#### Introduction to statistics: Linear models

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# Summary

- 1. We learnt about the single sample, two sample, and paired t-tests.
- 2. We learnt about Type I, II error (and power).
- 3. We learnt about Type M and Type S errors.

Now we are ready to look at linear modeling.

## Load Grodner and Gibson dataset

```
gge1crit<-read.table("data/grodnergibson05data.txt", header
head(gge1crit)
     subject item condition rawRT
##
## 6
                   objgap
                           320
          1 2 subjgap 424
## 19
          1 3
## 34
                  objgap 309
         1 4 subjgap 274
## 49
        1 5 objgap 333
## 68
               6
## 80
                   subjgap
                           266
```

# Compute means by factor level

Let's compute the means by factor levels:

The object relative mean is higher than the subject relative mean.

#### Paired t-test on the data

Correct t-test by subject and by items

This is how one would do a t-test CORRECTLY with such data, to compare means across conditions:

```
bysubj <- aggregate (rawRT subject + condition,
                   mean, data=gge1crit)
byitem <- aggregate (rawRT~item+condition, mean, data=gge1crit)
t.test(rawRT~condition,paired=TRUE,bysubj)$statistic
##
## 3.1093
t.test(rawRT~condition, paired=TRUE, byitem) $statistic
##
## 3.7542
```

#### Paired t-test on the data

Consider only by-subject analyses for now.

These are the means we are comparing by subject:

```
round(with(bysubj,
tapply(rawRT,condition,mean)))
## objgap subjgap
## 471 369
```

We can rewrite our best guess about how the object and subject relative clause reading time distributions like this:

Object relative:  $Normal(471, \hat{\sigma})$ 

Subject relative:  $Normal(471 - 102, \hat{\sigma})$ 

Note that the two distributions for object and subject relative are assumed to be independent. This is not true in our data as we get a data point each for each RC type from the same subject!

► The object relative's distribution can be written as a sum of two terms:

$$y = 471 + \epsilon \text{ where } \epsilon \sim Normal(0, \hat{\sigma})$$

► The subject relative's distribution can be written:

$$y = 471 - 102 + \epsilon \text{ where } \epsilon \sim Normal(0, \hat{\sigma})$$

Note that  $\hat{\sigma}=213$  because  $obs.t=\frac{\bar{x}}{s/\sqrt{n}}\Rightarrow s=\bar{x}\times\sqrt{n}/obs.t=-103\times\sqrt{42}/-3.109=213.$ 

The above statements describe a generative process for the data.

Now consider this **linear model**, which describes the rt in each row of the data frame as a function of condition.  $\epsilon$  is a random variable  $\epsilon \sim Normal(0,213)$ .

Object relative reading times:

$$rt = 471 + \epsilon \tag{1}$$

Subject relative reading times:

$$rt = 471 - 102 + \epsilon \tag{2}$$

When describing mean reading times, I can drop the  $\epsilon$ : Object relative reading times:

$$rt = 471 \tag{3}$$

Subject relative reading times:

$$rt = 471 - 102 (4)$$

The lm() function gives us these mean estimates from the data.

Object relative reading times:

$$rt = 471 \times \mathbf{1} - 102 \times \mathbf{0} + \epsilon \tag{5}$$

Subject relative reading times:

$$rt = 471 \times \mathbf{1} - 102 \times \mathbf{1} + \epsilon \tag{6}$$

So, object relatives are coded as 0, and subject relatives are coded as 1.

The lm() function sets up such a model.

With real data from the relative clause study:

```
contrasts(bysubj$condition)
           subjgap
##
## objgap
## subjgap
m0<-lm(rawRT~condition,bysubj)
round(summary(m0)$coefficients)[,1]
##
        (Intercept) conditionsubjgap
                471
                                 -102
##
```

The linear model gives us two numbers: object relative reading time (471), and the difference between object and subject relative (-102):

```
round(coef(m0))
## (Intercept) conditionsubjgap
## 471 -102
```

- 1. The **intercept** is giving us the mean of the objgap condition.
- 2. The **slope** is giving us the amount by which the subject relative is faster.

Note that the meaning of the intercept and slope depends on the ordering of the factor levels. We can make subject relative means be the intercept:

```
m1a<-lm(rawRT~condition,bysubj)
round(coef(m1a))
## (Intercept) conditionobjgap
## 369 102</pre>
```

Now the intercept is the subject relative clause mean. The slope is the increase in reading time for the object relative

condition.

In mathematical form, the model is:

$$rt = \beta_0 + \beta_1 condition + \epsilon \tag{7}$$

where

- $ightharpoonup eta_0$  is the mean for the object relative
- $\triangleright$   $\beta_1$  is the amount by which the object relative mean must be changed to obtain the mean for the subject relative.

