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Learning objectives and outline

Intro: hierarchical and longitudinal data

Fecal Fat example

Correlations within subjects

Random and fixed effects

## Session 9: Repeated Measures and Longitudinal Analysis I

Levi Waldron

CUNY SPH Biostatistics 2

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#### **Learning objectives**

#### Learning objectives:

- 1 Identify and define hierarchical and longitudinal data
- 2 Analyze correlated data using Analysis of Variance
- 3 Define and calculate Intraclass Correlation
- 4 Identify and define random and fixed effects

#### Textbook sections:

• Vittinghoff sections 7.1 (7.2-7.3 next class)

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#### **Outline**

- 1 Introduction to hierarchical and longitudinal data
- 2 Fecal Fat example
- 3 Correlations within subjects (ICC)
- 4 Random and fixed effects

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### Intro: hierarchical and longitudinal data

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## What are hierarchical and longitudinal data?

- Knee radiographs are taken yearly in order to understand the onset of osteoarthritis
- An indicator of heart damage is measured at 1, 3, and 6 days following a brain hemorrhage.
- Groups of patients in a urinary incontinence trial are assembled from different treatment centers
- Susceptibility to tuberculosis is measured in family members
- A study of the choice of type of surgery to treat a brain aneurysm either by clipping the base of the aneurysm or implanting a small coil. The study is conducted by measuring the type of surgery a patient receives from a number of surgeons at a number of different institutions.

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# What is the distinction between hierarchical and longitudinal data?

- Longitudinal data are repeated measures over time
- Longitudinal data are a type of hierarchical data
  - repeated measures are correlated, and nested within the observational unit (individual)
- Other non-longitudinal data can also be hierarchical

Definition: Hierarchical data are data (responses or predictors) collected from or specific to different levels within a study.

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## Important features of this type of data

- 1 The outcomes are correlated across observations
- 2 The predictor variables can be associated with different levels of a hierarchy. *e.g.* we might be interested in:
  - the volume of operations at the hospital,
  - whether it is a for-profit or not-for-profit hospital,
  - years of experience of the surgeon or where surgeons were trained,
  - how the choice of surgery type depends on the age and gender of the patient.

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#### A Repeated Measures Example

- Lack of digestive enzymes in the intestine can cause bowel absorption problems.
  - This will be indicated by excess fat in the feces.
  - Pancreatic enzyme supplements can alleviate the problem.
  - fecfat.csv: a study of fecal fat quantity (g/day) for individuals given each of a placebo and 3 types of pills

**Table 7.1** Fecal fat (g/day) for six subjects

number         None         Tablet         Capsule         Coated         Average           1         44.5         7.3         3.4         12.4         16.9           2         33.0         21.0         23.1         25.4         25.6           3         19.1         5.0         11.8         22.0         14.5           4         9.4         4.6         4.6         5.8         6.1           5         71.3         23.3         25.6         68.2         47.1           6         51.2         38.0         36.0         52.6         44.5	ct
2     33.0     21.0     23.1     25.4     25.6       3     19.1     5.0     11.8     22.0     14.5       4     9.4     4.6     4.6     5.8     6.1       5     71.3     23.3     25.6     68.2     47.1       6     51.2     38.0     36.0     52.6     44.5	age
3     19.1     5.0     11.8     22.0     14.5       4     9.4     4.6     4.6     5.8     6.1       5     71.3     23.3     25.6     68.2     47.1       6     51.2     38.0     36.0     52.6     44.5	
4 9.4 4.6 4.6 5.8 6.1 5 71.3 23.3 25.6 68.2 47.1 6 51.2 38.0 36.0 52.6 44.5	
5 71.3 23.3 25.6 68.2 47.1 6 51.2 38.0 36.0 52.6 44.5	
6 51.2 38.0 36.0 52.6 44.5	
D111 .	
Pill type	
average 38.1 16.5 17.4 31.1 25.8	

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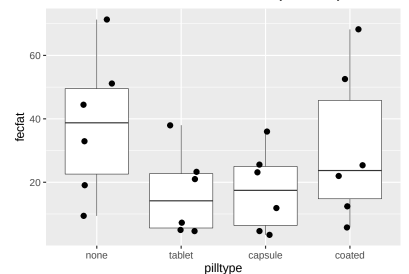
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## Option 1: non-hierarchical analysis (wrong)



