOrd1
##	104	dirImm1
##	105	dm1
##	106	dm2
##	107	dm3
##	108	dm6
##	109	dp2
##	110	dp3
##	111	droAna1
##	112	droAna2
##	113	droEre1
##	114	droGri1
##	115	droMoj1
##	116	droMoj2
##	117	droPer1
##	118	droSec1
##	119	droSim1
##	120	droSim2
##	121	droVir1
##	122	droVir2
##	123	droYak1
##	124	droYak2
##	125	eboVir3
##	126	echTel1
##	127	echTel2
##	128	egrGar1
##	129	enhLutNer1
##	130	equCab1
##	131	equCab2
##	132	equCab3
##	133	eriEur1
##	134	eriEur2
##	135	eurHel1
##	136	falChe1
##	137	falPer1
##	138	felCat3
##	100	1610919

##	139	felCat4
##	140	felCat5
##	141	felCat8
##	142	felCat9
##	143	ficAlb2
##	144	fr1
##	145	fr2
##	146	fr3
##	147	fulGla1
##	148	gadMor1
##	149	galGal2
	150	_
##		galGal3
##	151	galGal4
##	152	galGal5
##	153	galGal6
##	154	galVar1
##	155	gasAcu1
##	156	gavSte1
##	157	gbMeta
##	158	geoFor1
##	159	go
##	160	go080130
##	161	go140213
##	162	go150121
##	163	go180426
##	164	gorGor3
##	165	gorGor4
##	166	gorGor5
##	167	gorGor6
##	168	haeCon2
##	169	halAlb1
##	170	halLeu1
##	171	hetBac1
##	172	hetGla1
##	173	hetGla2
##	174	hg16
##	175	hg17
##	176	hg18
##	177	hg19
##	178	hg19Patch10
##	179	hg19Patch13
##	180	hg19_justpushed
##	181	hg38
##	182	hg38Patch11
##	183	hg38_justpushed
##	184	hgFixed
##	185	hgcentral
	186	<del>-</del>
##		hs1
##	187	information_schema
##	188	knownGeneKentHg19
##	189	knownGeneV39
##	190	knownGeneV43
##	191	knownGeneV44
##	192	knownGeneV45

##	193	knownGeneV46
##	194	knownGeneVM27
##	195	knownGeneVM30
##	196	knownGeneVM32
##	197	latCha1
##	198	lepDis1
##	199	letCam1
##	200	loaLoa1
##	201	loxAfr3
##	202	macEug1
##	203	macEug2
##	204	macFas5
##	205	manPen1
##	206	melGal1
##	207	melGal5
##	208	melHap1
##	209	melInc2
##	210	melUnd1
##	211	merNub1
##		mesUni1
##		micMur1
##		micMur2
##		mm10
##		mm39
##		mm39_justpushed
##		mm5
##	219	mm6
##	220	mm7
##	221	mm8
##		mm9
##		monDom1
##		monDom1
##		monDom4
##		mpxvRivers
##		mpxviiiveis musFur1
##	228	musruri myoLuc2
##	229	nanPar1
##	230	nasLar1
		nastari necAme1
##		neoSch1
##		
##		nipNip1
##		nomLeu1
##		nomLeu2 nomLeu3
##		
##	237	ochPri2
##		ochPri3
##		oncVol1
##		opiHoa1
##		oreNil1
##		oreNil2
##		oreNil3
##		ornAna1
##		ornAna2
##	246	oryCun2

