Intro to R Statistics

Summary

- ggplot() specifies what data use and what variables will be mapped to where
- inside ggplot(), mapping = aes(x = , y = , color =) specify what variables correspond to what aspects of the plot in general
- · layers of plots can be combined using the + at the **end** of lines
- use geom_line() and geom_point() to add lines and points
- sometimes you need to add a group element to mapping = aes() if your plot looks strange
- make sure you are plotting what you think you are by checking the numbers!
- facet_grid(~ variable) and facet_wrap(~variable) can be helpful to quickly split up your plot

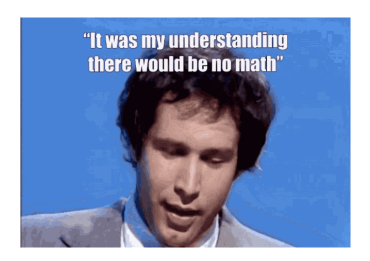
Summary

- the factor class allows us to have a different order from alphanumeric for categorical data
- we can change data to be a factor variable using mutate, the as_factor()
 (of forcats package) or factor() function and specifying the levels with the
 levels argument
- the fct_reorder({variable_to_reorder}, {variable_to_order_by})
 helps us reorder a variable by the values of another variable
- arranging, tabulating, and plotting the data will reflect the new order

Overview

We will cover how to use R to compute some of basic statistics and fit some basic statistical models.

- Correlation
- T-test
- · Linear Regression / Logistic Regression



Overview

We will focus on how to use R software to do these. We will be glossing over the statistical theory and "formulas" for these tests. Moreover, we do not claim the data we use for demonstration meet assumptions of the methods.

There are plenty of resources online for learning more about these methods, as well as dedicated Biostatistics series (at different advancement levels) at the JHU School of Public Health.

Check out <u>www.opencasestudies.org</u> for deep dives on some of the concepts covered here.

Correlation

Correlation

Function cor() computes correlation in R

```
cor(x, y = NULL, use = "everything",
   method = c("pearson", "kendall", "spearman"))
```

- provide two numeric vectors (arguments x, y), or
- provide a data.frame / tibble with numeric columns only
- by default, Pearson correlation coefficient is computed

Correlation

https://jhudatascience.org/intro_to_r/data/Charm_City_Circulator_Ridership.csv

```
library(jhur)
circ <- read circulator()</pre>
head(circ)
# A tibble: 6 \times 15
        date orangeBoardings orangeAlightings orangeAverage purpleBoardir
  day
  <chr> <chr>
                             <db1>
                                               <db1>
                                                             <db1>
                                                                              <dt
1 Monday 01/1...
                               877
                                                1027
                                                              952
2 Tuesday 01/1...
                               777
                                                 815
                                                              796
3 Wednesday 01/1...
                              1203
                                               1220
                                                             1212.
4 Thursday 01/1...
                              1194
                                               1233
                                                             1214.
5 Friday 01/1...
                              1645
                                               1643
                                                             1644
6 Saturday 01/1...
                              1457
                                               1524
                                                             1490.
# ... with 9 more variables: purpleAlightings <dbl>, purpleAverage <dbl>,
    greenBoardings <dbl>, greenAlightings <dbl>, greenAverage <dbl>,
    bannerBoardings <dbl>, bannerAlightings <dbl>, bannerAverage <dbl>,
#
    daily <dbl>
```

Correlation for two vectors

First, we compute correlation by providing two vectors.

Like other functions, if there are NAs, you get NA as the result. But if you specify use only the complete observations, then it will give you correlation using the non-missing data.

```
x <- circ %>% pull(orangeAverage)
y <- circ %>% pull(purpleAverage)

cor(x, y)

[1] NA

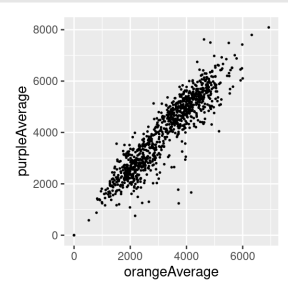
cor(x, y, use = "complete.obs")

[1] 0.9195356
```

Correlation for two vectors with plot

In plot form...

```
circ %>%
  ggplot(aes(x = orangeAverage, y = purpleAverage)) +
  geom_point(size = 0.3)
```



Correlation for data frame columns

0.8395806

0.5447031

We can compute correlation for all pairs of columns of a data frame / matrix. We typically just say, "compute correlation matrix".