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## Option 1: non-hierarchical analysis (wrong)

fit1way <- lm(fecfat ~ pilltype, data=dat)</pre>

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
pilltype	3	2008.60	669.53	1.86	0.1687
Residuals	20	7193.36	359.67		

Table 1: One-way analysis of variance table for fecal fat dataset

- Does not account for similarity of measurements within individual
- Would be correct if each treatment were given to a different individual

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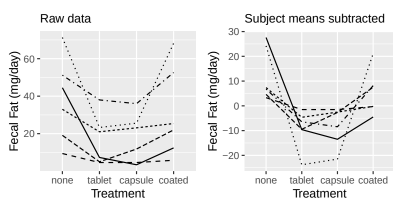
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### Option 2: 2-way AOV

- Accounts for individual differences in mean fecal fat
- Fits a coefficient for mean fecal fat per individual
- Getting closer



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### Option 2: 2-way AOV

fit1way <- lm(fecfat ~ pilltype, data=dat)</pre>

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
pilltype	3	2008.60	669.53	1.86	0.1687
Residuals	20	7193.36	359.67		

Table 2: One-way analysis of variance table for fecal fat dataset

fit2way <- lm(fecfat ~ subject + pilltype, data=dat)</pre>

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
subject	5	5588.38	1117.68	10.45	0.0002
pilltype	3	2008.60	669.53	6.26	0.0057
Residuals	15	1604.98	107.00		

**Table 3:** Two-way analysis of variance table. Note the similarity of the pilltype row.

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### What happened??

- 1-way ANOVA correctly estimates the effect of pill type
- However, 1-way ANOVA fails to accommodate the correlation within subjects
- 1-way ANOVA over-estimates the residual variance
  - under-estimates the significance of pill type

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## Regression models for 1 and 2-way ANOVA

Recall for ordinary multiple linear regression:

$$E[y|x] = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_p x_p$$

- $x_p$  are the predictors or independent variables
- *y* is the outcome, response, or dependent variable
- E[y|x] is the expected value of y given x
- $\beta_p$  are the regression coefficients

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## Regression models for 1 and 2-way ANOVA

One-way ANOVA (person i with pill type j):

$$FECFAT_{ij} = \text{fecal fat measurement for person i with pill typ}$$
  
=  $\mu + PILLTYPE_j + \epsilon_{ij}$ 

Two-way ANOVA:

$$FECFAT_{ij} = \mu + SUBJECT_i + PILLTYPE_j + \epsilon_{ij}$$

Assumption:  $\epsilon_{ij} \stackrel{iid}{\sim} N(0, \sigma_{\epsilon}^2)$ 

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### Correlations within subjects (ICC)

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### **Correlations within subjects**

- One-way ANOVA fails because it does not account for the correlation of measurements within-person
- How highly correlated are measurements on the same person? Consider subject i, pill types j and k:

$$corr(FECFAT_{ij}, FECFAT_{ik}) = \frac{cov(FECFAT_{ij}, FECFAT_{ik})}{sd(FECFAT_{ij})sd(FECFAT_{ik})}$$

\* This is a measure of how large the subject effect is, in relation to the error term

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### **Correlation within subjects**

$$cov(FECFAT_{ij}, FECFAT_{ik}) = cov(SUBJECT_i, SUBJECT_i)$$
  
=  $var(SUBJECT_i)$   
=  $\sigma_{subject}^2$ .(definition)

- Equality 1:
  - ullet  $\mu$  and  $\emph{pilltype}$  terms are assumed to be constant, so do not enter into covariance calculation
  - ullet residuals  $\epsilon$  are assumed to be independent
- Equality 2:
  - covariance with self is variance

Recall  $SUBJECT_i$  is the term for individual in 2-way AOV. Now  $\beta_i * subjectID$ , will later be treated as a **random variable** 

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### **Correlation within subjects**

Previous slide calculated *covariance* for numerator of correlation. Now calculate *variance* for the denominator  $(sd(FECFAT_{ij}) * sd(FECFAT_{ik}) = var(FECFAT_{ij}))$ 

$$var(FECFAT_{ij}) = var(SUBJECT_i, SUBJECT_i) + var(\epsilon_{ij})$$
  
=  $\sigma_{subject}^2 + \sigma_{\epsilon}^2$ .(definition)

- Difference is that the independent residuals do contribute to var(FECFAT<sub>ii</sub>)
- Variance is broken into componenets due to subject and residual variance

