# Variance-Covariance Matrices

Daniel Andersor Week 4

### Agenda

- Review Homework 1
- Review the most important parts of Week 3 content
- Discuss Gelman and Hill notation contrast with Raudenbush and Bryk
- Unstructured VCV Matrices and alternatives

### Learning Objectives

- Understand at least the basics of the GH notation and why I view it as preferable
- Gain a deeper understanding of how the residual structure is different in multilevel models
- Understand that there are methods for changing the residual structure, and understand when and why this might be preferable
- Have a basic understanding of implementing alternative methods

## Review Homework 1

# Review Week 3 content

### Data and model

extrav is a measure of extraversion.

What is this model fitting, in plain English?

### Model summary

#### arm::display(m)

```
## lmer(formula = popular ~ extrav + (extrav | class), data = popular,
## control = lmerControl(optimizer = "bobyga"))
##
             coef.est coef.se
## (Intercept) 2.46 0.20
## extrav 0.49 0.03
##
## Error terms:
## Groups Name Std.Dev. Corr
## class (Intercept) 1.73
    extrav 0.16 -0.97
##
## Residual
                    0.95
## ---
## number of obs: 2000, groups: class, 100
\#\# AIC = 5791.4, DIC = 5762
## deviance = 5770.7
```

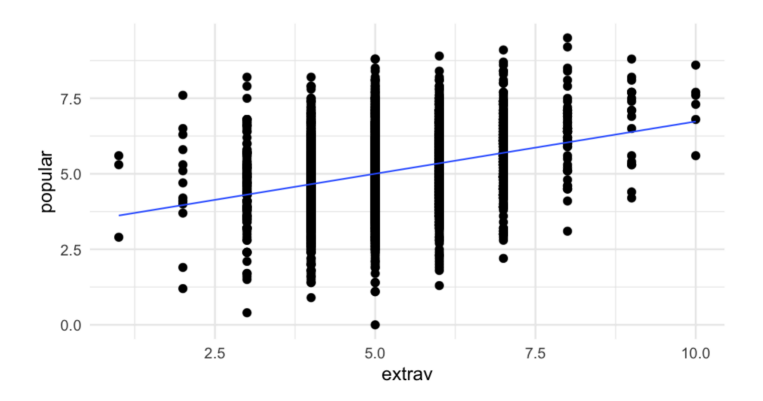
### Let's walk through

• By way of thinking through it, let's compare to simple linear regression

```
slr <- lm(popular ~ extrav, popular)</pre>
```

### Visually

```
ggplot(popular, aes(extrav, popular)) +
  geom_point() +
  geom_smooth(se = FALSE, method = "lm")
```



### Making predictions

$$\widehat{\text{popular}} = 3.27 + 0.35(\text{extrav})$$

#### Scores of 0 to 5

$$\widehat{\text{popular}} = 3.27 + 0.35 \times 0 = 3.27$$
 $\widehat{\text{popular}} = 3.27 + 0.35 \times 1 = 3.62$ 
 $\widehat{\text{popular}} = 3.27 + 0.35 \times 2 = 3.97$ 
 $\widehat{\text{popular}} = 3.27 + 0.35 \times 3 = 4.32$ 
 $\widehat{\text{popular}} = 3.27 + 0.35 \times 4 = 4.67$ 
 $\widehat{\text{popular}} = 3.27 + 0.35 \times 5 = 5.02$ 

### Now for the mlm

It's more complicated now for a couple of reasons

- Each class has their own intercept and slope. Which class is the student in?
- What if we want to make a prediction for someone outside our sample? Let's walk through 4 predictions

### Sample preds

```
sample_preds <- popular %>%
  group_by(class) %>%
  slice(1) %>%
  ungroup %>%
  slice(1:4)

sample_preds
```

```
## # A tibble: 4 x 7
## pupil class extrav sex texp popular popteach
## <dbl> 6.3
## 2 1 2 8 girl 14 6.4
## 3 1 3 5 boy 13 4.2
## 4 1 4 3 girl 20 4.100000
```

### Coefficients

$$egin{aligned} \widehat{ ext{popular}}_i &\sim N\left(2.46_{lpha_{j[i]}} + 0.49_{eta_{1j[i]}}( ext{extrav}), 0.95
ight) \ \left(egin{aligned} lpha_j \ eta_{1j} \end{aligned}
ight) &\sim N\left(\left(egin{aligned} 0 \ 0 \end{aligned}
ight), \left(egin{aligned} 1.73 & -0.97 \ -0.97 & 0.16 \end{aligned}
ight)
ight), ext{for class j} = 1, \ldots, J \end{aligned}$$

### Grab params

#### Fixed effects

#### classroom deviations

```
r <- ranef(m)
r
```

## Predictions depend on classroom

#### Fixed effect part

Works just like simple linear regression

```
sample_preds[1, ]
```

```
## # A tibble: 1 x 7
## pupil class extrav sex texp popular popteach
## <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> 6.3
```

$$\widehat{\text{popular}} = 2.46 + 0.49 \times 5 = 4.91$$

```
f[1] + f[2]*5
```

```
## (Intercept)
## 4.925309
```

### Random effects

We now have to add in the random effects for the corresponding classroom.

