dplyr and pipes

Install the tidyverse package set

Most usefull packages for modern R code: https://www.tidyverse.org/packages/ For those who learn to code R before 2015: http://kbroman.org/hipsteR/ Installs ggplot2, dplyr, tidyr, readr, purr, tibble and much more: 'colorspace', 'mnormt', 'bindr', 'RColorBrewer', 'dichromat', 'munsell', 'labeling', 'viridisLite', 'rematch', 'plyr', 'psych', 'reshape2', 'assertthat', 'bindrcpp', 'glue', 'pkgconfig', 'rlang', 'R6', 'Rcpp', 'BH', 'plogr', 'digest', 'gtable', 'scales', 'lazyeval', 'mime', 'curl', 'openssl', 'cellranger', 'stringi', 'selectr', 'tidyselect', 'broom', 'dplyr', 'forcats', 'ggplot2', 'haven', 'httr', 'hms', 'jsonlite', 'lubridate', 'magrittr', 'modelr', 'purrr', 'readr', 'readxl', 'stringr', 'tibble', 'rvest', 'tidyr', 'xml2'

```
# install.packages("tidyverse")
library(tidyverse)
library(magrittr)

# other useful depedencies
library(tableone)
library(psych)
library(fBasics)

# Sometimes function mask each other !
# tib %>% filter(Sex == "Male", Eye=="Brown") # produce error
filter = dplyr::filter
# tib %>% filter(Sex == "Male", Eye=="Brown") # now its ok
# vignette("dplyr")
```

Load some datesets

```
#install.packages("datasets")
# install.packages("mlbench")
# library(help= "mlbench") (Glass, BreastCancer, BostonHousin, Ionosphere, PimaIndiansDiabetes, Sonar)
library(datasets)
library(mlbench)
data("BostonHousing")
head(BostonHousing)
##
        crim zn indus chas
                                               dis rad tax ptratio
                                    rm age
                                                                        b
                             nox
## 1 0.00632 18 2.31
                        0 0.538 6.575 65.2 4.0900
                                                     1 296
                                                              15.3 396.90
## 2 0.02731 0 7.07
                         0 0.469 6.421 78.9 4.9671
                                                     2 242
                                                              17.8 396.90
## 3 0.02729 0 7.07
                         0 0.469 7.185 61.1 4.9671
                                                     2 242
                                                              17.8 392.83
## 4 0.03237 0 2.18
                        0 0.458 6.998 45.8 6.0622
                                                    3 222
                                                              18.7 394.63
## 5 0.06905 0 2.18
                         0 0.458 7.147 54.2 6.0622
                                                     3 222
                                                              18.7 396.90
## 6 0.02985 0 2.18
                         0 0.458 6.430 58.7 6.0622
                                                    3 222
                                                              18.7 394.12
##
    1stat medv
## 1 4.98 24.0
```

```
## 2 9.14 21.6
## 3 4.03 34.7
## 4 2.94 33.4
## 5 5.33 36.2
## 6 5.21 28.7
data("HairEyeColor")
HairEyeColor
## , , Sex = Male
##
##
         Eye
## Hair
          Brown Blue Hazel Green
             32
                       10
##
    Black
                 11
##
    Brown
             53
                  50
                        25
                             15
##
    Red
             10
                  10
                        7
                              7
    Blond
##
              3
                 30
                              8
##
## , , Sex = Female
##
##
         Eye
## Hair
          Brown Blue Hazel Green
##
    Black
             36
                  9
                        5
##
    Brown
             66
                  34
                        29
                             14
    Red
                  7
                        7
                              7
             16
##
    Blond
              4
                  64
                        5
                              8
data("mtcars")
head(mtcars)
##
                    mpg cyl disp hp drat wt qsec vs am gear carb
## Mazda RX4
                    21.0 6 160 110 3.90 2.620 16.46 0 1
## Mazda RX4 Wag
                    21.0 6 160 110 3.90 2.875 17.02 0 1
## Datsun 710
                    22.8 4 108 93 3.85 2.320 18.61 1 1
## Hornet 4 Drive
                    21.4 6 258 110 3.08 3.215 19.44 1 0
## Hornet Sportabout 18.7 8 360 175 3.15 3.440 17.02 0 0
## Valiant
                         6 225 105 2.76 3.460 20.22 1 0
                    18.1
data("swiss")
head(swiss)
##
               Fertility Agriculture Examination Education Catholic
## Courtelary
                    80.2
                               17.0
                                      15 12 9.96
## Delemont
                    83.1
                               45.1
                                                      9
                                                            84.84
                                             6
## Franches-Mnt
                    92.5
                               39.7
                                             5
                                                       5
                                                            93.40
## Moutier
                    85.8
                               36.5
                                             12
                                                      7
                                                            33.77
                                                     15
## Neuveville
                    76.9
                               43.5
                                            17
                                                            5.16
## Porrentruy
                    76.1
                               35.3
                                             9
                                                      7
                                                            90.57
               Infant.Mortality
## Courtelary
                          22.2
## Delemont
                          22.2
## Franches-Mnt
                          20.2
## Moutier
                          20.3
## Neuveville
                          20.6
## Porrentruy
                          26.6
```

```
data("BreastCancer")
head(BreastCancer)
         Id Cl.thickness Cell.size Cell.shape Marg.adhesion Epith.c.size
## 1 1000025
                       5
                                 1
## 2 1002945
                       5
                                                                      7
                                 4
                                                          5
                                                                      2
## 3 1015425
                       3
                                 1
                                            1
                                                         1
## 4 1016277
                       6
                                 8
                                                                      3
                                                         1
                                                                      2
## 5 1017023
                                 1
                                            1
                       8
                                10
## 6 1017122
                                           10
    Bare.nuclei Bl.cromatin Normal.nucleoli Mitoses
                                                       Class
## 1
             1
                     3
                                          1
                                                      benign
## 2
             10
                        3
                                          2
                                                       benign
## 3
             2
                         3
                                          1
                                                  1
                                                       benign
## 4
              4
                          3
                                          7
                                                       benign
## 5
                          3
              1
                                          1
                                                  1
                                                       benign
## 6
             10
                                                  1 malignant
```

