# Repeated Measures and Longitudinal Data Analysis II

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#### Welcome and outline - session 10

- Learning objectives
  - identify hierarchical data
  - define mixed effects models and population average models
  - perform model diagnostics for random effects models
  - interpret random intercepts and random slopes
- ▶ Vittinghoff sections 8.2, 8.3, 8.5

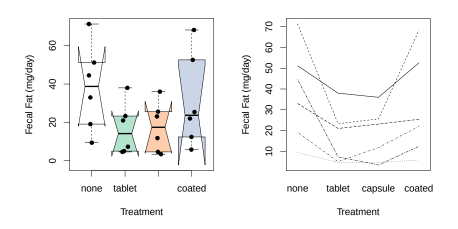
## Recall the simple repeated measures example: fecal fat

- ► 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

Subject	Pill type	Subject			
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
Pill type					
average	38.1	16.5	17.4	31.1	25.8

### Fecal fat dataset



### Analysis strategies for hierarchical data

- ► Analyses for each subgroup
  - e.g., look at each patient independently
  - doesn't work at all in this example, and in general is not an integrated analysis of the whole data
  - could sort of work for an example with many patients per doctor, a few doctors
- Analysis at the highest level in the hierarchy
  - ▶ first summarize data to highest level
  - doesn't work at all in this example
  - could sort of work for an example with many patients per doctor, a few doctors
- Analysis on "Derived Variables"
  - consider each treatment type separately, take differences in fat levels between treatment/control for each patient and use paired t-tests
  - can work, but not work for unbalanced groups (e.g. incomplete data for some participants)

# Better analysis strategies for hierarchical data

- Fixed effects models
- ► Random / mixed effects models
  - model certain regression coefficients (intercept, slopes) as random variables
- ► Population average models
  - using Generalized Estimating Equations (GEE)

## When is hierarchical analysis needed?

- ► Hierarchical analysis strategies are needed:
  - 1. when the correlation structure is of primary interest, *e.g.* familial aggregation of disease, or consistency of treatment within centers,
  - 2. when we wish to "borrow strength" across the levels of a hierarchy in order to improve estimates, and
  - 3. when dealing with unbalanced, correlated data.

## Fixed effects: two-way analysis of variance

► Two-way ANOVA (person *i* with pill type *j*):

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

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

	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 1: Two-way analysis of variance table. Equivalent to subtracting the mean fecal fat content for each individual.

\* Accounts for individual differences in mean fecal fat \* Fits a coefficient for mean fecal fat per individual

#### Mixed effects model

► Model looks like two-way ANOVA:

$$\textit{FECFAT}_{ij} = \beta_0 + \beta_{\textit{subjecti}} \textit{SUBJECT}_i + \beta_{\textit{pilltypej}} \textit{PILLTYPE}_j + \epsilon_{ij}$$

- ► Assumption:  $\epsilon_i \stackrel{iid}{\sim} N(0, \sigma_{\epsilon}^2)$
- ightharpoonup But instead of fitting a eta to each individual, we assume that the subject effects are selected from a distribution of possible subject effects:

$$\textit{FECFAT}_{ij} = \beta_0 + \textit{SUBJECT}_i + \beta_{\textit{pilltypej}} \textit{PILLTYPE}_j + \epsilon_{ij}$$

Where we assume:  $SUBJECT_i \stackrel{iid}{\sim} N(0, \tau_{00}^2)$ 

- ▶ This is a *mixed effects* model because:
  - ▶ the "true" intercept varies randomly from patient to patient
  - the "true" (population) coefficient of treatment is fixed (the same for everyone)

### Mixed effects model coefficients, variances, ICC

```
## Linear mixed-effects model fit by REML
     Data: fecfat
   Log-restricted-likelihood: -84.55594
   Fixed: fecfat ~ pilltype
##
       (Intercept) pilltypetablet pilltypecapsule pilltypecoated
         38.083334
                        -21.550001
                                        -20.666667
##
                                                          -7.016668
##
## Random effects:
## Formula: ~1 | subject
           (Intercept) Residual
## StdDev:
              15 89557 10 34403
##
## Number of Observations: 24
## Number of Groups: 6
ICC = 15.9^{2}/(15.9^{2} + 10.34^{2}) = 0.7 = 0.7.
```

- Recall ICC is a measure of how large the subject effect is, in relation to the error term
- Variances were estimated directly by the model!