#### head(r\$class)

```
## (Intercept) extrav

## 1 0.3409595 -0.02701468

## 2 -1.1779895 0.09697467

## 3 -0.6293700 0.05709718

## 4 1.0852971 -0.09995304

## 5 -0.1948964 0.02160180

## 6 -0.9833923 0.08376253
```

$$\widehat{\text{popular}} = (2.46 + 0.34) + (0.49 + -0.03) \times 5 = 5.10$$

### In code

```
class1 <- r$class[1, ]
class1

## (Intercept) extrav
## 1 0.3409595 -0.02701468

(f[1] + class1[1]) + (f[2] + class1[2])*5

## (Intercept)
## 1 5.131195</pre>
```

### Predictions

#### sample\_preds

```
## # A tibble: 4 x 7
## pupil class extrav sex texp popular popteach
## < dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> 6.3
## 2 1 2 8 girl 14 6.4
## 3 1 3 5 boy 13 4.2
## 4 1 4 3 girl 20 4.100000
```

#### $head(r\c class, n = 4)$

```
## (Intercept) extrav
## 1  0.3409595 -0.02701468
## 2  -1.1779895  0.09697467
## 3  -0.6293700  0.05709718
## 4  1.0852971 -0.09995304
```

#### fixef(m)

```
## (Intercept) extrav
## 2.4610234 0.4928571
```

$$\widehat{\text{popular}} = (2.46 + 0.34) + (0.49 + -0.03) \times 5 = 5.10$$

$$\widehat{\text{popular}} = (2.46 + -1.18) + (0.49 + 0.10) \times 8 = 6.00$$

$$\widehat{\text{popular}} = (2.46 + -0.63) + (0.49 + 0.06) \times 5 = 4.58$$

$$\widehat{\text{popular}} = (2.46 + 1.09) + (0.49 + -0.10) \times 3 = 4.72$$

predict(m, newdata = sample\_preds)

What if we want to make a prediction outside of our classrooms?

Fixed effects only

$$\widehat{\text{popular}} = 2.46 + 0.49 \times \text{extraversion}$$

### Plotting

We can use the **expand.grid()** function to create different conditions. Let's compare slopes across the first five classrooms

```
conditions <- expand.grid(extrav = 1:10, class = 1:5)</pre>
```

#### head(conditions)

```
## 1 extrav class
## 1 1 1 1
## 2 2 2 1
## 3 3 1
## 4 4 1
## 5 5 5 1
## 6 6 1
```

#### tail(conditions)

```
## 45 extrav class
## 46 6 5
## 47 7 5
## 48 8 5
## 49 9 5
## 50 10 5
```

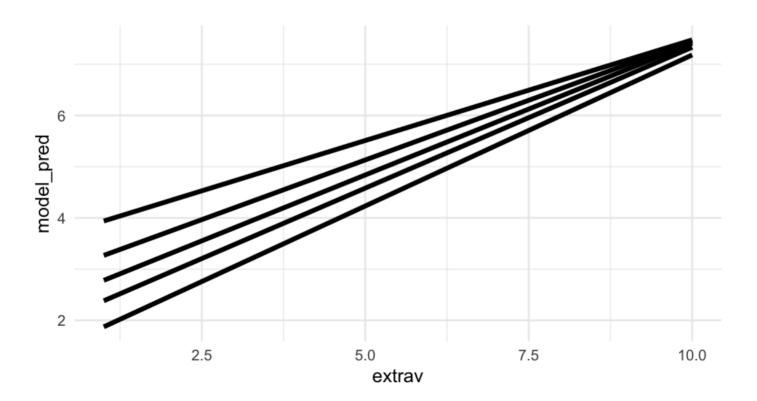
### Make predictions

```
conditions %>%
  mutate(model_pred = predict(m, newdata = conditions))
```

```
##
     extrav class model pred
## 1
                    3.267825
## 2
          2
                  3.733668
          3
## 3
                  4.199510
## 4
                1 4.665353
          5
## 5
                  5.131195
## 6
          6
                  5.597038
## 7
          7
                  6.062880
## 8
          8
                  6.528722
## 9
                  6.994565
## 10
         10
                  7.460407
                   1.872866
## 11
          2
                   2.462697
## 12
          3
                   3.052529
## 13
                   3.642361
## 14
           5
                   4.232193
## 15
          6
                   4.822025
## 16
          7
                   5.411856
## 17
          8
## 18
                   6.001688
## 19
                   6.591520
## 20
         10
                  7.181352
## 21
                    2.381608
## 22
                    2.931562
```

### Plot

```
conditions %>%
  mutate(model_pred = predict(m, newdata = conditions)) %>%
  ggplot(aes(extrav, model_pred)) +
  geom_line(aes(group = class))
```



### One more quick example

Model an interaction

### Model summary

#### arm::display(m2)

```
## lmer(formula = popular ~ extrav * sex + (extrav | class), data = popular
##
     control = lmerControl(optimizer = "bobyga"))
##
              coef.est coef.se
## (Intercept) 2.23 0.20
## extrav 0.41 0.03
## sexgirl 0.95 0.16
## extrav:sexgirl 0.06 0.03
##
## Error terms:
## Groups Name Std.Dev. Corr
## class (Intercept) 1.63
    extrav 0.18 -0.94
##
## Residual 0.74
## ---
## number of obs: 2000, groups: class, 100
\#\# AIC = 4890.7, DIC = 4836.6
## deviance = 4855.7
```

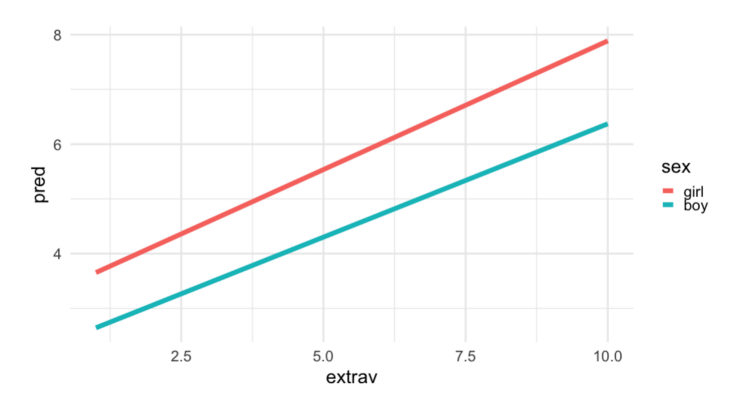
### Marginal effect

Let's look at the interaction between extraversion and sex

```
extrav sex class pred
##
         1 girl 0 3.654686
## 1
        2 girl 0 4.124926
3 girl 0 4.595165
## 2
## 3
## 4 4 girl 0 5.065404
## 5 5 girl 0 5.535643
## 6 6 girl 0 6.005883
## 7 7 girl 0 6.476122
## 8 8 girl 0 6.946361
## 9 9 girl 0 7.416600
    10 girl 0 7.886840
## 10
## 11
    1 boy 0 2.645470
     2 boy 0 3.059526
## 12
         3 boy 0 3.473583
## 13
```

