Statistical inference and linear models

November 20, 2018

If you get bored

- Go to the last slide for bonus exercises
- Work on code for your research and ask question during exercise time
- But try and keep an eye out, just in case



Disclaimer

- Assume you got lectures about statistics and
 - know why we need statistics
 - have heard of the general philosophy
- We may simplify to focus on practical aspects
- Correct us if we say something completely awful

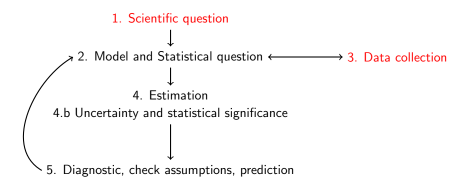
- Statistical inference
- 2 t-test, ANOVA, regression: all is one, one is all
- 3 Linear models in details
- Bonus fun

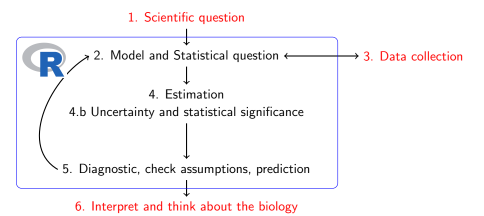
1. Scientific question

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 - \downarrow
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 - 4. Estimation
- 4.b Uncertainty and statistical significance





```
data("iris")
```

```
str(iris)
plot(iris)
```

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- Model and stat question:
 - ► Model:
 - * There is an intrinsic/expected sepal length value for a species; an individual value is the sum of this expectation and a random Gaussian deviation.
 - * $y_i = \mu_{species_i} + \epsilon_i$ with $\epsilon \sim N(0, \sigma^2)$
 - * t-test

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 - ★ t-test
 - Statistical question:
 - ★ Does sepal length differ significantly between the two taxa in our sample?
 - Is the observed difference between taxa likely if both taxa have the same intrinsic/expected value?
- Oata collection



One t-test for sepal length between setosa and versicolor:

```
t.test(x = iris$Sepal.Length[iris$Species == "setosa"],
        y = iris$Sepal.Length[iris$Species == "versicolor"])
Welch Two Sample t-test
data: iris$Sepal.Length[iris$Species == "setosa"] and iris$Sepal.Le
t = -10.521, df = 86.538, p-value < 2.2e-16
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-1.1057074 - 0.7542926
sample estimates:
mean of x mean of y
    5.006 5.936
```

When do we know it is different?

- Statistical estimation
 - a Estimation
 - \star Cannot know true difference $\mu_{\mathit{species}_1} \mu_{\mathit{species}_2}$
 - ★ Estimated difference = Mean₁ Mean₂
 - ★ Difference contains random variation
 - b Quantify uncertainty / Statistical significance

```
* t = \frac{\text{Mean}_1 - \text{Mean}_2}{\text{Variation}} \frac{\sqrt{\text{Sample Size}}}{\sqrt{2}}
```

- * We know exactly how t is distributed when $\mu_{species_1} \mu_{species_2} = 0$
- * Hence we know probability of $\geq t$ if $\mu_{species_1} \mu_{species_2} = 0$ (p-value)
- * Can derive confidence interval and standard error

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Less uncertainty with

- Larger absolute difference
- Smaller variability
- Larger sample size



When do we know it is different? Simulations

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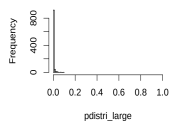
1. Larger absolute difference

```
nbsim <- 1000
pdistri_large <- vector(length = nbsim)</pre>
pdistri_small <- vector(length = nbsim)</pre>
for (i in 1:nbsim)
  x1 \leftarrow rnorm(n = 10, mean = 2, sd = 1)
  x2 \leftarrow rnorm(n = 10, mean = 4, sd = 1) \#large diff
  x3 \leftarrow rnorm(n = 10, mean = 2.5, sd = 1) #small diff
  out_large <- t.test(x1, x2)</pre>
  out_small <- t.test(x1, x3)</pre>
  pdistri_large[i] <-out_large$p.value
  pdistri_small[i] <-out_small$p.value
```

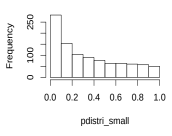
When do we know it is different?

```
par(mfrow=c(1,2), cex=2)
hist(pdistri_large, xlim=c(0,1),
     main=paste("Prop signif=",mean(pdistri_large<0.05)))</pre>
hist(pdistri_small, xlim=c(0,1),
     main=paste("Prop signif=",mean(pdistri_small<0.05)))</pre>
```

Prop signif= 0.992



Prop signif= 0.182



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When do we know it is different? Try it!

