BIOS 755: Introductory concepts

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Outline

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Introduction

Lecture goals:

- ▶ Give an overview of the objectives of longitudinal analysis and discuss features in the data.
- Longitudinal analysis is concerned with estimating how individuals change throughout the study.
- Examine the factors that are related to differences among individuals over time.
- Review features of longitudinal study designs.
- ▶ Introduce notation.

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Multicenter AIDS Cohort Study of HIV

- ▶ A cohort of 369 men was followed before and after HIV sero-conversion (which is the development of detectable specific antibodies to microorganisms in the blood serum as a result of infection or immunization)
- ▶ Important indicator of immune function is the CD4+ cell count
- ► CD4+ count was taken on each subject approximately every six months
- ▶ Over all 2376 observations

Objectives of Longitudinal Analysis

- ▶ Defining feature of longitudinal study: two or more observations taken on (at least) some subjects.
- ▶ Multiple measurements over time allow assessment of within-individual changes in the response variable.
- ▶ Thus, some main goals are to:
 - characterize (a loaded word) the change in the response over time.
 - determine whether changes are related to exposures of interest

Longitudinal Analysis

- ▶ One way to achieve this is to look at "change scores" or "difference scores," between pre- and post-treatment.
- ► There are different ways to form such a question
 - 1. are the *change scores* related to group.
 - 2. is the change in the scores related to covariates.
 - 3. is the final score related to covariates (path analysis).
- ▶ 1 could be answered using a paired t-test.
- ▶ 2 could be answered using standard linear regression techniques (possibly adjusting for the pre-score and differences in time).
- ▶ 3 also could use linear regression (adjusting for the pre-score and differences in time).

Longitudinal Analysis

- ► The usefulness of change score analysis is limited to situations with two measurements, with more than two measurements you will have more than 1 change score.
- Analyses with change scores should be used with caution as they
 - ► have been shown to lead to bias in observational (non-randomized) studies (reference), and
 - require complete data or multiple imputation.

Number of measurements

- ► The number of measurements can vary greatly from study to study and can be equally or unequally separated in time.
 - Pain measured every 15 minutes for 2 hours.
 - ▶ Height measured every 3-months until 3 yr old, then every year thereafter.
- ▶ Both of the above are considered to be **balanced**.
- Unbalanced data are very common in the health sciences.
 - individuals will miss schedule visits,
 - visits are not made exactly at the scheduled date,
 - timings are relative to a benchmark event (e.g. relative to sero-conversion), or
 - visits are themselves random (common in retrospective data).
- Missing data are the rule, not the exception, so unbalanced data are the rule.

Treatment of Lead-Exposed Chidren (TLC) Trial

Recall the TLC study (a balanced design)

- Exposure to lead during infancy is associated with substantial deficits in tests of cognitive ability
- ► Chelation treatment of children with high lead levels usually requires injections and hospitalization
- A new agent, Succimer, can be given orally
- Randomized trial examining changes in blood lead level during course of treatment
- ▶ 100 children randomized to placebo or succimer
- ▶ Measures of blood lead level at baseline, 1, 4 and 6 weeks

Data Notation

► Consider the following

 Y_{ij} = the jth measurement taken on unit i.

where
$$i = 1, 2, ..., N$$
 and $j = 1, 2, ..., n$

- ▶ In the TLC data each child had four measurements at baseline, 1, 4 and 6 weeks.
- ▶ Y_{ij} represents the **random** response of the *i*th child at measurement j, for j = 1, 2, 3, 4.
- y_{ij} denotes the realized value of Y_{ij} .
- $ightharpoonup t_{ij}$ is the time of the observation of the *i*th child at measurement *j*, for all *i*

$$t_{i1}=0, \quad t_{i2}=1, \quad t_{i3}=4, \quad t_4=6$$

Random Vectors

- ▶ It is convenient to represent all observations for a specific unit as a **random** vector.¹
- ▶ For the TLC data each child has,

$$oldsymbol{Y}_i = \left(egin{array}{c} Y_{i1} \ Y_{i2} \ Y_{i3} \ Y_{i4} \end{array}
ight)$$

the random vector for child i.

- Also use $Y_i = (Y_{i1}, Y_{i2}, Y_{i3}, Y_{i4})'$.
- ▶ In general, we have $\mathbf{Y}_i = (Y_{i1}, Y_{i2}, \dots, Y_{in})^t$.

¹see Gentle Introduction to Vectors and Matrices in appendix A

Expectations and mean

▶ The mean, average, or "expectation" of each response is

$$E(Y_{ij}) = \mu_{ij}$$

 μ_{ij} is the *conditional* mean at the *j*th occasion (i.e., conditional on covariate values).

Dependence and Correlation Introduction

- ► Two variables are said to be independent if the behavior of one variable does not depend on the value of another variable.
- ► For example, LDL cholesterol and sex are independent if the distribution of LDL cholesterol is the same for males and females.
- Longitudinal data methods do not make the assumption that the observations are independent.
- ► For example, if I tell you my LDL cholesterol at baseline was 80, what is a plausible range for this value at 4 weeks?
- What if it was 165 at baseline?