### Plot

```
ggplot(conditions2, aes(extrav, pred)) +
  geom_line(aes(color = sex))
```



## Questions?

## Notation

Introducing the Gelman and Hill notation

### Standard regression

Imagine we have a model like this

We would probably display this model like this

$$mpg = \alpha + \beta_1(disp) + \beta_2(hp) + \beta_3(drat) + \epsilon$$

What we often don't show, is the distributional assumption of the residuals

$$\epsilon \sim N\left(0,\sigma
ight)$$

### A different view

The model on the previous slide could also be displayed like this

$$\hat{y} = lpha + eta_1( ext{disp}) + eta_2( ext{hp}) + eta_3( ext{drat}) \ ext{mpg} \sim N\left(\hat{y}, \sigma
ight)$$

This makes the distributional assumptions clearer

Each mpg value is assumed generated from a normal distribution, with a mean structure according to  $\hat{y}$ , and an unknown standard deviation,  $\sigma$ .

### Simulate

If we have a solid understanding of the distributional properties, we can simulate new data from the model First let's set some population parameters

```
n <- 1000
intercept <- 100
b1 <- 5
b2 <- -3
b3 <- 0.5
sigma <- 4.5
```

### Simulate

Next create some variables. The standard deviations relate to the standard errors – more variance in the predictor leads to lower standard errors.

```
set.seed(123)
x1 <- rnorm(n, sd = 1)
x2 <- rnorm(n, sd = 2)
x3 <- rnorm(n, sd = 4)</pre>
```

### Create y-hat

```
yhat <- intercept + b1*x1 + b2*x2 + b3*x3
```

### Generate data & test

```
sim <- rnorm(n, yhat, sigma)</pre>
summary(lm(sim \sim x1 + x2 + x3))
##
## Call:
\#\# lm(formula = sim \sim x1 + x2 + x3)
##
## Residuals:
## Min 1Q Median 3Q Max
## -13.7528 -2.8505 0.0021 3.0387 13.0151
##
## Coefficients:
##
    Estimate Std. Error t value Pr(>|t|)
## (Intercept) 99.96508   0.14141   706.92   <2e-16 ***
## x1 4.99415 0.14306 34.91 <2e-16 ***
## x2 -3.01827 0.07027 -42.95 <2e-16 ***
           ## x3
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4.466 on 996 degrees of freedom
## Multiple R-squared: 0.7514, Adjusted R-squared: 0.7506
## F-statistic: 1003 on 3 and 996 DF, p-value: < 2.2e-16
```

### Generalizing

We can generalize this same basic approach to multilevel models

This is helpful because the error structure is more complicated

Using this approach helps us better understand the distributional assumptions of our model

### Simple example

I know we hate the HSB data but bear with me for a minute.

Consider this simple model

```
library(lme4)
library(equatiomatic)
hsb_m0 <- lmer(math ~ ses + (1|sch.id), data = hsb)</pre>
```

In Raudenbush and Bryk notation, the model on the prior slide would look like this

$$egin{aligned} \mathrm{math}_{ij} &= eta_{0j} + eta_{1j}(\mathrm{ses}) + e_{ij} \ eta_{0j} &= \gamma_{00} + u_{0j} \ eta_{1j} &= \gamma_{10} \end{aligned}$$

Generally, the distributional Put differently part is omitted, which in this case is

$$egin{aligned} E\left(e_{ij}
ight) &= 0, \mathrm{Var}\left(e_{ij}
ight) = \sigma^2 \ E\left(u_{0j}
ight) &= 0, \mathrm{Var}\left(u_{0j}
ight) = au_{00} \end{aligned}$$

$$e_{ij} \sim N\left(0, \sigma^2
ight) \ u_{0j} \sim N\left(0, au_{00}
ight)$$

#### G&H

In Gelman & Hill notation, this same model can be communicated as

$$egin{aligned} & \operatorname{math}_i \sim N\left(lpha_{j[i]} + eta_1(\operatorname{ses}), \sigma^2
ight) \ & lpha_j \sim N\left(\mu_{lpha_j}, \sigma^2_{lpha_j}
ight), ext{for sch.id j} = 1, \dots, & \mathrm{J} \end{aligned}$$

This notation communicates the distributional assumptions

We can also still easily see what levels the predictors are at

It does look a little more complex, but it's not hiding anything

If you properly understand the notation, you can simultate data assuming this data generating process (which we'll do later)

#### Bonus

It works really well to communicate model results

$$egin{aligned} \widehat{\mathrm{math}}_i \sim N \left( 12.66_{lpha_{j[i]}} + 2.39_{eta_1}(\mathrm{ses}), 6.09 
ight) \ lpha_j \sim N \left( 0, 2.18 
ight), ext{ for sch.id j} = 1, \ldots, \! \mathrm{J} \end{aligned}$$

#### Extra bonus!

You can use equatiomatic to give you the model formula. The above was generated with extract\_eq(hsb\_m0,
use\_coef = TRUE)

### Quick simulation

We'll go over this in more detail later, but I want to give you the general idea.