Manipulate data with dplyr and the pipe operator

```
Add a pipe operator : CTRL + SHIFT + M
tib = as.tibble(HairEyeColor)
# Filtering and selecting
tib %>% filter(Sex == "Male", Eye=="Brown")
## # A tibble: 4 x 4
     Hair
            Eye
                  Sex
     <chr> <chr> <chr> <chr> <dbl>
## 1 Black Brown Male
## 2 Brown Brown Male
                          53
      Red Brown Male
                          10
## 4 Blond Brown Male
                           3
tib %>% select(contains('S'))
## # A tibble: 32 x 1
##
        Sex
##
      <chr>
##
  1 Male
##
   2 Male
## 3 Male
## 4 Male
## 5 Male
## 6 Male
## 7 Male
## 8 Male
## 9 Male
## 10 Male
## # ... with 22 more rows
tib %>% select(Eye:n)
```

A tibble: 32 x 3

```
##
       Eve Sex
##
      <chr> <chr> <dbl>
## 1 Brown Male
## 2 Brown Male
                     53
   3 Brown Male
                     10
## 4 Brown Male
## 5 Blue Male
                    11
## 6 Blue Male
                     50
## 7 Blue Male
                    10
                    30
## 8 Blue Male
## 9 Hazel Male
                    10
## 10 Hazel Male
                     25
## # ... with 22 more rows
mtcars %>%
  group_by(cyl, am) %>%
  select(mpg, cyl, wt, am) %>%
  summarise(avgmpg = mean(mpg), avgwt = mean(wt)) %>%
  filter(avgmpg > 20)
## # A tibble: 3 x 4
## # Groups: cyl [2]
##
       cyl
             am
                 avgmpg
                            avgwt
##
     <dbl> <dbl>
                   <dbl>
                            <dbl>
              0 22.90000 2.93500
## 1
        4
## 2
              1 28.07500 2.04225
         4
         6
              1 20.56667 2.75500
# cw %>% group_by(sid) %>% summarise(accuracy = mean(correct)) %>% arrange(desc(accuracy))
# select(iris, contains("."))
# Select columns whose name contains a character string.
# select(iris, ends_with("Length"))
# Select columns whose name ends with a character string.
# select(iris, everything()) Select every column.
# select(iris, matches(".t."))
# Select columns whose name matches a regular expression.
# select(iris, num_range("x", 1:5)) Select columns named x1, x2, x3, x4, x5.
# select(iris, one_of(c("Species", "Genus")))
# Select columns whose names are in a group of names.
# select(iris, starts_with("Sepal"))
# Select columns whose name starts with a character string.
# select(iris, Sepal.Length:Petal.Width)
# Select all columns between Sepal.Length and Petal.Width (inclusive).
# select(iris, -Species)
# Select all columns except Species.
```