Columns must be all numeric!

greenAverage

bannerAverage

```
circ subset Average <- circ %>% select(ends with("Average"))
dim(circ_subset_Average)
[1] 1146
            4
cor_mat <- cor(circ_subset_Average, use = "complete.obs")</pre>
cor mat
              orangeAverage purpleAverage greenAverage bannerAverage
                                0.9078826
                                                           0.5447031
orangeAverage
                  1.0000000
                                             0.8395806
purpleAverage
                  0.9078826
                                1.0000000
                                             0.8665630
                                                           0.5213462
```

0.8665630

0.5213462

1.0000000

0.4533421

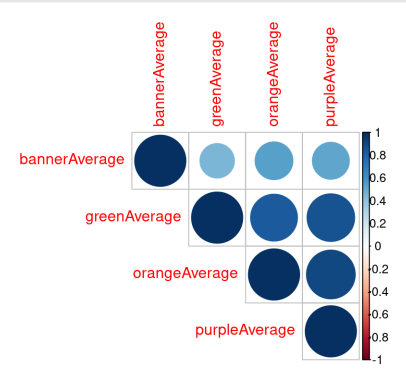
0.4533421

1.0000000

Correlation for data frame columns with plot

Google, "r correlation matrix plot"

```
library(corrplot)
corrplot(cor_mat, type = "upper", order = "hclust")
```



T-test

T-test

The commonly used are:

- one-sample t-test used to test mean of a variable in one group
- two-sample t-test used to test difference in means of a variable between two groups (if the "two groups" are data of the same individuals collected at 2 time points, we say it is two-sample paired t-test)

The t.test() function in R is one to address the above.

Running one-sample t-test

x <- circ %>% pull(orangeAverage)

It tests mean of a variable in one group. By default (i.e., without us explicitly specifying values of other arguments):

- tests whether a mean of a variable is equal to 0 (mu=0)
- uses "two sided" alternative (alternative = "two.sided")
- returns result assuming confidence level 0.95 (conf.level = 0.95)

Running two-sample t-test

It tests test difference in means of a variable between two groups. By default:

- tests whether difference in means of a variable is equal to 0 (mu=0)
- uses "two sided" alternative (alternative = "two.sided")
- returns result assuming confidence level 0.95 (conf.level = 0.95)
- assumes data are not paired (paired = FALSE)
- assumes true variance in the two groups is not equal (var.equal = FALSE)

```
x <- circ %>% pull(orangeAverage)
y <- circ %>% pull(purpleAverage)
t.test(x, y)
```

Welch Two Sample t-test

```
data: x and y
t = -17.076, df = 1984, p-value < 0.000000000000000022
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
   -1096.7602  -870.7867
sample estimates:
mean of x mean of y
3033.161  4016.935</pre>
```

T-test: retrieving information from the result with **broom** package

The broom package has a tidy() function that can organize results into a data frame so that they are easily manipulated (or nicely printed)

```
library(broom)
result <- t.test(x, y)
result tidy <- tidy(result)
result_tidy
# A tibble: 1 \times 10
 estimate estimate1 estimate2 statistic p.value parameter conf.low conf.high
                                <dbl>
                       <dbl>
                                         <dbl> <dbl> <dbl>
                                                                    <dbl>
    <dbl>
              <dbl>
    -984. 3033. 4017. -17.1 4.20e-61
                                                  1984. -1097.
                                                                    -871.
# ... with 2 more variables: method <chr>, alternative <chr>
```

P-value adjustment

You run an increased risk of Type I errors (a "false positive") when multiple hypotheses are tested simultaneously.