First, set some parameters

```
j <- 30 # 30 schools
nj <- 50 # 50 students per school</pre>
```

Next, simulate the school distribution

```
# School distribution
a_j <- rnorm(j, 0, 2.18)</pre>
```

For each school, simulate nj obs from leve 1 model, adding in the school deviation

There are lots of ways to do this — I'm using a for() loop here in an effort to be transparent

#### Test it out

```
sim_m0 <- lmer(score ~ ses + (1|scid), data = sim_df)
summary(sim_m0)</pre>
```

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: score ~ ses + (1 | scid)
##
     Data: sim df
##
## REML criterion at convergence: 9704.9
##
## Scaled residuals:
## Min 1Q Median 3Q Max
## -3.1418 -0.6848 0.0030 0.6552 3.5886
##
## Random effects:
## Groups Name Variance Std.Dev.
## scid (Intercept) 5.685 2.384
## Residual 36.187 6.016
## Number of obs: 1500, groups: scid, 30
##
## Fixed effects:
## Estimate Std. Error t value
## (Intercept) 12.3901 0.4622 26.81
## ses 2.4682 0.1562 15.80
##
## Correlation of Fixed Effects:
## (Intr)
```

## Expanding the model

Let's add a school-level predictor

```
hsb_m1 <- lmer(math ~ ses + sector + (1|sch.id), data = hsb)
extract_eq(hsb_m1)</pre>
```

$$egin{aligned} & \operatorname{math}_i \sim N\left(lpha_{j[i]} + eta_1(\operatorname{ses}), \sigma^2
ight) \ & lpha_j \sim N\left(\gamma_0^lpha + \gamma_1^lpha(\operatorname{sector}), \sigma_{lpha_j}^2
ight), ext{for sch.id j} = 1, \ldots, & ext{J} \end{aligned}$$

## Add in a random slope

hsb\_m2 <- lmer(math ~ ses + sector + (ses|sch.id), data = hsb)
extract\_eq(hsb\_m2)</pre>

$$egin{aligned} & \operatorname{math}_i \sim N\left(lpha_{j[i]} + eta_{1j[i]}(\operatorname{ses}), \sigma^2
ight) \ & \left(egin{aligned} lpha_j \ eta_{1j} \end{aligned}
ight) \sim N\left(\left(egin{aligned} \gamma_0^lpha + \gamma_1^lpha(\operatorname{sector}) \ \mu_{eta_{1j}} \end{aligned}
ight), \left(egin{aligned} \sigma_{lpha_j}^2 & 
ho_{lpha_jeta_{1j}} \ 
ho_{eta_{1j}lpha_j} & \sigma_{eta_{1j}}^2 \end{array}
ight)
ight), ext{ for sch.id } ext{j} = 1, \ldots, ext{J} \end{aligned}$$

#### Include interaction

Include **sector** as a predictor of the relation between **ses** and **math** 

$$egin{aligned} & \operatorname{math}_i \sim N\left(lpha_{j[i]} + eta_{1j[i]}(\operatorname{ses}), \sigma^2
ight) \ & \left(egin{array}{c} lpha_j \ eta_{1j} \end{array}
ight) \sim N\left(\left(egin{array}{c} \gamma_0^lpha + \gamma_1^lpha(\operatorname{sector}) \ \gamma_0^{eta_1} + \gamma_1^{eta_1}(\operatorname{sector}) \end{array}
ight), \left(egin{array}{c} \sigma_{lpha_j}^2 & 
ho_{lpha_jeta_{1j}} \ 
ho_{eta_{1j}lpha_j} & \sigma_{eta_{1j}}^2 \end{array}
ight) 
ight), ext{ for sch.id } ext{j} = 1, \ldots, ext{J} \end{aligned}$$

## Even more complicated

This model doesn't actually fit well – I omitted some convergence warnings

```
hsb_m4 <- lmer(
  math ~ ses * sector + minority + female + meanses + size +
    (ses + minority + female|sch.id),
  data = hsb
)
extract_eq(hsb_m4)</pre>
```

$$\begin{aligned} & \operatorname{math}_{i} \sim N\left(\mu,\sigma^{2}\right) \\ & \mu = \alpha_{j[i]} + \beta_{1j[i]}(\operatorname{ses}) + \beta_{2j[i]}(\operatorname{minority}) + \beta_{3j[i]}(\operatorname{female}) \\ & \begin{pmatrix} \alpha_{j} \\ \beta_{1j} \\ \beta_{2j} \\ \beta_{3j} \end{pmatrix} \sim N \begin{pmatrix} \begin{pmatrix} \gamma_{0}^{\alpha} + \gamma_{1}^{\alpha}(\operatorname{sector}) + \gamma_{2}^{\alpha}(\operatorname{meanses}) + \gamma_{3}^{\alpha}(\operatorname{size}) \\ \gamma_{0}^{\beta_{1}} + \gamma_{1}^{\beta_{1}}(\operatorname{sector}) \\ \mu_{\beta_{2j}} \\ \mu_{\beta_{3j}} \end{pmatrix}, \begin{pmatrix} \sigma_{\alpha_{j}}^{2} & \rho_{\alpha_{j}\beta_{1j}} & \rho_{\alpha_{j}\beta_{2j}} & \rho_{\alpha_{j}\beta_{3j}} \\ \rho_{\beta_{1j}\alpha_{j}} & \sigma_{\beta_{1j}}^{2} & \rho_{\beta_{1j}\beta_{2j}} & \rho_{\beta_{1j}\beta_{3j}} \\ \rho_{\beta_{2j}\alpha_{j}} & \rho_{\beta_{2j}\beta_{1j}} & \sigma_{\beta_{2j}}^{2} & \rho_{\beta_{2j}\beta_{3j}} \\ \rho_{\beta_{3j}\alpha_{j}} & \rho_{\beta_{3j}\beta_{1j}} & \rho_{\beta_{3j}\beta_{2j}} & \sigma_{\beta_{3j}}^{2} \end{pmatrix}, \text{ for sch.id } \mathbf{j} = 1, \dots, \mathbf{J} \end{aligned}$$

## Multiple levels

Let's go to a different dataset from equatiomatic

#### head(sim\_longitudinal)