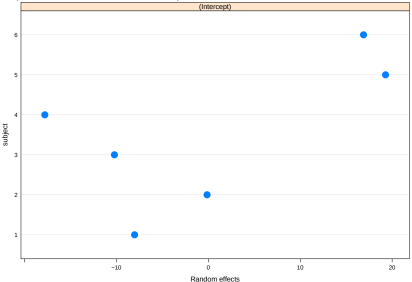
# Assumptions of the mixed model

$$\textit{FECFAT}_{ij} = \beta_0 + \textit{SUBJECT}_i + \beta_{\textit{pilltypej}} \textit{PILLTYPE}_j + \epsilon_{ij}$$

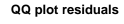
- Normally distributed residuals as in fixed effects model:
- Normally distributed latent variable:
  - ► SUBJECT<sub>i</sub>  $\stackrel{iid}{\sim} N(0, \tau_{00}^2)$

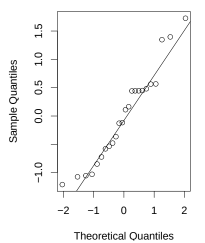
# Mixed effects model results (cont'd)

A plot of the random intercept:

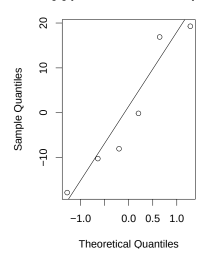


# Mixed effects model diagnostics





#### QQ plot random intercepts



#### Mixed effects model results

```
## Linear mixed-effects model fit by REML
## Data: fecfat
         ATC
                  BIC
                         logLik
    181.1119 187.0863 -84.55594
##
## Random effects:
## Formula: ~1 | subject
         (Intercept) Residual
## StdDev: 15.89557 10.34403
##
## Fixed effects: fecfat ~ pilltype
##
                      Value Std.Error DF t-value p-value
## (Intercept)
                 38.08333 7.742396 15 4.918805 0.0002
## pilltypetablet -21.55000 5.972127 15 -3.608430 0.0026
## pilltypecapsule -20.66667 5.972127 15 -3.460521 0.0035
## pilltypecoated -7.01667 5.972127 15 -1.174903 0.2583
## Correlation:
##
                  (Intr) plltypt plltypcp
## pilltypetablet -0.386
## pilltypecapsule -0.386 0.500
## pilltypecoated -0.386 0.500 0.500
##
## Standardized Within-Group Residuals:
##
           Min
                         01
                                     Med
                                                   03
                                                              Max
## -1.210052934 -0.615068039 -0.002727166 0.457105344 1.725618643
##
## Number of Observations: 24
## Number of Groups: 6
```

# Mixed effects model results (cont'd)

- ▶ Note on correlation of the estimator of the fixed effects
  - high correlations may (but not necessarily) be due to collinearity
  - not usually useful, not included in output of some packages

## Mixed effects model results (cont'd)

Inference for variance terms (and fixed effects):

```
## Approximate 95% confidence intervals
##
   Fixed effects:
##
                      lower
                                            upper
## (Intercept)
                  21.58081 38.083334 54.585860
## pilltypetablet -34.27929 -21.550001 -8.820714
## pilltypecapsule -33.39595 -20.666667 -7.937381
## pilltypecoated -19.74595 -7.016668 5.712618
## attr(,"label")
## [1] "Fixed effects:"
##
   Random Effects:
    Level: subject
##
                     lower
                              est.
                                        upper
## sd((Intercept)) 8.001171 15.89557 31.57904
##
   Within-group standard error:
##
       lower
  7.232404 10.344027 14.794374
```

- Would conclude that variation of the intercept between subjects is non-zero
  - not attributable to within-subject variation

### Longitudinal data

- Interested in the change in the value of a variable within a "subject"
- Collect data repeatedly through time.
- For hierarchical longitudinal analysis to be effective,
   before/after measurements need to be positively correlated

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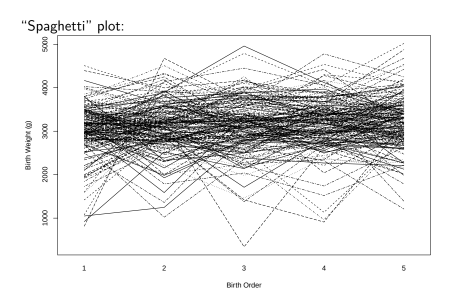
### Longitudinal data examples