The null hypothesis is that the difference in means between the two relative clause types  $\beta_1$  is:

$$H_0: \beta_1 = 0$$

The **contrast coding** determines the meaning of the  $\beta$  parameters:

We will make a distinction between the unknown true mean  $\beta_0, \beta_1$  and the estimated mean from the data  $\hat{\beta}_0, \hat{\beta}_1$ .

- lacksquare Estimated mean object relative processing time:  $\hat{eta}_0=471$  .
- Estimated mean subject relative processing time:

$$\hat{\beta}_0 + \hat{\beta}_1 = 471 + -102 = 369.$$

Reparameterizing the linear model with sum contrast coding

In mathematical form, the model is:

$$rt = \beta_0 + \beta_1 condition + \epsilon$$
 (8)

We can change the **contrast coding** to change the meaning of the  $\beta$  parameters:

```
## new contrast coding:
bysubj$cond<-ifelse(bysubj$condition=="objgap",1,-1)</pre>
```

Reparameterizing the linear model with sum contrast coding

```
xtabs(~cond+condition,bysubj)

## condition

## cond objgap subjgap

## -1 0 42

## 1 42 0
```

Reparameterizing the linear model with sum contrast coding

Now the model parameters have a different meaning:

```
m1<-lm(rawRT~cond,bysubj)
round(coef(m1))
## (Intercept) cond
## 420 51</pre>
```

Reparameterizing the linear model with sum contrast coding

- **E**stimated **grand mean** processing time:  $\hat{\beta}_0 = 420$ .
- Estimated mean object relative processing time:  $\hat{\beta}_0 + \hat{\beta}_1 = 420 + 51 = 471$ .
- Estimated mean subject relative processing time:  $\hat{\beta}_0 \hat{\beta}_1 = 420 51 = 369$ .

This kind of parameterization is called **sum-to-zero contrast** or more simply **sum contrast** coding. This is the coding we will use.

#### The null hypothesis for the slope

The null hypothesis for the slope is

$$H_0: \mathbf{1} \times \mu_{obj} + (-\mathbf{1} \times) \mu_{subj} = 0$$
 (9)

The sum contrasts are referring to the  $\pm 1$  terms in the null hypothesis:

- ▶ object relative: +1
- ▶ subject relative: -1

Now the model is:

Object relative reading times:

$$rt = 420 \times \mathbf{1} + 51 \times \mathbf{1} + \epsilon \tag{10}$$

Subject relative reading times:

$$rt = 420 \times \mathbf{1} + 51 \times -\mathbf{1} + \epsilon \tag{11}$$

So, object relatives are coded as 1, and subject relatives are coded as -1.

The model is:

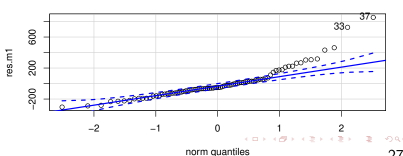
$$rt = \beta_0 + \beta_1 + \epsilon \text{ where } \epsilon \sim Normal(0, \sigma)$$
 (12)

It is an assumption of the linear model that the residuals are (approximately) normally distributed. We can check that this assumption is met:

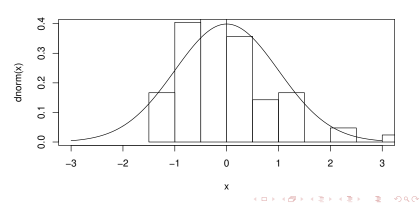
```
## residuals:
res.m1<-residuals(m1)</pre>
```

Plot the residuals by comparing them to the standard normal distribution (Normal(0,1)):

## [1] 37 33



Another way to visualize it (not the standard way):



#### Log transformation

A log-transform improves the normality of residuals:

```
m1log<-lm(log(rawRT)~cond,bysubj)
round(coef(m1log),4)

## (Intercept) cond
## 5.9488 0.0843</pre>
```

#### Log transformation

- **E**stimated **grand mean** processing time:  $\hat{\beta}_0 = 5.9488$  .
- Estimated mean object relative processing time:

$$\hat{\beta}_0 + \hat{\beta}_1 = 5.9488 + 0.0843 = 6.0331.$$

Estimated mean subject relative processing time:

$$\hat{\beta}_0 - \hat{\beta}_1 = 5.9488 - 0.0843 = 5.8645.$$

The model is:

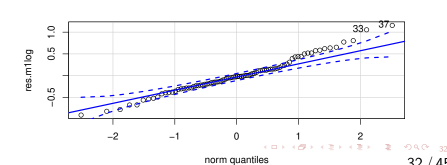
$$\log rt = \beta_0 + \beta_1 + \epsilon \tag{13}$$

Now check the residuals:

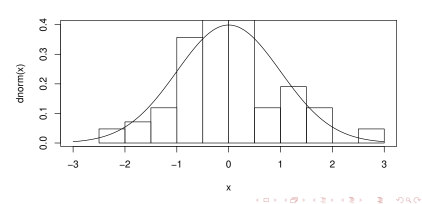
```
## residuals:
res.m1log<-residuals(m1log)</pre>
```

Plot the residuals by comparing them to the standard normal distribution (Normal(0,1)):

## [1] 37 33



Another way to visualize it (not the standard way):



Log transformation: recovering estimates on ms scale

- Estimated mean object relative processing time:  $\hat{\beta}_0 + \hat{\beta}_1 = 5.9488 + 0.0843 = 6.0331$ .
- Estimated mean subject relative processing time:  $\hat{\beta}_0 \hat{\beta}_1 = 5.9488 0.0843 = 5.8645$ .