```
## # A tibble: 6 x 8
  # Groups: school [1]
      sid school district group treatment prop low wave score
##
##
  <int> <int> <int> <chr> <fct>
                                        <dbl> <dbl> <dbl>
                                  0.1428571
## 1
                                                0 102.2686
                   1 medium 1
## 2
             1 1 medium 1
                              0.1428571 1 102.0135
             1 1 medium 1 0.1428571 2 102.5216
1 medium 1 0.1428571 3 102.2792
## 3
## 4
## 5
                    1 medium 1 0.1428571
                                                4 102.2834
                    1 medium 1 0.1428571
## 6
                                                5 102.7963
```

#### Four levels

#### Model doesn't really fit again

```
sl_m <- lmer(
  score ~ wave*treatment + group + prop_low +
      (wave|sid) + (wave + treatment| school) + (1|district),
  data = sim_longitudinal
)
extract_eq(sl_m)</pre>
```

$$egin{aligned} & \operatorname{score}_i \sim N\left(lpha_{j[i],k[i],l[i]} + eta_{1j[i],k[i]}(\operatorname{wave}),\sigma^2
ight) \ & \left(egin{aligned} lpha_j \ eta_{1j} \end{aligned}
ight) \sim N\left(\left(egin{aligned} \gamma_0^lpha + \gamma_{1k[i]}^lpha(\operatorname{treatment}_1) + \gamma_2^lpha(\operatorname{group}_{\operatorname{low}}) + \gamma_3^lpha(\operatorname{group}_{\operatorname{medium}}) \ \end{pmatrix}, \left(egin{aligned} \sigma_{lpha_j}^2 & 
ho_{lpha_jeta_{1j}} \ 
ho_{eta_{1j}lpha_{1j}} \end{array}
ight) \end{aligned}
ight), ext{for sid } ext{j} = 1, \dots \ & \left(egin{aligned} lpha_k \ eta_{1k} \ eta_{1k} \ \end{pmatrix} \sim N\left(\left(egin{aligned} \gamma_0^lpha + \gamma_1^lpha(\operatorname{prop\_low}) \ \mu_{eta_{1k}} \ \mu_{\gamma_{1k}} \end{array}
ight), \left(egin{aligned} \sigma_{lpha_k}^2 & 
ho_{lpha_keta_{1k}} & 
ho_{lpha_k\gamma_{1k}} \ eta_{\beta_{1k}} & \sigma_{\beta_{1k}\gamma_{1k}} \ eta_{\gamma_{1k}} \ \end{pmatrix} 
ight), ext{for school } ext{k} = 1, \dots, ext{K} \ & lpha_l \sim N\left(\mu_{lpha_l}, \sigma_{lpha_l}^2\right), ext{for district } ext{l} = 1, \dots, ext{L} \end{aligned}$$

## Residual structures

#### Data

#### Willett, 1988

- n = 35 people
- Each completed a cognitive inventory on "opposites naming"
- At first time point, participants also completed a general cognitive measure

## Read in data

```
willett <- read_csv(here::here("data", "willett-1988.csv"))
willett</pre>
```

```
## # A tibble: 140 \times 4
##
        id time
                  opp
                        cog
##
     <dbl> <dbl> <dbl> <dbl> <dbl>
## 1
                  205
                      137
##
   2
                  217
                      137
##
   3
                  268
                      137
##
   4
                  302
                      137
##
   5
                  219 123
## 6
                  243 123
##
   7
                  279 123
##
                  302
                      123
##
   9
              0 142
                      129
## 10
                  212
                        129
## # ... with 130 more rows
```

#### Standard OLS

- We have four observations per participant.
- If we fit a standard OLS model, it would look like this

```
bad <- lm(opp ~ time, data = willett)
summary(bad)</pre>
```

```
##
## Call:
## lm(formula = opp ~ time, data = willett)
##
## Residuals:
##
     Min 1Q Median 3Q Max
## -88.374 -25.584 1.186 28.926 64.746
##
## Coefficients:
##
    Estimate Std. Error t value Pr(>|t|)
## (Intercept) 164.374 5.035 32.65 <2e-16 ***
       26.960 2.691 10.02 <2e-16 ***
## time
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 35.6 on 138 degrees of freedom
## Multiple R-squared: 0.421, Adjusted R-squared: 0.4168
```

## Assumptions

As we discussed previously, this model looks like this

$$opp = \alpha + \beta_1(time) + \epsilon$$

where

$$\epsilon \sim (0,\sigma)$$

#### Individual level residuals

We can expand our notation, so it looks like a multivariate normal distribution

$$egin{pmatrix} \epsilon_1 \ \epsilon_2 \ \epsilon_3 \ dots \ \epsilon_n \end{pmatrix} \sim egin{pmatrix} 0 \ 0 \ 0 \ 0 \ 0 \ 0 \ \sigma_\epsilon & 0 & 0 \ 0 \ 0 & \sigma_\epsilon & 0 & 0 \ 0 & \sigma_\epsilon & 0 & 0 \ 0 & 0 & \sigma_\epsilon & 0 & 0 \ 0 & 0 & \sigma_\epsilon & 0 & 0 \ \end{bmatrix}$$

This is where the i.i.d. part comes in. The residuals are assumed independent and identically distributed.

#### Multilevel model

Very regularly, there are reasons to believe the *i.i.d.* assumption is violated. Consider our current case, with 4 time points for each individual.

- Is an observation for one time point for one individual independent from the other observations for that individual?
- Rather than estimating a single residual variance, we estimate an additional components associated with individuals, leading to a *block* diagonal structure

## Block diagonal

$$\begin{pmatrix} \epsilon_{11} \\ \epsilon_{12} \\ \epsilon_{13} \\ \epsilon_{14} \\ \epsilon_{21} \\ \epsilon_{22} \\ \epsilon_{23} \\ \epsilon_{24} \\ \vdots \\ \epsilon_{n1} \\ \epsilon_{n2} \\ \epsilon_{n3} \\ \epsilon_{n4} \end{pmatrix} \sim \begin{pmatrix} \epsilon_{11} \\ \epsilon_{12} \\ \sigma_{0} \\ \sigma_{$$

Correlations for off-diagonals estimated

Same variance components for all blocks

Off diagonals are still zero

## Homogeneity of variance

As mentioned on the previous slide, we assume the same variance components across all student

This is referred to as the homogeneity of variance assumption – although the block (often referred to as the composite residual) may be heteroscedastic and dependent within a grouping factor (i.e., people) the entire error structure is repeated identically across units (i.e., people)

## Block diagonal

Because of the homogeneity of variance assumption, we can re-express our block diagonal design as follows

$$r\sim N egin{bmatrix} oldsymbol{\Sigma_r} & oldsymbol{0} & oldsymbol{\Omega} & oldsymbol{\Sigma_r} & oldsymbol{0} & \dots & oldsymbol{0} \ oldsymbol{0} & oldsymbol{\Omega} & oldsymbol{\Sigma_r} & \dots & oldsymbol{0} \ dots & dots & dots & \ddots & dots \ oldsymbol{0} & oldsymbol{0} & oldsymbol{0} & \dots & oldsymbol{\Sigma_r} \ \end{pmatrix}$$

## Composite residual

We then define the composite residual, which is common across units

$$m{\Sigma_r} = egin{bmatrix} \sigma_{11} & \sigma_{12} & \sigma_{13} & \sigma_{14} \ \sigma_{21} & \sigma_{22} & \sigma_{23} & \sigma_{24} \ \sigma_{31} & \sigma_{32} & \sigma_{33} & \sigma_{34} \ \sigma_{41} & \sigma_{42} & \sigma_{43} & \sigma_{44} \end{bmatrix}$$

## Let's try!

Let's fit a parallel slopes model with the Willett data. You try first.