Assignment

```
df <- mpg %>% filter(cty > 20, cyl == 4) %>% select(1:2)
mpg %>% filter(cty > 20, cyl == 4) %>% select(1:2) -> df
```

```
df %<>% filter(manufacturer != "honda")
df %>% table %>% as.tibble %>% filter(n!=0)
## # A tibble: 12 x 3
##
     manufacturer
                      model
##
           <chr>
                       <chr> <int>
## 1
            audi
                           a4
## 2
          nissan
                      altima
## 3
          toyota
                                 4
                       camry
## 4
           toyota camry solara
                                 4
## 5
                      corolla
                                 5
           toyota
## 6 volkswagen
                          gti
                                 3
## 7
           subaru impreza awd
                                 1
                        jetta
## 8
      volkswagen
                                 4
       chevrolet
## 9
                       malibu
                                 1
## 10
       volkswagen new beetle
                                 3
                                 2
## 11
       volkswagen
                       passat
## 12
          hyundai
                       sonata
                                 2
all_letters <- c(letters, LETTERS) %>%
   sort %>%
   write.csv(file = "all_letters.csv")
```

Using functional sequences

```
library(magrittr) # needed to include the pipe operators
library(lubridate) # in tidyverse
##
## Attaching package: 'lubridate'
## The following object is masked from 'package:base':
##
##
       date
read_year <- . %>% as.character %>% as.Date %>% year
# Creating a dataset
df <- data.frame(now = "2015-11-11", before = "2012-01-01")
#
           now
                   before
# 1 2015-11-11 2012-01-01
# Example 1: applying `read_year` to a single character-vector
df$now %>% read_year
## [1] 2015
# [1] 2015
# Example 2: applying `read_year` to all columns of `df`
df %% lapply(read_year) %% as.data.frame # implicit `lapply(df, read_year)
##
     now before
## 1 2015
           2012
```

```
# now before
# 1 2015  2012

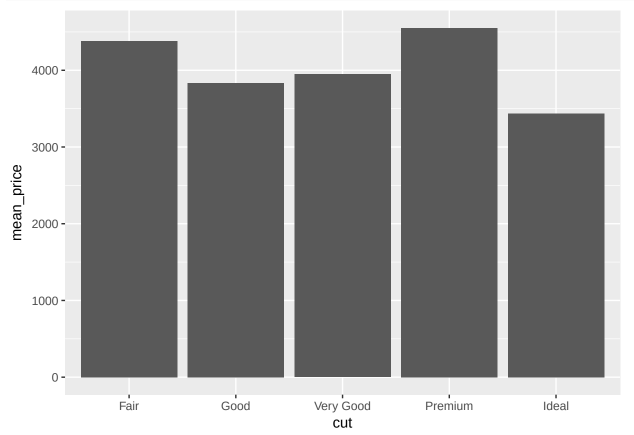
# Example 3: same as above using `mutate_all`
library(dplyr)
df %>% mutate_all(funs(read_year))

## now before
## 1 2015  2012

# if an older version of dplyr use `mutate_each`
# now before
# 1 2015  2012
```

Plots

```
library(ggplot2)
diamonds %>%
    filter(depth > 60) %>%
    group_by(cut) %>%
    summarize(mean_price = mean(price)) %>%
    ggplot(aes(x = cut, y = mean_price)) +
        geom_bar(stat = "identity")
```



Basic renaming

```
require(pander)
## Loading required package: pander
# example of variable renaming
breastcancer = BreastCancer %>% rename_all(tolower)
cat("toLower", pandoc.list(names(breastcancer)))

    id

   • cl.thickness
   • cell.size
   • cell.shape
  • marg.adhesion
   • epith.c.size
   • bare.nuclei
   • bl.cromatin
   • normal.nucleoli
   • mitoses
   • class
toLower
# you could also use base functions
breastcancer %<>% setNames(tolower(gsub("\\.", "_", names(.))))
cat("setNames(tolower(gsub(.)))", pandoc.list(names(breastcancer)))
   • id
  • cl thickness
   • cell size
  • cell_shape

    marg_adhesion

  • epith_c_size
   • bare nuclei
   \bullet bl_cromatin
   • normal nucleoli
   • mitoses
   • class
setNames(tolower(gsub(.)))
```