- ► Example 1: a measure of sleepiness before and after administration of treatment or placebo
- Example 2: Study of Osteoporotic Fractores (SOF dataset)
  - ▶ 9,704 women tracked with clinical visits every two years
  - Bone Mineral Density (BMD), Body Mass Index (BMI), many other variables
- Questions for Example 2:
  - Is change in BMD related to age at menopause? This is a time-invariant predictor, age at menopause, with time-dependent changes in the outcome, BMD.
  - 2. Is change in BMD related to change in BMI? This is an analysis relating a time-varying predictor, BMI, with changes in the outcome, BMD. BMI varies quite a lot between women, but also varies within a woman over time.

# Longitudinal data examples (cont'd)

- birthweight and birth order
- provides birthweights and order of infants from mothers who had 5 children in Georgia
  - interested in whether birthweight of babies changes with order
  - whether this difference depends on the mother's age at first childbirth or on the weight of initial baby.

# Georgia Birthweights dataset

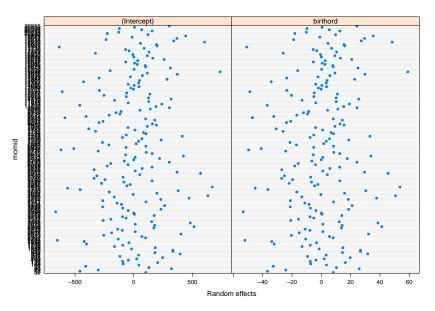


### Georgia Birthweights dataset

- Do birth weights or the effect of birth order vary by mother?
  - random intercept, random slope

Note: the control argument increases the max # iterations to allow convergence. If msVerbose=TRUE, it produces verbose output.

# Georgia Birthweights dataset (cont'd)



# Georgia Birthweights dataset (cont'd)

summary(gafit1)

```
## Linear mixed-effects model fit by REML
## Data: ga
##
          ATC
                  BIC
                        logLik
   15321.23 15350.67 -7654.616
##
## Random effects:
## Formula: ~birthord | momid
## Structure: General positive-definite, Log-Cholesky parametrization
##
               StdDev
                        Corr
## (Intercept) 296.36304 (Intr)
## birthord 23,82664 0.996
## Residual
              443 43313
##
## Fixed effects: bweight ~ birthord
##
                 Value Std.Error DF t-value p-value
## (Intercept) 2995.640 38.99534 799 76.82046
## birthord
                46.608 10.05758 799 4.63411
## Correlation:
           (Intr)
##
## birthord -0.662
##
## Standardized Within-Group Residuals:
##
           Min
                       Q1
                                  Med
                                                          Max
## -5.32850276 -0.42195082 0.04103012 0.53113344 3.25512201
##
## Number of Observations: 1000
## Number of Groups: 200
```

# Georgia Birthweights dataset (cont'd)

```
intervals(gafit1)
```

```
## Approximate 95% confidence intervals
##
   Fixed effects:
##
                    lower
                              est.
                                        upper
## (Intercept) 2919.09459 2995.640 3072.18541
## birthord
                 26.86559
                            46.608
                                     66.35041
## attr(,"label")
## [1] "Fixed effects:"
##
   Random Effects:
   Level: momid
##
                                  lower
                                                        upper
## sd((Intercept))
                             227.421283 296.3630360 386.20418
## sd(birthord)
                               9.351472 23.8266383 60.70795
## cor((Intercept),birthord) -1.000000 0.9964047 1.00000
##
   Within-group standard error:
      lower
                est.
                        upper
## 422 2138 443 4331 465 7189
```

- Do birth weights or the effect of birth order vary by mother?
  - yes: both standard deviations are non-zero

### Population Average Models

- ► An alternative to random / mixed-effects models that is more robust to assumptions of:
  - distribution of random effects
  - correlation structure
- Estimates correlation structure from the data rather than assuming normality
  - Requires a fair bit more clusters than observations per cluster
- Estimates regression coefficients and robust standard errors
  - commonly by Generalized Estimating Equations (GEE)

# Population Average Models

Compare mixed model multiple linear regression:

$$E[Y_{ij}|X_{ij}] = \beta_0 + \alpha_{0j} + \beta_1 X_{ij}, \alpha_{0j} \sim N(0, \sigma)$$

for subject i in group j.

to a population average model:

$$E[Y_{ij}|X_{ij}] = \beta_0^* + \beta_1^* X_{ij}$$

- ▶ Interpretations of  $\beta^*$  and  $\beta$  are equivalent
- Numerically equivalent for linear and log-linear models (if specification of mixed model is correct), but not for logistic link.