```
w0 <- lmer(opp ~ time + (1|id), willett)
```

What does the residual variance-covariance look like? Let's use **sundry** to pull it

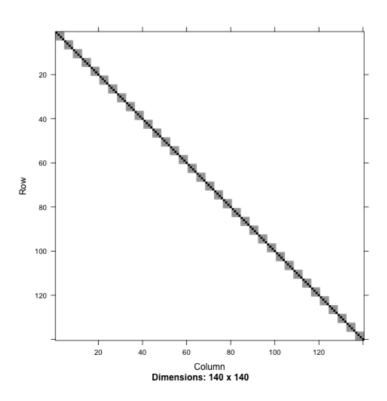
```
library(sundry)
w0_rvcv <- pull_residual_vcov(w0)</pre>
```



## lmage

Sparse matrix - we can view it with image()

image(w0\_rvcv)



## Pull first few rows/cols

```
w0_rvcv[1:8, 1:8]
```

#### Structure

On the previous slide, note the values on the diagonal are all the same, as are all the off-diagonals

 This is because we've only estimated one additional variance component

## Understanding these numbers

Let's look at the model output

#### arm::display(w0)

The diagonal values were 1280.7065389 while the off diagonal values were 904.8053852

Let's extract the variance components from our model.

```
vars_w0 <- as.data.frame(VarCorr(w0))
vars_w0</pre>
```

```
## grp var1 var2 vcov sdcor
## 1 id (Intercept) <NA> 904.8054 30.07998
## 2 Residual <NA> <NA> 375.9012 19.38817
```

Notice anything?

The diagonals are given by **sum(vars\_w0)\$vcov** while the off-diagonals are just the intercept variance

## Including more complexity

Try estimating this model now, then look at the residual variance—covariance matrix again

$$egin{aligned} \operatorname{opp}_i &\sim N\left(lpha_{j[i]} + eta_{1j[i]}( ext{time}), \sigma^2
ight) \ \left(egin{aligned} lpha_j \ eta_{1j} \end{aligned}
ight) &\sim N\left(\left(egin{aligned} \mu_{lpha_j} \ \mu_{eta_{1j}} \end{aligned}
ight), \left(egin{aligned} \sigma_{lpha_j}^2 & 
ho_{lpha_jeta_{1j}} \ 
ho_{eta_{1j}lpha_j} & \sigma_{eta_{1j}}^2 \end{array}
ight)
ight), ext{ for id } ext{j} = 1, \ldots, ext{J} \end{aligned}$$



## The composite residual

```
w1 <- lmer(opp ~ time + (time|id), willett)
w1_rvcv <- pull_residual_vcov(w1)
w1_rvcv[1:4, 1:4]</pre>
```

```
## 4 x 4 sparse Matrix of class "dgCMatrix"

## 1 2 3 4

## 1 1358.2469 1019.5095 840.2515 660.9934

## 2 1019.5095 1132.1294 925.7905 878.9310

## 3 840.2515 925.7905 1170.8089 1096.8686

## 4 660.9934 878.9310 1096.8686 1474.2856
```

#### Unstructured

The model we fit has an *unstructured* variance co-variance matrix. While each block is the same, every element of the block is now estimated.

### What are these numbers?

They are the variance components, re-expressed as a composite residual

The diagonal is given by

$$\sigma^2+\sigma_{lpha_j}^2+2\sigma_{01}^2w_i+\sigma_{eta_1}^2w_i^2$$

where  $oldsymbol{w}$  represents the given wave (for our example)

Let's do this "by hand"

## Get the pieces

```
vars_w1 <- as.data.frame(VarCorr(w1))

# get the pieces
int_var <- vars_w1$vcov[1]
slope_var <- vars_w1$vcov[2]
covar <- vars_w1$vcov[3]
residual <- vars_w1$vcov[4]</pre>
```

### Calculate

```
diag(w1_rvcv[1:4, 1:4])
## [1] 1358.247 1132.129 1170.809 1474.286
residual + int_var
## [1] 1358.247
residual + int_var + 2*covar + slope_var
## [1] 1132.129
residual + int_var + (2*covar)*2 + slope_var*2^2
## [1] 1170.809
residual + int_var + (2*covar)*3 + slope_var*3^2
## [1] 1474.286
```

## Off-diagonals

The off-diagonals are given by

$$\sigma_{lpha_j}^2 + \sigma_{01}(t_i + t_i') + \sigma_{eta_1}^2 t_i t_i'$$

#### Calculate a few