Reshaping

Base R function for reshaping datasets is reshape Some very usefull parameters

```
x = rnorm(4), y = runif(4))
reshape(df, timevar = "visit", idvar = "id", direction = "wide")
##
        x.Before y.Before
                              x.After
                                         y.After
     1 0.1094677 0.1292292 -1.710644 0.2408263
     2 -0.9349554 0.6174639 -1.466715 0.3039000
## 5 3 0.1094677 0.1292292 -1.710644 0.2408263
## 7 4 -0.9349554 0.6174639 -1.466715 0.3039000
reshape(df, timevar = "visit", idvar = "id", direction = "wide", v.names = "x")
## Warning in reshapeWide(data, idvar = idvar, timevar = timevar, varying =
## varying, : some constant variables (y) are really varying
##
                   x.Before
                               x.After
## 1
     1 0.1292292 0.1094677 -1.710644
     2 0.6174639 -0.9349554 -1.466715
## 5 3 0.1292292 0.1094677 -1.710644
## 7 4 0.6174639 -0.9349554 -1.466715
df3 <- data.frame(id = 1:4,
                 age = c(40,50,60,50),
                 dose1 = c(1,2,1,2),
                 dose2 = c(2,1,2,1),
                 dose4 = c(3,3,3,3))
reshape(df3, direction = "long", varying = 3:5, sep = "")
##
       id age time dose
## 1.1 1
          40
                 1
## 2.1
       2
          50
                 1
## 3.1 3
          60
                      1
                 1
## 4.1
          50
       4
                1
## 1.2
       1
          40
                2
                     2
## 2.2
       2
                2
          50
                     1
## 3.2
       3
                2
                     2
          60
## 4.2 4
          50
                2
                     1
## 1.4
          40
                     3
       1
                 4
       2
                     3
## 2.4
          50
                4
                      3
## 3.4
       3
          60
## 4.4
       4 50
                      3
```

Use TidyR / Reshape2

Best organization for datasets

Rules:

- Each variable has its own column
- Each observation in its own row
- Each value is placed in its own cell

Functions gather, spread, separate, and unite are the building block of the tidyR library Here parameters are very simple so you have to have a good understanding of how you want to reshape your data.