##	247	oryLat2
##	248	otoGar3
##	249	oviAri1
##	250	oviAri3
##	251	oviAri4
##	252	panPan1
##	253	panPan2
##	254	panPan3
##	255	panRed1
##	256	panTro1
##	257	panTro2
##	258	panTro3
##	259	panTro4
##	260	panTro5
##	261	panTro6
##	262	papAnu2
##	263	papAnu4
##	264	papHam1
##	265	pelCri1
##	266	pelSin1
##	267	performance_schema
##		petMar1
##	269	petMar2
##	270	petMar3
##	271	phaCar1
##	272	phaLep1
##	273	phoRub1
##	274	picPub1
##	275	ponAbe2
##	276	ponAbe3
##	277	priExs1
##	278	priPac1
##	279	priPac3
##	280	proCap1
##	281	proteins120806
##	282	proteins121210
##	283	proteins140122
##	284	proteins150225
##	285	proteins160229
##	286	proteins180404
##	287	proteome
##	288	pteGut1
##	289	pteVam1
##	290	pygAde1
##	291	pytBiv1
##	292	rheMac1
##	293	rheMac10
##	294	rheMac2
##	295	rheMac3
##	296	rheMac8
##	297	rhiRox1
##	298	rn3
##	299	rn4
##	300	rn5

"" 004	
## 301	rn6
## 302	rn7
## 303 ## 304	sacCer1
	sacCer2
## 305 ## 306	sacCer3
## 307	saiBol1 sarHar1
## 308	sarnari serCan1
## 309	sorAra1
## 310	sorAra2
## 311	sp120323
## 312	sp120323
## 313	sp121210
## 314	sp140122
## 315	sp160229
## 316	sp180404
## 317	spico404 speTri2
## 318	strCam1
## 319	strPur1
## 320	strPur2
## 321	strRat2
## 322	susScr11
## 323	susScr2
## 324	susScr3
## 325	taeGut1
## 326	taeGut2
## 327	tarSyr1
## 328	tarSyr2
## 329	tauEry1
## 330	tetNig1
## 331	tetNig2
## 332	thaSir1
## 333	tinGut2
## 334	triMan1
## 335	triSpi1
## 336	triSui1
## 337	tupBel1
## 338	turTru2
## 339	tytAlb1
## 340	uniProt
## 341	vicPac1
## 342	vicPac2
## 343	visiGene
## 344	wuhCor1
## 345	xenLae2
## 346	xenTro1
## 347	xenTro10
## 348	xenTro2
## 349	xenTro3
## 350	xenTro7
## 351	xenTro9
## 352	zonAlb1

```
hg19 <- dbConnect(MySQL(), user="genome", db = "hg19", host="genome-mysql.cse.ucsc.edu") # connection t
allTables <- dbListTables(hg19) # extract names of all tables
length(allTables)
## [1] 12690
allTables[1:5]
## [1] "HInv"
                      "HInvGeneMrna" "acembly"
                                                     "acemblyClass" "acemblyPep"
dbListFields(hg19, "affyU133Plus2") # fields/columns of table affyU133Plus2 (is a microarray)
## [1] "bin"
                      "matches"
                                     "misMatches"
                                                   "repMatches"
                                                                 "nCount"
## [6] "qNumInsert"
                      "qBaseInsert" "tNumInsert"
                                                   "tBaseInsert" "strand"
## [11] "qName"
                      "qSize"
                                     "qStart"
                                                   "qEnd"
                                                                  "tName"
## [16] "tSize"
                      "tStart"
                                     "tEnd"
                                                   "blockCount"
                                                                 "blockSizes"
                      "tStarts"
## [21] "qStarts"
dbGetQuery(hg19, "select count(*) from affyU133Plus2") # how many rows/record. Pass mySQL command to co
##
     count(*)
## 1
        58463
library(knitr) # to supress warnings
## Warning: package 'knitr' was built under R version 4.4.2
suppressWarnings({ affyData <- dbReadTable(hg19, "affyU133Plus2") }) # import data from table in mySQL</pre>
head(affyData)
##
     bin matches misMatches repMatches nCount qNumInsert qBaseInsert tNumInsert
## 1 585
             530
                          4
                                      0
                                            23
                                                        3
## 2 585
            3355
                                           109
                                                                   67
                                                                                9
                         17
                                      0
                                                        9
## 3 585
            4156
                         14
                                      0
                                            83
                                                       16
                                                                   18
                                                                                2
                                                                   42
                                                                                3
## 4 585
            4667
                          9
                                      0
                                            68
                                                       21
## 5 585
            5180
                         14
                                      0
                                           167
                                                       10
                                                                   38
                                                                                1
## 6 585
             468
                          5
                                      0
                                            14
                                                        0
                                                                    0
     tBaseInsert strand
                               qName qSize qStart qEnd tName
##
                                                                  tSize tStart
## 1
             898
                         225995_x_at
                                       637
                                                 5 603 chr1 249250621
                                                                         14361
                                                                         14381
## 2
           11621
                         225035_x_at 3635
                                                 0 3548 chr1 249250621
## 3
              93
                         226340_x_at 4318
                                                 3 4274 chr1 249250621 14399
## 4
            5743
                      - 1557034_s_at 4834
                                                48 4834
                                                         chr1 249250621 14406
## 5
              29
                                                 0 5399 chr1 249250621 19688
                           231811_at 5399
## 6
               0
                           236841_at
                                       487
                                                   487 chr1 249250621 27542
      tEnd blockCount
##
## 1 15816
## 2 29483
                   17
## 3 18745
                   18
```

