

# Crossed Random Effects for Other Repeated Measures Designs

- Topics:
  - **ANOVA for repeated measures**
  - MLM for repeated measures

# Analytic Toolbox of the Experimental Psychologist

- Our friend, analysis of variance (ANOVA)
  - Between-group (*aka* between-subject, independent IV)
  - Within-group (*aka* within-subject, dependent, repeated measures IV)
  - Split-plot (*aka* mixed design of between- and within-group IVs)
- Expandable to include:
  - multiple IVs (factorial ANOVA)
  - main effects of continuous covariates (ANCOVA)
  - multiple outcomes (MANOVA/MANCOVA)

# ANOVA works well when...

- Experimental stimuli are **controlled** and **exchangeable**
  - Controlled → Constructed, not sampled from a population
  - Exchangeable → Stimuli vary only in dimensions of interest
  - ...What to do with non-exchangeable stimuli (e.g., words, scenes)?
- Experimental manipulations create **discrete conditions**
  - e.g., set size of 3 vs. 6 vs. 9 items
  - e.g., response compatible vs. incompatible distractors
  - ...What to do with *continuous* item predictors (e.g., time, salience)?
- One has **complete data**
  - e.g., if outcome is RT and accuracy is near ceiling
  - e.g., if responses are missing for no systematic reason
  - ...What if data are not missing completely at random (e.g., inaccuracy)?

# Motivating Example: Psycholinguistic Study Designs

- Word Recognition Tasks (e.g., Lexical Decision)
  - Word lists are constructed based on targeted dimensions while controlling for other relevant dimensions
  - Outcome = RT to decide if the stimulus is a word or non-word (accuracy is usually near ceiling)
- Tests of effects of experimental treatment are typically conducted with the person as the unit of analysis...
  - Average the responses over words within conditions
    - Contentious fights with reviewers about adequacy of experimental control when using real words as stimuli
    - Long history of debate as to how words as experimental stimuli should be analyzed...  $F_1$  ANOVA or  $F_2$  ANOVA (or both)?
    - $F_1$  only creates a “Language-as-Fixed-Effects Fallacy” (Clark, 1973)

# ANOVAs on Summary Data

## Original Data per Subject

	B1	B2
A1	Trial 001	Trial 101
	Trial 002	Trial 102
	.....	.....
	Trial 100	Trial 200
A2	Trial 201	Trial 301
	Trial 202	Trial 302
	.....	.....
	Trial 300	Trial 400



## Subject Summary Data

	B1	B2
A1	Mean (A1, B1)	Mean (A1, B2)
A2	Mean (A2, B1)	Mean (A2, B2)

## "F<sub>1</sub>" Repeated Measures ANOVA on *N* subjects:

$$RT_{cs} = \gamma_0 + \gamma_1 A_c + \gamma_2 B_c + \gamma_3 A_c B_c + \mathbf{U}_{0s} + e_{cs}$$

## "F<sub>2</sub>" Between-Groups ANOVA on *T* trials:

$$RT_t = \gamma_0 + \gamma_1 A_t + \gamma_2 B_t + \gamma_3 A_t B_t + e_t$$

## Trial Summary Data

	B1
A1, B1	Trial 001 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) Trial 002 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) ..... Trial 100
A1, B2	Trial 101 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) Trial 102 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) ..... Trial 200
A2, B1	Trial 201 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) Trial 202 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) ..... Trial 300
A2, B2	Trial 301 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) Trial 302 = Mean(Subject 1, Subject 2,... Subject <i>N</i> ) ..... Trial 400

# Choosing Amongst ANOVA Models

- $F_1$  RM ANOVA on **subject** summary data:
  - Assumes trials are fixed—within-condition **trial** variability is gone
- $F_2$  ANOVA on **trial** summary data:
  - Assumes persons are fixed—within-trial **subject** variability is gone
- Proposed ANOVA-based resolutions:
  - **F'** → quasi-F test that treats both trials and subjects as random (Clark, 1973), but requires complete data (least squares)
  - **Min F'** → lower-bound of F' derived from  $F_1$  and  $F_2$  results, which does not require complete data, but is (too) conservative
  - **$F_1 \times F_2$  criterion** → effects are only “real” if they are significant in **both  $F_1$  and  $F_2$  models** (aka, death knell for psycholinguists)
  - But neither model is complete (two wrongs don't make a right)...