Sometimes what you really want to do is create new variables: for that you need all the smaller functions to make it longer first, unite variables, then spread it out so that each variable has its own column.

```
stocks = data.frame(
  time = as.Date('2009-01-01') + 0:9,
  X = rnorm(10, 0, 1),
 Y = rnorm(10, 0, 2),
  Z = rnorm(10, 0, 4)
head(stocks)
##
          time
## 1 2009-01-01 -0.6619441 -2.835567 3.5165349
## 2 2009-01-02 -2.0694765 -1.776735 -0.2858571
## 3 2009-01-03 -0.9062100 3.194403 1.0223352
## 4 2009-01-04 0.3172697 -2.198807 8.7843088
## 5 2009-01-05 -1.3213492 1.613739 -2.1011186
## 6 2009-01-06 -1.5920188 -3.006474 -1.3597501
# Gather all the selected variables in two columns stock and price
# you can use the minus sign to exclude columns from the selection
stocksm = stocks %>% gather(stock, price, -time)
stocksm %>% head
##
           time stock
                           price
## 1 2009-01-01
                   X -0.6619441
                   X -2.0694765
## 2 2009-01-02
## 3 2009-01-03
                   X -0.9062100
                   X 0.3172697
## 4 2009-01-04
## 5 2009-01-05
                   X -1.3213492
                   X -1.5920188
## 6 2009-01-06
# You can invert what you just did using spread
stocksm %>% spread(stock, price) %>% head
##
                                   Y
                                              Ζ
          time
                         X
## 1 2009-01-01 -0.6619441 -2.835567 3.5165349
## 2 2009-01-02 -2.0694765 -1.776735 -0.2858571
## 3 2009-01-03 -0.9062100 3.194403 1.0223352
## 4 2009-01-04 0.3172697 -2.198807 8.7843088
## 5 2009-01-05 -1.3213492 1.613739 -2.1011186
## 6 2009-01-06 -1.5920188 -3.006474 -1.3597501
# But you can also spread based on another variable
stocksm %>% spread(time, price) %>% head
##
     stock 2009-01-01 2009-01-02 2009-01-03 2009-01-04 2009-01-05 2009-01-06
## 1
        X -0.6619441 -2.0694765 -0.906210 0.3172697
                                                       -1.321349
                                                                   -1.592019
## 2
         Y -2.8355666 -1.7767351
                                   3.194403 -2.1988068
                                                         1.613739
                                                                   -3.006474
## 3
        Z 3.5165349 -0.2858571
                                   1.022335 8.7843088 -2.101119 -1.359750
     2009-01-07 2009-01-08 2009-01-09 2009-01-10
## 1 -1.2628544 -2.304841 0.4334867
                                      -0.173297
## 2 -0.1993526
                -2.125841 -1.0827912
                                       -1.958514
## 3 1.2422260 -2.814044 5.3319974
                                        1.382366
# Use 'convert = TRUE' to produce variables of mixed type
df \leftarrow data.frame(id = rep(c(1, 2), each = 3),
                 var = c("Sepal.Length", "Species_num"),
```

```
value = c(5.1, "setosa", 1, 7.0, "versicolor", 2))
df
##
    id
                var
                         value
## 1 1 Sepal.Length
                           5.1
## 2 1
            Species
                        setosa
## 3 1 Species_num
                             1
## 4 2 Sepal.Length
                             7
## 5 2
            Species versicolor
## 6 2 Species_num
df %>% spread(var, value)
    id Sepal.Length
                       Species Species num
## 1 1
                5.1
                        setosa
## 2 2
                  7 versicolor
df %>% spread(var, value, convert = TRUE)
    id Sepal.Length
                       Species Species_num
## 1 1
                5.1
                        setosa
## 2 2
                7.0 versicolor
# Unite several columns into one
storms %>% unite(date, day:year, sep = "-")
## # A tibble: 10,010 x 11
##
      name
                date hour
                             lat long
                                                    status category wind
##
               <chr> <dbl> <dbl> <dbl>
   * <chr>
                                                     <chr>
                                                              <ord> <int>
                       0 27.5 -79.0 tropical depression
       Amy 27-6-1975
                                                                 -1
                                                                       25
  1
##
       Amy 27-6-1975
                         6 28.5 -79.0 tropical depression
                                                                       25
                                                                 -1
##
  3
       Amy 27-6-1975
                      12 29.5 -79.0 tropical depression
                                                                 -1
                                                                       25
## 4
       Amy 27-6-1975 18 30.5 -79.0 tropical depression
                                                                 -1
                                                                       25
## 5
       Amy 28-6-1975 0 31.5 -78.8 tropical depression
                                                                       25
                                                                -1
                        6 32.4 -78.7 tropical depression
## 6
       Amy 28-6-1975
                                                                 -1
                                                                       25
##
  7
       Amy 28-6-1975 12 33.3 -78.0 tropical depression
                                                                 -1
                                                                       25
##
       Amy 28-6-1975 18 34.0 -77.0 tropical depression
                                                                 -1
                                                                       30
## 9
       Amy 29-6-1975
                        0 34.4 -75.8
                                            tropical storm
                                                                  0
                                                                       35
## 10
       Amy 29-6-1975
                         6 34.0 -74.8
                                            tropical storm
                                                                  0
                                                                       40
## # ... with 10,000 more rows, and 3 more variables: pressure <int>,
## # ts_diameter <dbl>, hu_diameter <dbl>
# Separate values into new column based on pattern
example <- tibble(`N [ears]` = c("173", "60", "54 [96]", "168 [328]", "906 [1685]"),
                  '% Otorrhea' = c("58.61%", "13.30%", "11.11%", "52.38%", "14.79% [10.45%]"))
example %>%
   separate(`N [ears]`, into = c("N_patients", "N_ears"), sep = "\\s\\[", fill = "right") %>%
    separate('% Otorrhea', into = c("pct_patients", "pct_ears"), sep = "\\s\\[", fill = "right") %>%
   mutate_each(funs(parse_number))
## `mutate_each()` is deprecated.
## Use `mutate_all()`, `mutate_at()` or `mutate_if()` instead.
## To map `funs` over all variables, use `mutate_all()`
## # A tibble: 5 x 4
   N_patients N_ears pct_patients pct_ears
```

```
##
         <dbl> <dbl>
                              <dbl>
                                       <dbl>
## 1
            173
                   NΑ
                              58.61
                                          NΑ
## 2
            60
                   NA
                              13.30
                                          NA
## 3
            54
                   96
                              11.11
                                         NA
## 4
            168
                   328
                              52.38
                                          NA
## 5
            906
                              14.79
                                       10.45
                  1685
# Exercice
# We will try to do what the reshape function did for us on a
df \leftarrow data.frame(id = rep(1:4, rep(2,4)),
                 visit = rep(c("Before", "After"), 4),
                 x = rnorm(4), y = runif(4)
reshape(df, timevar = "visit", idvar = "id", direction = "wide")
           x.Before y.Before
                                 x.After
                                            y.After
## 1 1 0.002239927 0.9390014 -0.7418142 0.92362157
## 3 2 1.977745827 0.2196017 -0.2763168 0.03209948
## 5 3 0.002239927 0.9390014 -0.7418142 0.92362157
## 7 4 1.977745827 0.2196017 -0.2763168 0.03209948
# SOLUTION
# first gather variales into two columns called variable and value
df %<>% gather(variable, value, -(c("id", "visit"))) %>% print
##
      id visit variable
                                value
                     x 0.002239927
## 1
       1 Before
## 2
      1 After
                      x -0.741814188
## 3
       2 Before
                      x 1.977745827
## 4
       2 After
                      x -0.276316788
## 5
       3 Before
                      x 0.002239927
## 6
       3 After
                      x -0.741814188
## 7
                      x 1.977745827
       4 Before
## 8
     4 After
                      x -0.276316788
## 9
       1 Before
                      y 0.939001360
                      у 0.923621567
## 10 1 After
## 11 2 Before
                      у 0.219601650
## 12 2 After
                         0.032099484
                      У
## 13 3 Before
                      y 0.939001360
                      у 0.923621567
## 14 3 After
## 15 4 Before
                         0.219601650
                      У
                      y 0.032099484
## 16 4 After
# Then create the new variables X before X after and Y before Y after by uniting the two columns
df %<>% unite(variable, variable, visit, sep = "_") %>% print
##
      id variable
                        value
## 1
       1 x_Before 0.002239927
      1 x_After -0.741814188
## 3
       2 x_Before 1.977745827
      2 x_After -0.276316788
       3 x_Before 0.002239927
## 5
## 6
      3 x_After -0.741814188
## 7
       4 x Before 1.977745827
## 8
       4 x After -0.276316788
      1 y_Before 0.939001360
## 9
```

```
## 10 1 y_After 0.923621567
## 11 2 y_Before 0.219601650
## 12 2 y After 0.032099484
## 13 3 y_Before 0.939001360
## 14 3 y_After 0.923621567
## 15 4 y Before 0.219601650
## 16 4 y_After 0.032099484
# Finally spread into one column per variable
df %<>% spread(variable, value) %>% print
   id
          x_After
                     x_Before
                                y_After y_Before
## 1 1 -0.7418142 0.002239927 0.92362157 0.9390014
## 2 2 -0.2763168 1.977745827 0.03209948 0.2196017
## 3 3 -0.7418142 0.002239927 0.92362157 0.9390014
## 4 4 -0.2763168 1.977745827 0.03209948 0.2196017
melt and cast are the two main functions of reshape2
# convert matrix to data.frame
HairEyeColor %>% reshape2::melt() -> tib
tib = as.tibble(HairEyeColor) # !! Does not transforms character columns as factor
```