Variance and Covariance

▶ Along with the expectation we'll use variance

$$var(Y_{ij}) = E(Y_{ij} - \mu_{ij})^2 = \sigma_j^2$$

- ▶ The standard deviation is $\sqrt{\sigma_j^2} = \sigma_j$.
- ▶ Covariance: a measure of how two random variables vary together.
- Mathematically this is expressed as,

$$cov(Y_{ij}, Y_{ik}) = E\{(Y_{ij} - \mu_{ij})(Y_{ik} - \mu_{ik})\} = \sigma_{jk}$$

Covariance Matix

▶ The covariance matrix of $Y_i = (Y_{i1}, Y_{i2}, ..., Y_{in})'$

$$\mathsf{Cov}(oldsymbol{Y}_i) = \left(egin{array}{cccc} \sigma_1^2 & \sigma_{12} & \dots & \sigma_{1n} \ \sigma_{21} & \sigma_2^2 & \dots & \sigma_{2n} \ dots & dots & \ddots & dots \ \sigma_{n1} & \sigma_{n2} & \dots & \sigma_n^2 \end{array}
ight) = oldsymbol{\Sigma}$$

▶ You will sometimes see $\sigma_j^2 = \text{var}(Y_{ij}) = \text{cov}(Y_{ij}, Y_{ij}) = \sigma_{jj}$.

Covariance to correlation

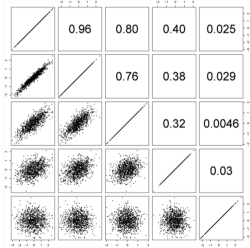
- ► The covariance values are hard to interpret since their magnitude is dependent on the variance of the variables.
- Covariance is commonly standardized to correlation.
- The population correlation of two elements is

$$\rho_{jk} = \frac{\sigma_{jk}}{\sqrt{\sigma_j^2 \sigma_k^2}}$$

Which gives the correlation matrix

$$\mathsf{Corr}(\boldsymbol{Y}_i) = \begin{pmatrix} 1 & \rho_{12} & \dots & \rho_{1n} \\ \rho_{21} & 1 & \dots & \rho_{2n} \\ \vdots & \vdots & \ddots & \vdots \\ \rho_{n1} & \rho_{n2} & \dots & 1 \end{pmatrix}$$

Correlation Matix Example



Objectives of TLC Trial

- ► Goal: determine whether the new treatment reduces blood lead levels over time relative to placebo.
- Let μ_{jS} and μ_{jP} are the mean levels at occasion j for succimer and placebo groups, respectively.
- ▶ Different ways to answer this question:
 - 1. $H_0: \mu_{jS} = \mu_{jP}$ for all j = 1, ..., 4.
 - 2. $H_0: \mu_{jS} \mu_{1S} = \mu_{jP} \mu_{1P}$ for j = 2, 3, 4.

Correlation in TLC

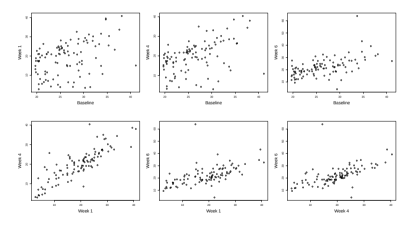


Figure: Correlation for 50 children in the placebo group

Estimated covariance & correlation matrices

▶ The estimated covariance matrix for the TLC

$$\mathsf{Cov}(\boldsymbol{Y}_i) = \begin{pmatrix} 25.2 & 22.8 & 24.3 & 21.4 \\ 22.8 & 29.8 & 27.0 & 23.4 \\ 24.3 & 27.0 & 33.1 & 28.2 \\ 21.4 & 23.4 & 28.2 & 31.8 \end{pmatrix}$$

▶ The estimated correlation matrix for the TLC

$$\mathsf{Corr}(\boldsymbol{Y}_i) = \begin{pmatrix} 1 & 0.83 & 0.84 & 0.76 \\ 0.83 & 1 & 0.86 & 0.76 \\ 0.84 & 0.86 & 1 & 0.87 \\ 0.76 & 0.76 & 0.87 & 1 \end{pmatrix}$$

Time plot for TLC

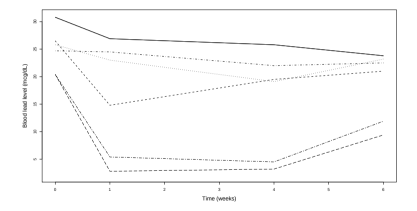


Figure: Time plot for 6 subjects

Correlation "truths"

- 1. the correlations are positive
- 2. the correlations often decrease with increasing time separation
- 3. the correlations between repeated measures rarely ever approach zero
- 4. the correlation between a pair of repeated measures taken very closely rarely approaches one.

Sources of variability

Three potential sources of variability

- between-individual heterogeneity
- within-individual biological variation
- measuement error