## 4 24893

23

```
## 5 25078
                   11
## 6 28029
                    1
##
                                                                       blockSizes
## 1
                                                                93,144,229,70,21,
## 2
                  73,375,71,165,303,360,198,661,201,1,260,250,74,73,98,155,163,
## 3
                      690, 10, 32, 33, 376, 4, 5, 15, 5, 11, 7, 41, 277, 859, 141, 51, 443, 1253,
## 4 99,352,286,24,49,14,6,5,8,149,14,44,98,12,10,355,837,59,8,1500,133,624,58,
## 5
                                             131,26,1300,6,4,11,4,7,358,3359,155,
## 6
                                                                              487,
##
## 1
                                                                                           34,132,278,541,
## 2
                             87,165,540,647,818,1123,1484,1682,2343,2545,2546,2808,3058,3133,3206,3317,3
## 3
                       44,735,746,779,813,1190,1195,1201,1217,1223,1235,1243,1285,1564,2423,2565,2617,3
## 4 0,99,452,739,764,814,829,836,842,851,1001,1016,1061,1160,1173,1184,1540,2381,2441,2450,3951,4103,4
## 5
                                                           0,132,159,1460,1467,1472,1484,1489,1497,1856,5
## 6
##
## 1
## 2
                                          14381,14454,14969,15075,15240,15543,15903,16104,16853,17054,17
## 3
                                    14399, 15089, 15099, 15131, 15164, 15540, 15544, 15549, 15564, 15569, 15580, 15
## 4 14406,20227,20579,20865,20889,20938,20952,20958,20963,20971,21120,21134,21178,21276,21288,21298,21
                                                                                19688,19819,19845,21145,21
## 5
## 6
suppressWarnings({ query <- dbSendQuery(hg19, "select * from affyU133Plus2 where misMatches between 1 a
affyMis <- fetch(query); quantile(affyMis$misMatches) # fetch subset from database and run quantile on
##
        25% 50%
                   75% 100%
     0%
##
      1
           1
                2
                     2
affyMisSmall <- fetch(query,n=10); dbClearResult(query); # fetch top 10 records
## [1] TRUE
dim(affyMisSmall)
## [1] 10 22
dbDisconnect(hg19) # CLOSE CONNECTION
## [1] TRUE
```

qSt.

## Just HDF5 Stuff

- Large data sets, store range of data types, Hierarchical Data Format.
- Groups of 0+ data sets and their metadata: group header (with name and list of attributes) and group symbol table (with list of objects in groups).