Basic stats

```
# Linear regression
# Logistic regression
```

Pandoc

```
require(fBasics)
mpg %>% mutate all(as.numeric) %>% basicStats %>% t %>% pandoc.table
## Warning in evalq(as.numeric(manufacturer), <environment>): NAs introduced
## by coercion
## Warning in evalq(as.numeric(model), <environment>): NAs introduced by
## coercion
## Warning in evalq(as.numeric(trans), <environment>): NAs introduced by
## coercion
## Warning in evalq(as.numeric(drv), <environment>): NAs introduced by
## coercion
## Warning in evalq(as.numeric(fl), <environment>): NAs introduced by coercion
## Warning in evalq(as.numeric(class), <environment>): NAs introduced by
## coercion
## Warning in min(X): no non-missing arguments to min; returning Inf
## Warning in max(X): no non-missing arguments to max; returning -Inf
## Warning in min(X): no non-missing arguments to min; returning Inf
```

```
## Warning in max(X): no non-missing arguments to max; returning -Inf
## Warning in min(X): no non-missing arguments to min; returning Inf
## Warning in max(X): no non-missing arguments to max; returning -Inf
## Warning in min(X): no non-missing arguments to min; returning Inf
## Warning in max(X): no non-missing arguments to max; returning -Inf
## Warning in min(X): no non-missing arguments to min; returning Inf
## Warning in max(X): no non-missing arguments to max; returning -Inf
```

Table 1: Table continues below

	nobs	NAs	Minimum	Maximum	1. Quartile	3. Quartile
manufacturer	234	234	Inf	-Inf	NA	NA
model	234	234	Inf	-Inf	NA	NA
displ	234	0	1.6	7	2.4	4.6
year	234	0	1999	2008	1999	2008
cyl	234	0	4	8	4	8
trans	234	234	Inf	-Inf	NA	NA
${f drv}$	234	131	4	4	4	4
$\operatorname{\mathbf{cty}}$	234	0	9	35	14	19
hwy	234	0	12	44	18	27
fl	234	234	Inf	-Inf	NA	NA
${f class}$	234	234	Inf	-Inf	NA	NA