Use the p.adjust() function on a vector of p values. Use method = to specify the adjustment method:

```
my_pvalues <- c(0.049, 0.001, 0.31, 0.00001)
p.adjust(my_pvalues, method = "BH") # Benjamini Hochberg

[1] 0.06533333 0.00200000 0.31000000 0.00004000
p.adjust(my_pvalues, method = "bonferroni")

[1] 0.19600 0.00400 1.00000 0.00004</pre>
```

Some other statistical tests

- wilcox.test() Wilcoxon signed rank test, Wilcoxon rank sum test
- shapiro.test() Shapiro test
- · ks.test() Kolmogorov-Smirnov test
- var.test() Fisher's F-Test
- chisq.test() Chi-squared test
- aov() Analysis of Variance (ANOVA)

Lab Part 1

Class Website

Lab

Regression

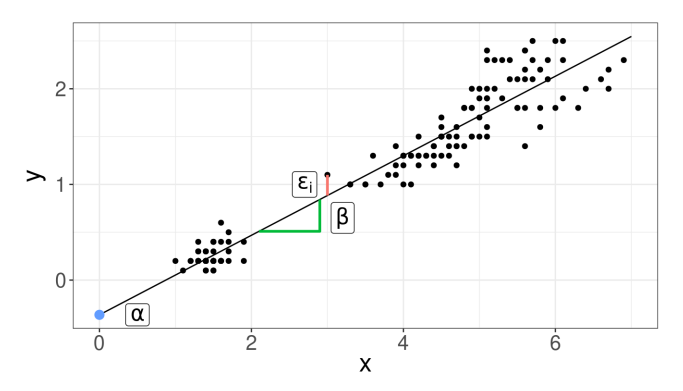
Linear regression is a method to model the relationship between a response and one or more explanatory variables.

We provide a little notation here so some of the commands are easier to put in the proper context.

$$y_i = \alpha + \beta x_i + \varepsilon_i$$

where:

- · y_i is the outcome for person i
- α is the intercept
- β is the slope
- · x_i is the predictor for person i
- · $arepsilon_i$ is the residual variation for person i



Linear regression is a method to model the relationship between a response and one or more explanatory variables.

We provide a little notation here so some of the commands are easier to put in the proper context.

$$y_i = \alpha + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$

where:

- · y_i is the outcome for person i
- α is the intercept
- \cdot β_1 , β_2 , β_2 are the slopes for variables x_{i1} , x_{i2} , x_{i3}
- $\cdot \;\; x_{i1}$, x_{i2} , x_{i3} are the predictors for person i
- · $arepsilon_i$ is the residual variation for person i

Linear regression fit in R

To fit regression models in R, we use the function glm() (Generalized Linear Model).

We typically provide two arguments:

- formula model formula written using names of columns in our data
- · data our data frame

Linear regression fit in R: model formula

Model formula

$$y_i = \alpha + \beta x_i + \varepsilon_i$$

In R translates to

Linear regression fit in R: model formula

Model formula

$$y_i = \alpha + \beta x_i + \varepsilon_i$$

In R translates to

In practice, y and x are replaced with the names of columns from our data set.

For example, if we want to fit a regression model where outcome is income and predictor is years_of_education, our formula would be:

Linear regression fit in R: model formula

Model formula

$$y_i = \alpha + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$

In R translates to

$$y \sim x1 + x2 + x3$$

In practice, y and x1, x2, x3 are replaced with the names of columns from our data set.

For example, if we want to fit a regression model where outcome is income and predictors are years_of_education, age, location then our formula would be:

We will use data about emergency room doctor complaints.

"Data was recorded on 44 doctors working in an emergency service at a hospital to study the factors affecting the number of complaints received."

```
# install.packages("faraway")
library(faraway)

data(esdcomp)
esdcomp
```

	visits	complaints	residency	gender	revenue	hours	
1	2014	2	Y	F	263.03	1287.25	
2	3091	3	N	M	334.94	1588.00	
3	879	1	Υ	M	206.42	705.25	
4	1780	1	N	M	226.32	1005.50	
5	3646	11	N	M	288.91	1667.25	
6	2690	1	N	M	275.94	1517.75	
7	1864	2	Υ	M	295.71	967.00	
8	2782	6	N	M	224.91	1609.25	
9	3071	9	N	F	249.32	1747.75	
10	1502	3	Υ	M	269.00	906.25	
11	2438	2	N	F	225.61	1787.75	
12	2278	2	N	M	212.43	1480.50	
13	2458	5	N	M	211.05	1733.50	
14	2269	2	N	F	213.23	1847.25	
15	2431	7	N	M	257.30	1433.00	
16	3010	2	Υ	M	326.49	1520.00	
17	2234	5	Υ	М	290.53	1404.75	

Linear regression: model fitting

We fit linear regression model with the number of patient visits (visits) as an outcome and total number of hours worked (hours) as a predictor.