- Datasets are multi-dimensional array of data elements with their metadata: header (with name, datatype, dataspace, and storage layout) and data array (containing data).
- http://www.hdfgroup.org/
- Install: if (!require("BiocManager", quietly = TRUE)) install.packages("BiocManager"); BiocManager::install(version = "3.20"); BiocManager::install(pkgs=c("rhdf5"));
- https://www.bioconductor.org/packages/release/bioc/html/rhdf5.html

```
library(rhdf5)
## Warning: package 'rhdf5' was built under R version 4.4.2
file.remove("example.h5")
## [1] TRUE
created = h5createFile("example.h5") # create hdf5 file
created
## [1] TRUE
created = h5createGroup("example.h5", "foo") # greate group with given name inside h5
created = h5createGroup("example.h5","baa")
created = h5createGroup("example.h5", "foo/foobaa") # create subgroup
h5ls("example.h5") # list components of h5 file
                       otype dclass dim
##
     group
             name
              baa H5I GROUP
        /
              foo H5I_GROUP
## 1
         /
## 2 /foo foobaa H5I GROUP
A <- matrix(1:10,nrow=5,ncol=2)
h5write(A, "example.h5", "foo/A") # write object to group
B \leftarrow array(seq(0.1,2.0,by=0.1),dim=c(5,2,2))
attr(B, "scale") <- "liter" # can include metadata</pre>
h5write(B, "example.h5", "foo/foobaa/B")
h5ls("example.h5")
##
           group
                               otype dclass
                                                    dim
                    name
## 0
                           H5I GROUP
                    baa
## 1
               /
                           H5I_GROUP
                    foo
## 2
                       A H5I_DATASET INTEGER
            /foo
                                                  5 x 2
                           H5I_GROUP
## 3
            /foo foobaa
## 4 /foo/foobaa
                      B H5I_DATASET
                                       FLOAT 5 x 2 x 2
df <- data.frame(1L:5L,seq(0,1,length.out=5), c("ab","cde","fghi","a","s"), stringsAsFactors = FALSE)</pre>
h5write(df, "example.h5", "df") # write data set to top level group
h5ls("example.h5")
```

```
##
                                        dclass
                                                      dim
           group
                    name
                               otype
## 0
                           H5I_GROUP
                     baa
## 1
                      df H5I_DATASET COMPOUND
                                                        5
                           H5I_GROUP
## 2
                     foo
## 3
            /foo
                       A H5I_DATASET
                                       INTEGER
                                                    5 x 2
                           H5I GROUP
## 4
            /foo foobaa
                                         FLOAT 5 x 2 x 2
## 5 /foo/foobaa
                       B H5I_DATASET
h5read("example.h5", "foo/A") # read data from hf file
##
        [,1] [,2]
## [1,]
           1
## [2,]
                7
           2
## [3,]
           3
                8
## [4,]
           4
                9
## [5,]
h5write(c(12,13,14), "example.h5", "foo/A", index=list(1:3,1)) # can read and write (in this case edit) in
h5read("example.h5","foo/A")
```

```
##
         [,1] [,2]
## [1,]
           12
                  7
## [2,]
           13
## [3,]
           14
                  8
## [4,]
            4
                  9
## [5,]
            5
                 10
```

## HTML and Webscraping

- Webscraping: programatically extracting data from HTML code of websites. Be careful of Terms of Service and speed of scraping...
- http://en.wikipedia.org/wiki/Web\_scraping
- http://www.r-bloggers.com/?s=Web+Scraping
- http://cran.r-project.org/web/packages/httr/httr.pdf

```
con = url("https://scholar.google.com/citations?user=HI-I6COAAAAJ&hl=en") # open connection to website
htmlCode = readLines(con) # extract file lines into a char vector

## Warning in readLines(con): incomplete final line found on
## 'https://scholar.google.com/citations?user=HI-I6COAAAAJ&hl=en'

close(con) # REMEMBER
htmlCode[1] # Hard to read
```