Table 2: Table continues below

	Mean	Median	Sum	SE Mean	LCL Mean	UCL Mean
manufacturer	NA	NA	0	NA	NA	NA
\mathbf{model}	NA	NA	0	NA	NA	NA
$\operatorname{\mathbf{displ}}$	3.472	3.3	812.4	0.08446	3.305	3.638
year	2004	2004	468819	0.2948	2003	2004
cyl	5.889	6	1378	0.1053	5.681	6.096
trans	NA	NA	0	NA	NA	NA
drv	4	4	412	0	4	4
$\operatorname{\mathbf{cty}}$	16.86	17	3945	0.2782	16.31	17.41
hwy	23.44	24	5485	0.3893	22.67	24.21
fl	NA	NA	0	NA	NA	NA
${f class}$	NA	NA	0	NA	NA	NA

	Variance	Stdev	Skewness	Kurtosis
manufacturer	NA	NA	NA	NA
${f model}$	NA	NA	NA	NA
${f displ}$	1.669	1.292	0.4386	-0.9106
year	20.34	4.51	0	-2.009
\mathbf{cyl}	2.597	1.612	0.1123	-1.464
${f trans}$	NA	NA	NA	NA
\mathbf{drv}	0	0	NA	NA
\mathbf{cty}	18.11	4.256	0.7864	1.431
$\mathbf{h}\mathbf{w}\mathbf{y}$	35.46	5.955	0.3645	0.1369

	Variance	Stdev	Skewness	Kurtosis
fl	NA	NA	NA	NA
class	NA	NA	NA	NA

df2latex

A couple of xx2latex function allows to transform data into a latex format that can help produce clean graphics. To show the graphics or in general to use the output of a code chunk as if you wrote the RMarkdown code directly, use the chunk option results="asis"

```
require(fBasics)
require(psych)
# some utilities
to.latex.table = function (data, ...) {
    data %>%
        df2latex(silent = T, file="tempcorrtest.txt", ...) %>%
        gsub(pattern = "\\begin{tabular}", replacement = "\\resizebox{\\textwidth}{!}{\\begin{tabular}"
        gsub(pattern = "\\end{tabular}", replacement = "\\end{tabular}}", ., fixed = T) %>%
       paste0(collapse = " ")
}
corr.test.2latex = function (data, variables = NULL, method="spearman", ...) {
    if (is.null(variables)) { variables = names(data)}
   rs = corr.test(data %>% select(variables), method=method) #find the probabilities of the correlatio
    cp =corr.p(r=rs$r, n = NROW(data))
    cpp = rs$r
    cpp[cp$p>0.05] = 3
    colnames(cpp) = paste0(1:NROW(cpp), ".")
    rownames(cpp) = paste(colnames(cpp), gsub("_", " ", rownames(cpp), perl=T))
   cpp[upper.tri(cpp)] = NA
    cpp %>%
        df2latex(silent = T, file="tempcorrtest.txt", ...) %>%
        gsub(pattern = "3.00", replacement = ".", ., fixed = T) %>%
       gsub(pattern = "\begin{tabular}", replacement = "\resizebox{\\textwidth}{!}{\\begin{tabular}"
        gsub(pattern = "\\end{tabular}", replacement = "\\end{tabular}}", ., fixed = T) %>%
       paste0(collapse = " ")
}
shapiro.test.p = function (x) { shapiro.test(x)$p.value }
shapiro.test.star = function (x) {if (shapiro.test(x)$p.value<0.05) { stars.pval(shapiro.test(x)$p.value
cat(mpg %>% mutate_all(as.numeric) %>% basicStats %>% t %>% to.latex.table)
cat(breastcancer %>% select(-id) %>% mutate_all(as.numeric) %>% corr.test.2latex)
```

Table 4: df2latex

Variable	nobs	NAs	Minmm	Maxmm	1.Qrt	3.Qrt	Mean	Medin	Sum	SEMen	LCLMn	UCLMn	Varnc	Stdev	Skwns	Krtss
manufacturer	234.00	234.00	Inf	-Inf			NaN		0.00						NaN	NaN
model	234.00	234.00	Inf	-Inf			NaN		0.00						NaN	NaN
displ	234.00	0.00	1.60	7.00	2.40	4.60	3.47	3.30	812.40	0.08	3.31	3.64	1.67	1.29	0.44	-0.91
year	234.00	0.00	1999.00	2008.00	1999.00	2008.00	2003.50	2003.50	468819.00	0.29	2002.92	2004.08	20.34	4.51	0.00	-2.01
cyl	234.00	0.00	4.00	8.00	4.00	8.00	5.89	6.00	1378.00	0.11	5.68	6.10	2.60	1.61	0.11	-1.46
trans	234.00	234.00	Inf	-Inf			NaN		0.00						NaN	NaN
drv	234.00	131.00	4.00	4.00	4.00	4.00	4.00	4.00	412.00	0.00	4.00	4.00	0.00	0.00	NaN	NaN
cty	234.00	0.00	9.00	35.00	14.00	19.00	16.86	17.00	3945.00	0.28	16.31	17.41	18.11	4.26	0.79	1.43
hwy	234.00	0.00	12.00	44.00	18.00	27.00	23.44	24.00	5485.00	0.39	22.67	24.21	35.46	5.95	0.36	0.14
fl	234.00	234.00	Inf	-Inf			NaN		0.00						NaN	NaN
class	234.00	234.00	Inf	-Inf			NaN		0.00						NaN	NaN