Exercise

Check the effect of smaller variability and/or larger sample size.

By the way, what are these p-values?

Probability for a summary statistic to be greater or equal to the observed summary statistic, when the null-hypothesis of a given statistical model is true.

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- \bullet Depends on the null-hypothesis (H_0) of a given model with assumptions
- Uniform distribution under H_0 ...
- ... hence proportion(significance under H_0) = significance threshold

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NB: Focus on p-value criticized, but common and they are no more evil than other misused statistics!

T-test exercise: p-values and simulations

```
t.test(x = ..., y=..., var.equal = TRUE)
t.test(x = ..., y=..., var.equal = FALSE)
```

What if variance are different by chance only?

```
set.seed(1234)
var(rnorm(20, mean = 0, sd = 1))
[1] 1.027806
var(rnorm(20, mean = 0, sd = 1))
[1] 0.6265501
```

Exercise

What option is more correct for var.equal?

- Statistical inference
- 2 t-test, ANOVA, regression: all is one, one is all
- Linear models in details
- Bonus fun

A small example

Animal behavior in response to weather Load data:

```
getwd()
setwd()
```

```
dat.behav <- read.csv(file = "datbehav.csv") # path to file</pre>
```

A small example

Animal behavior in response to weather Load data:

```
getwd()
setwd()
```

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dat.behav <- read.csv(file = "datbehav.csv") # path to file</pre>
```

STEP 1: have a look at your data

```
str(dat.behav)
summary(dat.behav)
plot(dat.behav)
```

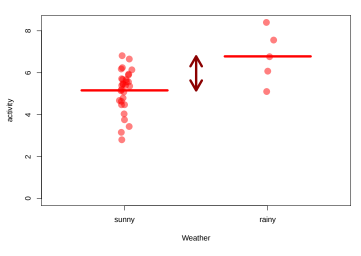
t-test

```
fitstudent <- t.test(x = dat.behav$activity[dat.behav$weather==
                                               "rainy"],
                     y = dat.behav$activity[dat.behav$weather==
                                               "sunny"],
                     var.equal = TRUE)
print(fitstudent)
Two Sample t-test
data: dat.behav$activity[dat.behav$weather == "rainy"] and dat.behav
t = 3.2752, df = 33, p-value = 0.002485
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 0.6138373 2.6270325
sample estimates:
mean of x mean of y
 6.781476 5.161041
```

Linear models

t-test, graphically

Difference between means



ANOVA

- What is the fastest way to get row averages in a data-frame?
- Create a function called colVars, like colMeans but for variance
- Oreate nice plots to visualize iris data (ideally journal-quality)

```
fitanova <- aov(data = dat.behav, formula = activity ~ weather)

summary(fitanova)

Df Sum Sq Mean Sq F value Pr(>F)

weather 1 11.25 11.253 10.73 0.00248 **

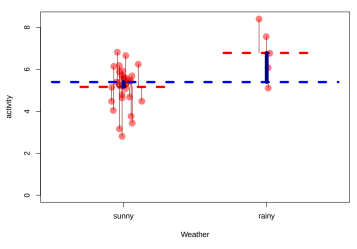
Residuals 33 34.62 1.049

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

ANOVA, graphically

Variance decomposition



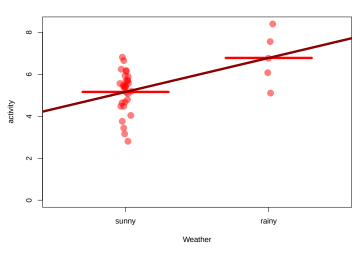
Linear regression

```
fitlm <- lm(data = dat.behav, formula = activity ~ weather)
summary(fitlm)
Call:
lm(formula = activity ~ weather, data = dat.behav)
Residuals:
   Min 1Q Median 3Q Max
-2.3547 -0.6028 0.2346 0.6419 1.6534
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 6.7815 0.4581 14.805 3.94e-16 ***
weathersunny -1.6204 0.4948 -3.275 0.00248 **
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Residual standard error: 1.024 on 33 degrees of freedom
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Regression, graphically

Rate of change



NB: aov() vs. anova()