## [1] "<!doctype html><html><head><title>Jeff Leek - Google Scholar</title><meta http-equiv=\"Content-

```
# alternate method: XML
library(XML)
url <- "http://web.archive.org/web/20130207021632/http://scholar.google.com:80/citations?user=HI-I6COAA
html <- htmlTreeParse(url,useInternalNodes = TRUE) # parsing file using internal nodes
xpathSApply(html, "//title", xmlValue) # use sApply for clean vector. Accessing title
## [1] "Jeff Leek - Google Scholar Citations"
xpathSApply(html, "//td[@id='col-citedby']", xmlValue) # accessing elements of table
   [1] "Cited by" "339"
                              "173"
                                          "140"
                                                     "133"
                                                                 "107"
                                                                 "10"
  [7] "95"
                   "79"
                               "79"
                                          "54"
                                                     "17"
##
                                          "8"
                                                                 "6"
                   "8"
                              "8"
                                                     "8"
## [13] "9"
## [19] "6"
                   "5"
                               "3"
# alternate method: httr and GET
library(httr)
html2 <- GET("https://scholar.google.com/citations?user=HI-I6COAAAAJ&hl=en")
content2 <- content(html2, as = "text") # extract content from HTML page as one text string</pre>
parsedHtml <- htmlParse(content2, asText = TRUE) # parse the text, same as xml package result
xpathSApply(parsedHtml, "//title", xmlValue)
## [1] "Jeff Leek - Google Scholar"
# passwords
url <- "http://httpbin.org/basic-auth/user/passwd"</pre>
pg1 <- GET(url)
pg1 # not authenticated, passes status 401
## Response [http://httpbin.org/basic-auth/user/passwd]
     Date: 2025-01-26 16:36
##
##
     Status: 401
##
     Content-Type: <unknown>
## <EMPTY BODY>
pg2 <- GET(url, authenticate("user", "passwd")) # sends authentication
pg2
## Response [http://httpbin.org/basic-auth/user/passwd]
##
    Date: 2025-01-26 16:36
     Status: 200
##
##
     Content-Type: application/json
     Size: 47 B
##
## {
##
     "authenticated": true,
##
     "user": "user"
## }
```

#### APIs

- Application programming interfaces.
- Developed platform will have GET request URLs (and parameters), which you use in httr to get data. Need a API/dev account, need to submit a request (for each project!). Receive: consumer key, consumer secret, request token URL, and Authorization URL.
- Able to GET, POST, PUT, DELETE with httr if authorized.
- Need oauth in most cases, sometimes username/password allowed.
- Sites: Facebook, Google, Twitter, GitHub...

```
# in personal vault :)
rm(list=ls())
```

#### Other Sources

Packages for Data Storage Mechanisms - Search Google: "data storage mechanism R package"

- ?connections to get info on creating connection (CLOSE)
- foreign package: loads data from Minitab, S, SAS, SPSS, Stata, Systat. read.lang. http://cran.r-project.org/web/packages/foreign/foreign.pdf
- RPostresSQL: https://code.google.com/p/rpostgresql/, http://cran.r-project.org/web/packages/RPostgreSQL/RPostgreSQL.pdf
- RODBC (PostgreQL, MySQL, Microsoft Access, SQLite): http://cran.r-project.org/web/packages/RODBC/vignettes/RODBC.pdf, http://cran.r-project.org/web/packages/RODBC/RODBC.pdf
- RMongo/rmongodb: http://cran.r-project.org/web/packages/RMongo/RMongo.pdf, http://www.r-bloggers.com/r-and-mongodb

Image Data - jpeg: http://cran.r-project.org/web/packages/jpeg/index.html

- readbitmap: http://cran.r-project.org/web/packages/readbitmap/index.html
- png: http://cran.r-project.org/web/packages/png/index.html
- EBImage (Bioconductor): http://www.bioconductor.org/packages/2.13/bioc/html/EBImage.html

 ${\it GIS~Data-raster:~http://cran.r-project.org/web/packages/raster/index.html}$ 

- 'sf' and 'terra'

Musical Data - tuneR: http://cran.r-project.org/web/packages/tuneR/

- seewave: http://rug.mnhn.fr/seewave