Table 5: df2latex

A table from the psych package in R

Variable	1.	2.	3.	4.	5.	6.	7.	8.	9.	10.
1. cl thickness	1.00									
2. cell size	0.67	1.00								
3. cell shape	0.66	0.89	1.00							
4. marg adhesion	0.54	0.74	0.71	1.00						
5. epith c size	0.58	0.79	0.76	0.67	1.00					
6. bare nuclei	0.59	0.77	0.75	0.70	0.69	1.00				
7. bl cromatin	0.54	0.72	0.69	0.62	0.64	0.68	1.00			
8. normal nucleoli	0.57	0.76	0.73	0.63	0.71	0.66	0.66	1.00		
9. mitoses	0.42	0.51	0.47	0.45	0.48	0.47	0.39	0.50	1.00	
10. class	0.68	0.86	0.84	0.73	0.76	0.84	0.74	0.74	0.53	1.00

Tables

A useful package to create tables is tableone. install.packages("tableone")

```
library(tableone)
generate_table = function(data, columns = NULL, strata_g = NULL) {
  if (is.null(columns)) { columns = names(data) }
  columns = unique(columns)
  catVar = NULL
  sapply(columns, function(name) {
    if (class(data[[name]])%in%"factor") {
      catVar <<- c(catVar, name)</pre>
    }
  })
  data_filtered = data %>% select(unique(c(columns,strata_g)))
  if (is.null(strata_g)) {
   table_g <- CreateTableOne(vars = columns, data = data_filtered, factorVars = catVar, includeNA = T)
  } else {
    table_g <- CreateTableOne(columns, data_filtered, catVar, strata = strata_g, includeNA = T)
  }
  mat_g <- print(table_g, exact = "stage", quote = FALSE, noSpaces = TRUE, printToggle = FALSE)</pre>
 return(mat_g)
}
```

```
breastcancer %<>% mutate_each(funs(as.numeric), -id, -class)

## `mutate_each()` is deprecated.

## Use `mutate_all()`, `mutate_at()` or `mutate_if()` instead.

## To map `funs` over a selection of variables, use `mutate_at()`

breastcancer %>% select(-id) %>% generate_table(strata_g = "class")

## Stratified by class

## Benign malignant p test

## n "458" "241" "" ""
```

```
"458"
                                                "241"
                                 "2.96 (1.67)" "7.20 (2.43)" "<0.001" ""
     cl_thickness (mean (sd))
##
                                 "1.33 (0.91)" "6.57 (2.72)" "<0.001" ""
##
     cell size (mean (sd))
##
     cell_shape (mean (sd))
                                 "1.44 (1.00)" "6.56 (2.56)" "<0.001" ""
                                 "1.36 (1.00)" "5.55 (3.21)" "<0.001" ""
##
    marg_adhesion (mean (sd))
                                 "2.12 (0.92)" "5.30 (2.45)" "<0.001" ""
##
     epith_c_size (mean (sd))
    bare_nuclei (mean (sd))
                                 "1.35 (1.18)" "7.63 (3.12)" "<0.001" ""
##
                                 "2.10 (1.08)" "5.98 (2.27)" "<0.001" ""
    bl_cromatin (mean (sd))
##
##
    normal nucleoli (mean (sd)) "1.29 (1.06)" "5.86 (3.35)" "<0.001" ""
                                 "1.06 (0.50)" "2.53 (2.39)" "<0.001" ""
##
     mitoses (mean (sd))
     class = malignant (%)
                                               "241 (100.0)" "<0.001" ""
                                 "0 (0.0)"
```

Recode

http://dplyr.tidyverse.org/reference/recode.html

TidyR

```
# nested dataframes
# data %>% group_by(x) %>% nest()
```

BROOM

http://varianceexplained.org/r/broom-intro/

Survival

https://github.com/pavopax/gists/blob/master/survival-in-R.md

Purrr