Note that exp(log(rt)) = rt.

To get the mean estimates on the raw ms scale, we just exponentiate both sides of the equation:

$$exp(\log rt) = exp(\beta_0 + \beta_1)$$

Log transformation: recovering estimates on ms scale

- Estimated mean object relative processing time:  $exp(\hat{\beta}_0 + \hat{\beta}_1) = exp(5.9488 + 0.0843) = 417.$
- Estimated mean subject relative processing time:  $exp(\hat{\beta}_0 \hat{\beta}_1) = exp(5.9488 0.0843) = 352.$

The difference in reading time is 417-352=65 ms (cf. 102 ms on raw scale).

# Linear models: Summary

#### Summary

Using the relative clause example, we learnt

- the meaning of treatment contrast coding.
- ▶ the meaning of sum contrast coding. This is the coding we will use.

For a more comprehensive discussion of contrast coding, read this paper:

How to capitalize on a priori contrasts in linear (mixed) models: A tutorial, Schad, Hohenstein, Vasishth, Kliegl. Download from: https://arxiv.org/abs/1807.10451

### Paired t-tests vs linear models

The linear mixed model

The paired t-test and the linear model t-test values don't match:

### Paired t-tests vs linear models

The linear mixed model

This is because the linear model implements the **unpaired** (i.e., two sample) t-test:

```
round(summary(m0)$coefficients,2)[,c(1:3)]
##
                     Estimate Std. Error t value
  (Intercept)
                      471.36
                                   31.13 15.14
## conditionsubjgap -102.29 44.02 -2.32
round(t.test(rawRT~condition,bysubj,
             paired=FALSE) $statistic, 2)
##
## 2.32
                                 <□ > <□ > < = > < = > < = > < > < ( >
```

### Paired t-tests vs linear models

The linear mixed model

The paired t-test has an equivalent in the linear modeling framework:

```
m0.lmer<-lmer(rawRT~condition+(1|subject),bysubj)
summary(m0.lmer)$coefficients

## Estimate Std. Error t value
## (Intercept) 471.36 31.128 15.1428
## conditionsubjgap -102.29 32.896 -3.1093
```

We turn to the linear mixed model in the next lecture.

- 1. In our relative clause example, the 'predictor' is categorial.
- 2. What about when we have continuous predictors?
- 3. For example, we have instructors' "beauty" levels measured on a continuous scale as predictors of their teaching evaluations.
- 4. Beauty levels are centered; this means that a beauty level of 0 means average beauty level. This is a data set from a paper by Hamermesh and Parker (Beauty in the Classroom: Instructors' Pulchritude and Putative Pedagogical Productivity," Economics of Education Review, August 2005). I got the data from Gelman and Hill (2007).

```
bdata <- read.table("data/beauty.txt",header=T)</pre>
head(bdata)
##
     beauty evaluation
## 1 0.20157
                    4.3
                   4.5
## 2 -0.82608
## 3 -0.66033
                  3.7
                 4.3
## 4 -0.76631
## 5 1.42145
                  4.4
## 6 0.50022
                    4.2
plot(evaluation~beauty,bdata)
```

Note that the beauty scores are centered to have mean (approximately) 0:

```
summary(bdata$beauty)
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -1.5388 -0.7446 -0.1564 -0.0883 0.4572 1.8817
```

One model we can fit:

$$y = \beta_0 + \epsilon$$

```
m2<-lm(evaluation~1,bdata)
mean(bdata$evaluation)

## [1] 3.9983

round(summary(m2)$coefficients,digits=3)

## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.998 0.026 155.05 0
```

This model is only estimating the grand mean of evaluation scores.

$$y = \beta_0 + \beta_1 x + \epsilon$$

```
m3<-lm(evaluation~beauty,bdata)
round(summary(m3)$coefficients,digits=3)

## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.010 0.026 157.205 0

## beauty 0.133 0.032 4.133 0
```

The intercept now means: the expected evaluation score given an average beauty level.

The slope means: the expected increase in evaluation with **one unit** increase in beauty.

# Summary

We now know how to fit

- 1. Simple linear models with a categorical predictor (relative clause data)
- 2. Simple linear models with a continuous predictor.