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#### **Intraclass Correlation**

The correlation between two treatments j and k across subjects i is:

$$corr(FECFAT_{ij}, FECFAT_{ik}) = \frac{cov(FECFAT_{ij}, FECFAT_{ik})}{sd(FECFAT_{ij})sd(FECFAT_{ik})}$$

$$= \frac{\sigma_{subj}^{2}}{\sigma_{subj}^{2} + \sigma_{\epsilon}^{2}}$$

$$ICC = \frac{\tau_{00}^{2}}{\tau_{00}^{2} + \sigma_{\epsilon}^{2}}$$

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## Intuition behind correlations within subjects

**Table 7.1** Fecal fat (g/day) for six subjects

Subject	Pill type				Cubicat
Subject number	None	Tablet	Capsule	Coated	Subject Average
1	44.5	7.3	3.4	12.4	16.9
2	33.0	21.0	23.1	25.4	25.6
3	19.1	5.0	11.8	22.0	14.5
4	9.4	4.6	4.6	5.8	6.1
5	71.3	23.3	25.6	68.2	47.1
6	51.2	38.0	36.0	52.6	44.5
Pill type					
average	38.1	16.5	17.4	31.1	25.8

Figure 2: Fecal Fat dataset

Variance of the subject averages (279.4) is increased by correlation of measurements within individual

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[1] 106.9989

## Calculation of correlations within subjects (ICC)

What is your estimate of the variability due to subjects, from the 2-way ANOVA?

```
sum(residuals(fit2way)^2) / 15 / 4 #df=15, divided by 4 pilltypes
## [1] 26.74972
279.419 - 26.75 #var(SUBJECT_i)
## [1] 252.669
Residual variance is:
sum(residuals(fit2way)^2) / 15 #df=15
```

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## Calculation of correlations within subjects (ICC)

Finally calculate ICC:

$$ICC = \frac{\sigma_{subj}^2}{\sigma_{subj}^2 + \sigma_{\epsilon}^2}$$
$$= \frac{253}{253 + 107} = 0.70$$

This calculation will become easier when we learn to estimate *random coefficients* in directly in the regression model.

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## The next step: a mixed effects model

• Two-way ANOVA is a fixed effects model:

$$FECFAT_{ij} = \beta_0 + \beta_{subjecti}SUBJECT_i + \beta_{pilltypej}PILLTYPE_j + \epsilon_{ij}$$

- Assumption:  $\epsilon_i \stackrel{iid}{\sim} N(0, \sigma_{\epsilon}^2)$
- Instead of fitting a  $\beta_{subjecti}$  to each individual, assume that subject effects are selected from a distribution of possible subject effects:

$$FECFAT_{ij} = \mu + SUBJECT_i + \beta_{pilltypej}PILLTYPE_j + \epsilon_{ij}$$
  
where  $SUBJECT_i \stackrel{iid}{\sim} N(0, \sigma_{cubi}^2)$ 

- Here subject is a *random* effect, and pill type is a *fixed* effect.
- This is also a random intercept model

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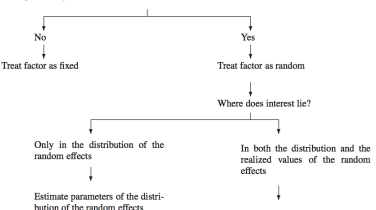
Random and fixed effects

#### Random and fixed effects

7.6 Re-Analysis of the Georgia Babies Data Set

Table 7.14 Decision tree for deciding between fixed and random

Is it reasonable to assume levels of the factor come from a probability distribution?



Estimate parameters of the distribution of the random effects

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## Summary: correlations within subjects

- Subject-to-subject variability simultaneously raises or lowers all the observations on a subject
  - induces correlation of within-subject measurements
- Variability of individual measurements can be separated into that due to subjects and that left to residual variance.
  - $var(FECFAT_{ij}) = \sigma_{subj}^2 + \sigma_{\epsilon}^2$
- 2-way ANOVA does not directly estimate variability due to subjects
  - variance of coefficients for individual is not too far off

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### Summary: hierarchical data

- Estimates of coefficients (or "effect sizes") are unchanged by hierarchical modeling
- Ignoring within-subject correlations results in incorrect estimates of variance, F statistics, p-values
  - not always "conservative"
- Intraclass Correlation (ICC) provides a measure of correlation induced by grouping
- Should be able to recognize fixed and random effects