```
w1 rvcv[1:4, 1:4]
## 4 x 4 sparse Matrix of class "dqCMatrix"
##
## 1 1358.2469 1019.5095 840.2515 660.9934
## 2 1019.5095 1132.1294 925.7905 878.9310
## 3 840.2515 925.7905 1170.8089 1096.8686
## 4 660.9934 878.9310 1096.8686 1474.2856
int_var + covar*(1 + 0) + slope_var*1*0
## [1] 1019.51
int var + covar*(2 + 1) + slope var*2*1
## [1] 925.7905
int_var + covar*(3 + 2) + slope_var*3*2
## [1] 1096.869
int_var + covar*(2 + 0) + slope_var*2*0
```

# Positing other structures

# The possibilities

There are a number of alternative structures. We'll talk about a few here.

If you want to go deeper, I suggest Singer & Willett, Chapter 7

Code to fit models with each type of structure, using the same Willett data we're using today, is available here

## Structures we'll fit

- Unstructured (default with Ime4, we've already seen this)
- Autoregressive
- Heterogeneous autoregressive
- Toeplitz

Outside of unstructured, we'll need to use the **nlme** package to estimate other structures

We'll also use a generalized least squares estimator, rather than maximum likelihood

# Autoregressive

# Autoregressive

- There are many types of autoregressive structures
  - If you took a class on time—series data you'd learn about others
- What we'll talk about is referred to as an AR1 structure
- Variances (on the diagonal) are constant
- Includes constant "band-diagonals"

# Autoregressive

$$oldsymbol{\Sigma_r} = egin{bmatrix} \sigma^2 & \sigma^2 
ho & \sigma^2 
ho^2 & \sigma^2 
ho^3 \ \sigma^2 
ho & \sigma^2 & \sigma^2 
ho & \sigma^2 
ho^2 \ \sigma^2 
ho^2 & \sigma^2 
ho & \sigma^2 & \sigma^2 
ho \ \sigma^2 
ho^3 & \sigma^2 
ho^2 & \sigma^2 
ho & \sigma^2 \end{bmatrix}$$

- Each band is forced to be lower than the prior by a constant fraction
  - $\circ$  estimated autocorrelation parameter ho. The error variance is multiplied by ho for the first diagonal, by  $ho^2$  for the second, etc.
- Uses only two variance components

## Fit

#### First load nlme

```
library(nlme)
```

We'll use the gls() function. The interface is, overall, fairly similar to **Ime4** 

# Summary

#### summary(ar)

```
## Generalized least squares fit by REML
## Model: opp ~ time
## Data: willett
##
        AIC BIC logLik
## 1281.465 1293.174 -636.7327
##
## Correlation Structure: AR(1)
## Formula: ~1 | id
## Parameter estimate(s):
##
       Phi
## 0.8249118
##
## Coefficients:
##
                Value Std.Error t-value p-value
## (Intercept) 164.33842 6.136372 26.78104
## time 27.19786 1.919857 14.16661
##
## Correlation:
## (Intr)
## time -0.469
##
## Standardized residuals:
##
         Min
                     O1 Med
                                           Q3
                                                    Max
## -2.42825488 -0.71561388 0.03192973 0.78792605 1.76110779
```

## Extract composite residual

```
cm_ar <- corMatrix(ar$modelStruct$corStruct) # all of them
cr_ar <- cm_ar[[1]] # just the first (they're all the same)
cr_ar</pre>
```

```
## [,1] [,2] [,3] [,4]
## [1,] 1.0000000 0.8249118 0.6804795 0.5613356
## [2,] 0.8249118 1.0000000 0.8249118 0.6804795
## [3,] 0.6804795 0.8249118 1.0000000 0.8249118
## [4,] 0.5613356 0.6804795 0.8249118 1.0000000
```

Multiply the correlation matrix by the model residual variance to get the covariance matrix

```
cr_ar * sigma(ar)^2
```

```
## [,1] [,2] [,3] [,4]

## [1,] 1323.4596 1091.7375 900.5872 742.9050

## [2,] 1091.7375 1323.4596 1091.7375 900.5872

## [3,] 900.5872 1091.7375 1323.4596 1091.7375

## [4,] 742.9050 900.5872 1091.7375 1323.4596
```

# Confirming calculations

```
sigma(ar)^2
## [1] 1323.46
sigma(ar) ^2*0.8249118
## [1] 1091.737
sigma(ar)^2*0.8249118^2
## [1] 900.5871
sigma(ar)^2*0.8249118^3
## [1] 742.9049
```

# Heterogenous autoregressive

- Same as autorgressive but allows each variance to differ
- Still one ho estimated
  - Same "decay" across band diagonals
- Band diagonals no longer equivalent, because different variances

# Heterogenous autoregressive

$$oldsymbol{\Sigma_r} = egin{bmatrix} \sigma_1^2 & \sigma_1\sigma_2
ho & \sigma_1\sigma_3
ho^2 & \sigma_1\sigma_4
ho^2 \ \sigma_2\sigma_1
ho & \sigma_2^2 & \sigma_2\sigma_3
ho & \sigma_2\sigma_4
ho^2 \ \sigma_3\sigma_1
ho^2 & \sigma_3\sigma_2
ho & \sigma_3^2 & \sigma_3\sigma_4
ho \ \sigma_4\sigma_1
ho^3 & \sigma_4\sigma_2
ho^2 & \sigma_4\sigma_3
ho & \sigma_4^2 \end{bmatrix}$$

### Fit

Note - varIdent specifies different variances for each wave (variances of the identity matrix)

```
har <- gls(
  opp ~ time,
  data = willett,
  correlation = corAR1(form = ~ 1|id),
  weights = varIdent(form = ~1|time)
)</pre>
```

# Summary

#### summary(har)