Linear regression: model summary

The summary() function returns a list that shows us some more detail

```
summary(fit)
Call:
glm(formula = visits ~ hours, data = esdcomp)
Deviance Residuals:
            10 Median 30
   Min
                                    Max
-797.48 -196.35 -43.15 169.58 864.65
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 140.288 242.723 0.578
                                               0.566
             1.584 0.167 9.488 0.00000000000526 ***
hours
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 128155.3)
   Null deviance: 16919101 on 43 degrees of freedom
Residual deviance: 5382524 on 42 degrees of freedom
AIC: 646.3
Number of Fisher Scoring iterations: 2
```

Linear regression: multiple predictors

Let's try adding another explanatory variable to our model, dollars per hour earned by the doctor (`revenue``).

```
fit2 <- glm(visits ~ hours + revenue, data = esdcomp)</pre>
summary(fit2)
Call:
glm(formula = visits ~ hours + revenue, data = esdcomp)
Deviance Residuals:
   Min
             10 Median
                               30
                                      Max
-420.57 -166.87 -17.45 140.79 616.20
Coefficients:
             Estimate Std. Error t value
                                                    Pr(>|t|)
(Intercept) -2078.1369 327.9157 -6.337
                                               0.00000014326 ***
hours
               1.6179
                      0.1081 14.968 < 0.0000000000000000 ***
                      1.0828 7.706
              8.3437
                                               0.00000000169 ***
revenue
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 53620.97)
   Null deviance: 16919101 on 43 degrees of freedom
Residual deviance: 2198460 on 41 degrees of freedom
AIC: 608.91
Number of Fisher Scoring iterations: 2
```

Linear regression: factors

Factors get special treatment in regression models - lowest level of the factor is the comparison group, and all other factors are relative to its values.

residency takes values Y or N to indicate whether the doctor is a resident.

esdcomp %>% count(residency)

```
residency n
1 N 24
2 Y 20
```

Linear regression: factors

```
fit_3 <- glm(visits ~ residency, data = esdcomp)</pre>
summary(fit_3)
Call:
glm(formula = visits ~ residency, data = esdcomp)
Deviance Residuals:
    Min
              10
                  Median
                                         Max
                                3Q
-1356.30
        -371.55 -62.79 400.46 1527.70
Coefficients:
          Estimate Std. Error t value
                                              Pr(>|t|)
(Intercept) 2510.8
                       residencyY -275.5
                       187.4 -1.47
                                                 0.149
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 383122.6)
   Null deviance: 16919101 on 43 degrees of freedom
Residual deviance: 16091148 on 42 degrees of freedom
AIC: 694.49
Number of Fisher Scoring iterations: 2
```

Generalized Linear Models (GLMs)

Generalized Linear Models (GLMs) allow for fitting regressions for non-continuous/normal outcomes. Examples include: logistic regression, Poisson regression.

Add the **family** argument – a description of the error distribution and link function to be used in the model. These include:

- binomial(link = "logit")
- poisson(link = "log"), and other.

See ?family documentation for details of family functions.

Logistic regression

We will use data about breast cancer tumors.

"The purpose of this study was to determine whether a new procedure called fine needle aspiration which draws only a small sample of tissue could be effective in determining tumor status."