### Fit a population average model

```
gafit.gee <- gee::gee(bweight ~ birthord, corstr="exchangeable",
                     id=momid, data=ga)
summary(gafit.gee)
##
## GEE: GENERALIZED LINEAR MODELS FOR DEPENDENT DATA
   gee S-function, version 4.13 modified 98/01/27 (1998)
##
## Model:
## Link:
                              Identity
## Variance to Mean Relation: Gaussian
## Correlation Structure:
                              Exchangeable
##
## Call:
## gee::gee(formula = bweight ~ birthord, id = momid, data = ga,
      corstr = "exchangeable")
##
##
## Summary of Residuals:
##
        Min
                         Median
                                       30
                                                Max
                   10
## -2795 464 -299 126 48 840 341 144 1824 536
##
##
## Coefficients:
              Estimate Naive S.E. Naive z Robust S.E. Robust z
## (Intercept) 2995.640 41.973695 71.369462 38.808066 77.191170
## hirthord 46 608 9 958128 4 680398 9 996256 4 662546
##
## Estimated Scale Parameter: 332525.3
## Number of Iterations: 1
##
## Working Correlation
##
            [.1]
                      [,2]
                                Γ.31
                                          Γ.47
                                                    Γ.51
## [1.] 1.0000000 0.4035684 0.4035684 0.4035684 0.4035684
```

# GEE working correlation types

- Must make some assumption about form of correlation among observations from the same subject, same hospital, etc
- Independence
  - no correlation between measurements within group
- Exchangeable
  - all correlations (except those variables with themselves) are a common value
  - nothing to distinguish one member of a cluster from another
  - appropriate in the absence of other data structures such as measurements taken through time or space
- Auto-regressive
  - observations taken more closely in time are more highly correlated

## GEE working correlation types

- Unstructured
  - estimates a separate correlation between observations taken on each pair of "times"
- Non-stationary
  - similar to unstructured, but assumes all correlations for pairs separated far enough in time are zero
- Stationary
  - e.g. stationary of order 2: observations taken at time points 1 and 3 have the same correlation as time points 2 and 4
  - but this might be different from the correlation between observations taken at times 2 and 3
  - correlations for observations 3 or more time periods apart assumed to be zero

Fewer assumptions requires more data, and good assumptions improve results

# Help in choosing a method

Characteristic	Marginal	Fixed-effect	Mixed-effect
Distinguishes observations belonging to the same or different subjects	Yes <sup>a</sup>	Yes	Yes
Reliant on distribution of subject-specific effects	No	No	Yes
Subjects considered a sample from a population larger than the sample itself	Yes <sup>a</sup>	No	Yes
Computation handles few subjects well	No	Yes	No
Computation handles a very large number of subjects well	Yes	No	Yes
Noisy for few observations per subject	No	Yes	No
Computation handles a large number of observations per subject	Depends <sup>b</sup>	Yes	Yes
Accommodates variable observations per subject	Yes	Yes	Yes

Note: aOnly for calculation of standard errors.

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Figure 2: decision table

<sup>&</sup>lt;sup>b</sup>Problems can arise under some specifications of the working covariance structure and depending on the estimation method used.

#### Conclusions

- Ignoring within-subject correlations can produce very wrong results, and is not always "conservative"
- Hierarchical analysis strategies are needed for any of:
  - When the correlation structure is of primary interest, e.g. familial aggregation of disease, or consistency of treatment within centers,
  - 2. When we wish to "borrow strength" across the levels of a hierarchy in order to improve estimates, and
  - 3. When dealing with unbalanced correlated data. E.g., no requirement that each Georgia mother have exactly 5 children.
- Population average models provide a robust alternative to mixed models
  - ► for one level of hierarchy

#### Lab exercise

For the fecal fat dataset: 1. Fit a linear model with random coefficients for pills, and summarize the output 2. Estimate confidence intervals for the coefficients of this model, and interpret them 3. Calculate the ICC 4. Create residuals plots for this model and interpret 5. Fit population average models for the fecal fat dataset using at least two different working correlation structures. Compare the results to each other and to the mixed model