```
aov(data = dat.behav, formula = activity ~ weather)
anova(fitlm)
```

All is one...

All is one...

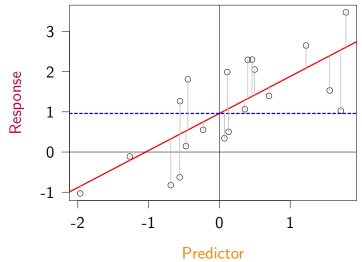
...but lm() rules!

- t-test, ANOVA, regression and others can be mathematically equivalent
- In R, lm() and related functions can do them all...
- ...and much more!

- Statistical inference
- 2 t-test, ANOVA, regression: all is one, one is all
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A simple linear model

$Response = Intercept + Slope \times Predictor + Error$



A simple linear model

```
Response = Intercept + Slope \times Predictor + Error
```

In R:

```
lm(response ~ 1 + predictor1 + predictor2, data=data)
# equivalent to
lm(response ~ predictor1 + predictor2, data=data)
```

- Intercept can be explicit or implicit
- Can remove intercept with $\ldots \sim 0 + \ldots$
- Error is implicit
- Feed the option data= to keep code short, reliable and flexible
- Order of predictors do not matter



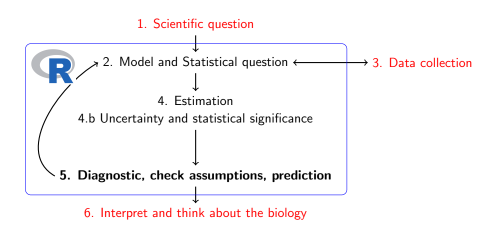
Interpretation

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Im vs. plot

- Fit a linear model $y \sim x$ for each of the four "distri"
- Plot the relationship $y \sim x$ for each of the four "distri"
- Can we trust these models? For what? I expect more than "it's all bullshit"

General approach



Not necessarily wrong, but typical interpretation assumes:

Linear combination of parameters (including transformation, polynoms, interactions...)
 Risk: biologically meaningless

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- Homoscedasticity (constant error variance)
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 Risk: Over-optimistic uncertainty, unreliable predictions
- Independence of error
 Risk: Bias and over-optimistic uncertainty

Diagnostic: summary and plot

```
lm1 <- lm(y ~ x , data=Ans[Ans$distri==1,])
lm2 <- lm(y ~ x , data=Ans[Ans$distri==2,])</pre>
```

```
summary(lm1)
par(mfrow=c(2,2))
plot(lm1)
```

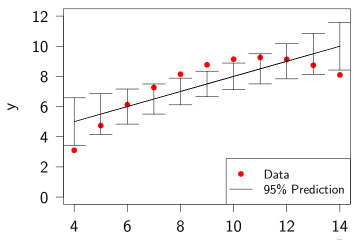
```
summary(lm2)
plot(lm2)
par(mfrow=c(1,1))
```

Diagnostic: prediction

```
pred2 <- predict(lm2, se.fit = TRUE, interval = "confidence")
pred2 <- cbind(Ans[Ans$distri==2,], pred2)</pre>
```

Diagnostic: prediction

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Linear models

Practice Im() with parasites

What explains variation in parasitic load?

You collected ecto-parasites on some furry large mammals at three locations. Parasites break easily when we collect them and are impossible to count, so we decide to measure parasitic load as their mass. Why do some mammals have larger parasitic load?

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- Load the Para.csv data (don't forget: str(), summary(), plot()...)
- Model Parasite_Mass using lm()
- Find what variables predict Parasite_Mass
- How good are your models? Assumptions? Prediction?
- What biological interpretation can you imagine?

- Statistical inference
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Extra exercises

General R coding

- What is the fastest way to get row averages in a data-frame?
- Create a function called colVars, like colMeans but for variance
- Create nice plots to visualize iris data (ideally journal-quality)

Linear models

- Load Cdata.csv, fit models of y predited by x1 and x2, or x2 and x3. Something is weird, what is going on? What to do?
- For model that can be fitted with t.test, aov, and Im, is one of the function faster?
- Write your own code to obtain a prediction from a lm (that is, a simpler version of the predict function), with confidence interval. (extra toughness: do it using the matrix formulation of the analytical solution to a linear model)