data(wbca)
wbca

	Class	Adhes	BNucl	Chrom	Epith	Mitos	NNucl	Thick	UShap	USize
1	1	1	1	3	2	1	1	5	1	1
2	1	5	10	3	7	1	2	5	4	4
3	1	1	2	3	2	1	1	3	1	1
4	1	1	4	3	3	1	7	6	8	8
5	1	3	1	3	2	1	1	4	1	1
6	0	8	10	9	7	1	7	8	10	10
7	1	1	10	3	2	1	1	1	1	1
8	1	1	1	3	2	1	1	2	2	1
9	1	1	1	1	2	5	1	2	1	1
10	1	1	1	2	2	1	1	4	1	2
11	1	1	1	3	1	1	1	1	1	1
12	1	1	1	2	2	1	1	2	1	1
13	0	3	3	4	2	1	4	5	3	3
14	1	1	3	3	2	1	1	1	1	1
15	0	10	9	5	7	4	5	8	5	7
16	0	4	1	4	6	1	3	7	6	4
17	1	1	1	2	2	1	1	4	1	1
18	1	1	1	3	2	1	1	4	1	1
19	0	6	10	4	4	2	1	10	7	7
20	1	1	1	3	2	1	1	6	1	1
21	0	10	10	5	5	4	4	7	2	3
22	0	3	7	7	6	1	10	10	5	5
23	1	1	1	2	2	1	1	3	1	1
24	1	1	1	3	2	1	1	1	1	1
25	0	4	7	3	2	1	6	5	3	2
26	1	1	1	2	1	1	1	3	1	2
27	1	1	1	2	2	1	1	5	1	1

Logistic regression

Class is a 0/1-valued variable indicating if the tumor was malignant (0 if malignant, 1 if benign).

```
binom_fit <- glm(Class ~ UShap + USize, data = wbca, family = binomial())</pre>
summary(binom_fit)
Call:
glm(formula = Class ~ UShap + USize, family = binomial(), data = wbca)
Deviance Residuals:
   Min
             10 Median
                              3Q
                                      Max
-2.8262 -0.0171 0.1929 0.1929 4.0082
Coefficients:
           Estimate Std. Error z value
                                                  Pr(>|z|)
(Intercept) 5.6868
                     0.4359 13.047 < 0.00000000000000000 ***
            -0.8431 0.1593 -5.292
UShap
                                               0.000000121 ***
            -0.8686 0.1690 -5.139
                                               0.000000277 ***
USize
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 881.39 on 680 degrees of freedom
Residual deviance: 218.28 on 678 degrees of freedom
AIC: 224.28
Number of Fisher Scoring iterations: 7
```

Odds ratios

This data shows whether people became ill after eating ice cream in the 1940s.

head(ice_cream) ill vanilla.ice.cream 1 1 1 ice_cream %>% count(ill, vanilla.ice.cream) ill vanilla.ice.cream 0 0 18 0 1 1 1 11 1 43

Odds ratios

attr(, "method")

Use oddsratio(x, y) from the epitools() package to calculate

```
library(epitools)
response <- ice_cream %>% pull(ill)
predictor <- ice_cream %>% pull(vanilla.ice.cream)
oddsratio(predictor, response)
$data
        Outcome
Predictor 0 1 Total
         18 3
                  21
         11 43
                  54
   Total 29 46 75
$measure
        odds ratio with 95% C.I.
Predictor estimate
                     lower
                              upper
       0 1.00000
                        NA
                                 NA
       1 21.40719 5.927963 109.4384
$p.value
        two-sided
                            fisher.exact
Predictor
              midp.exact
                                             chi.square
                      NA
                                                     NA
       1 0.0000002698215 0.00000002597451 0.0000001813314
$correction
[1] FALSE
```

Final note

Some final notes:

- Researcher's responsibility to understand the statistical method they use underlying assumptions, correct interpretation of method results
- Researcher's responsibility to understand the R software they use meaning of function's arguments and meaning of function's output elements

Summary

- Use cor() to calculate correlation between two vectors. corrplot() is nice for a quick visualization!
- t.test() tests the difference in means between two vectors
- glm() fits regression models:
 - Use the formula = argument to specify the model (e.g., y ~ x or y ~ x1
 + x2 using column names)
 - Use data = to indicate the dataset
 - Use family = binomial() to do a logistic regression
 - summary() gives useful statistics
- oddsratio() from the epitools package can calculate odds ratios
- this is just the tip of the iceberg!

Lab Part 2

Class Website

Lab



Image by Gerd Altmann from Pixabay

Extra Slides

Adding correlation value to a plot

Note that you can add the correlation value to a plot, via the annotate().

```
cor_val <- cor(x, y, use = "complete.obs")
cor_val_label <- paste0("r = ", round(cor_val, 3))

circ %>%
    ggplot(aes(x = orangeAverage, y = purpleAverage)) +
    geom_point(size = 0.3) +
    annotate("text", x = 2000, y = 7500, label = cor_val_label, size = 5)
```