```
## Generalized least squares fit by REML
## Model: opp ~ time
## Data: willett
##
        AIC BIC logLik
## 1285.76 1306.25 -635.8798
##
## Correlation Structure: AR(1)
## Formula: ~1 | id
## Parameter estimate(s):
##
       Phi
## 0.8173622
## Variance function:
## Structure: Different standard deviations per stratum
## Formula: ~1 | time
## Parameter estimates:
## 1.000000 0.915959 0.985068 1.045260
##
## Coefficients:
##
                 Value Std.Error t-value p-value
## (Intercept) 164.63344 5.959533 27.62523
## time 27.11552 1.984807 13.66154
##
## Correlation:
```

# Extract/compute composite residual

The below is fairly complicated, but you can work it out if you go line by line

```
cm_har <- corMatrix(har$modelStruct$corStruct)[[1]]
var_struct <- har$modelStruct$varStruct
vars <- coef(var_struct, unconstrained = FALSE, allCoef = TRUE)
vars <- matrix(vars, ncol = 1)

cm_har * sigma(har)^2 *
  (vars %*% t(vars)) # multiply by a mat of vars</pre>
```

```
## [,1] [,2] [,3] [,4]

## [1,] 1308.6660 979.7593 861.2399 746.9588

## [2,] 979.7593 1097.9457 965.1295 837.0629

## [3,] 861.2399 965.1295 1269.8757 1101.3712

## [4,] 746.9588 837.0629 1101.3712 1429.8058
```

# Toeplitz

- Constant variance
- Has identical band-diagonals, like autoregressive
- Relaxes assumption of each band being a parallel by a common fraction of the prior band
  - Each band determined empirically by the data

A bit of a compromise between prior two

## Toeplitz

$$m{\Sigma_r} = egin{bmatrix} \sigma^2 & \sigma^2_1 & \sigma^2_2 & \sigma^2_3 \ \sigma^2_1 & \sigma^2 & \sigma^2_1 & \sigma^2_2 \ \sigma^2_2 & \sigma^2_1 & \sigma^2 & \sigma^2_1 \ \sigma^2_2 & \sigma^2_2 & \sigma^2_1 & \sigma^2 \end{bmatrix}$$

#### Fit

# Summary

#### summary(toep)

```
## Generalized least squares fit by REML
## Model: opp ~ time
## Data: willett
##
        AIC BIC logLik
## 1277.979 1295.543 -632.9896
##
## Correlation Structure: ARMA(3,0)
## Formula: ~1 | id
## Parameter estimate(s):
##
       Phi1 Phi2 Phi3
##
   0.8039121 0.3665122 -0.3950326
##
## Coefficients:
##
                Value Std.Error t-value p-value
## (Intercept) 165.11855 6.122841 26.96764
## time 26.91997 2.070391 13.00236
##
## Correlation:
## (Intr)
## time -0.507
##
## Standardized residuals:
##
         Min
                     O1 Med
                                          03
                                                   Max
## -2.44029024 -0.71984566 0.01373249 0.77304950 1.75580973
```

# Extract/compute composite residual

Same as with autoregressive – just multiply the correlation matrix by the residual variance

```
cr_toep <- corMatrix(toep$modelStruct$corStruct)[[1]]
cr_toep * sigma(toep)^2</pre>
```

```
## [,1] [,2] [,3] [,4]

## [1,] 1333.6848 1105.7350 940.9241 634.8366

## [2,] 1105.7350 1333.6848 1105.7350 940.9241

## [3,] 940.9241 1105.7350 1333.6848 1105.7350

## [4,] 634.8366 940.9241 1105.7350 1333.6848
```

# Comparing fits

```
library(performance)
compare_performance(ar, har, toep, w1,
                  metrics = c("AIC", "BIC"),
                  rank = TRUE) %>%
  as_tibble()
## Warning: Could not get model data.
## Warning: Could not get model data.
## Warning: Could not get model data.
## Warning: When comparing models, please note that probably not all models
## data.
## # A tibble: 4 x 5
## Name Model
                  AIC BIC Performance Score
## <chr> <dbl> <dbl>
                                            <dbl>
## 1 toep gls 1277.979 1295.543
                                      0.9094409
## 2 w1 lmerMod 1278.823 1296.473 0.8196721
## 3 ar qls 1281.465 1293.174 0.7759608
## 4 har gls 1285.760 1306.250
```

	Unstructured	Autoregressive (AR)	Heterogeneous AR	Toeplitz
(Intercept)	164.374	164.338	164.633	165.119
	(6.119)	(6.136)	(5.960)	(6.123)
time	26.960	27.198	27.116	26.920
	(2.167)	(1.920)	(1.985)	(2.070)
sd(Intercept)	34.623			
cor_(Intercept).time	-0.450			
sdtime	11.506			
sd_Observation	12.629			
AIC	1278.8	1281.5	1285.8	1278.0
BIC	1296.5	1293.2	1306.3	1295.5
Log.Lik.	-633.411	-636.733	-635.880	-632.990
REMLcrit	1266.823			

# Stepping

Why do we care about all of this?

# Overfitting

- If a simpler model fits the data just as well, it should be preferred
- Models that are overfit have "learned too much" from the data and won't generalize well
- Can think of it as fitting to the errors in the data, rather than the "true" patterns found in the population

# Convergence issues

- As your models get increasingly complicated, you're likely to run into convergence issues.
- Simplifying your residual variance—covariance structure may help
  - Keeps the basic model structure intact

Aside – see here for convergence troubleshooting with **Ime4**. The allFit() function is often helpful but very computationally intensive.

# Summary

- As is probably fairly evident from the code there are many more structures you could explore. However, most are not implemented through **Ime4**.
- Simplifying the residual variance—covariance can sometimes lead to better fitting models
- May also be helpful with convergence and avoid overfitting

# Next time: Modeling growth (part