# Sources of Variance (Clark, 1973)

$t = \#conditions, i = \#items, s = \#subjects$

Label		DF	Expected Mean Square
T	Treatments (t)	$t-1$	$\sigma_e^2 + \sigma_{S \times I}^2 + i\sigma_{T \times S}^2 + \text{---} + s\sigma_I^2 + i\sigma_T^2$
I w T	Items (i) within Treatments	$t(i-1)$	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + \text{---} + s\sigma_I^2 + \text{---}$
S	Subjects (s)	$s-1$	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + t\sigma_S^2 + \text{---} + \text{---}$
T x S	Treatments by Subjects	$(t-1)(s-1)$	$\sigma_e^2 + \sigma_{S \times I}^2 + i\sigma_{T \times S}^2 + \text{---} + \text{---} + \text{---}$
S x I w T	Subjects by Items within Treatments	$t(i-1)(s-1)$	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + \text{---} + \text{---} + \text{---}$

# Effect of Treatment via $F_1$ ANOVA

*T numerator should differ from TxS denominator by 1 term*

Label		DF	Expected Mean Square
T	Treatments (t)	t-1	$\sigma_e^2 + \sigma_{S \times I}^2 + i\sigma_{T \times S}^2 + \text{---} + \boxed{s\sigma_I^2} + \boxed{is\sigma_T^2}$
I w T	Items (i) within Treatments	t(i-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + \text{---} + s\sigma_I^2 + \text{---}$
S	Subjects (s)	s-1	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + t\sigma_S^2 + \text{---} + \text{---}$
T x S	Treatments by Subjects	(t-1)(s-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + i\sigma_{T \times S}^2 + \text{---} + \text{---} + \text{---}$
S x I w T	Subjects by Items within Treatments	t(i-1)(s-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + \text{---} + \text{---} + \text{---} + \text{---}$



# Effect of Treatment via $F_2$ ANOVA

*T numerator should differ from I x T denominator by 1 term*

Label		DF	Expected Mean Square
T	Treatments (t)	t-1	$\sigma_e^2 + \sigma_{S \times I}^2 + \boxed{i\sigma_{T \times S}^2} + \_\_\_\_\_\_ + s\sigma_I^2 + \boxed{is\sigma_T^2}$
I w T	Items (i) within Treatments	t(i-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + \_\_\_\_\_\_ + \_\_\_\_\_\_ + s\sigma_I^2 + \_\_\_\_\_\_$
S	Subjects (s)	s-1	$\sigma_e^2 + \sigma_{S \times I}^2 + \_\_\_\_\_\_ + t\sigma_S^2 + \_\_\_\_\_\_ + \_\_\_\_\_\_$
T x S	Treatments by Subjects	(t-1)(s-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + i\sigma_{T \times S}^2 + \_\_\_\_\_\_ + \_\_\_\_\_\_ + \_\_\_\_\_\_$
S x I w T	Subjects by Items within Treatments	t(i-1)(s-1)	$\sigma_e^2 + \sigma_{S \times I}^2 + \_\_\_\_\_\_ + \_\_\_\_\_\_ + \_\_\_\_\_\_ + \_\_\_\_\_\_$

# Simultaneous Quasi-F Ratio (F')

- F' was proposed by Clark (1973) as a quasi-F test that treats both items and subjects as random factors

$$F'(df_{\text{num}}, df_{\text{den}}) = \frac{MS_T + MS_{S \times I}}{MS_{T \times S} + MS_I}$$

$$\text{where } df_{\text{num}} = \frac{(MS_T + MS_{S \times I})^2}{\frac{MS_T}{df_T} + \frac{MS_{S \times I}}{df_{S \times I}}} \text{ and } df_{\text{den}} = \frac{(MS_{T \times S} + MS_I)^2}{\frac{MS_{T \times S}}{df_{T \times S}} + \frac{MS_I}{df_I}}$$

$$F'(df_{\text{num}}, df_{\text{den}}) = \frac{(2 * \sigma_e^2) + (2 * \sigma_{S \times I}^2) + (\#I * \sigma_{T \times S}^2) + (\#S * \sigma_I^2) + (\#I * \#S * \sigma_T^2)}{(2 * \sigma_e^2) + (2 * \sigma_{S \times I}^2) + (\#I * \sigma_{T \times S}^2) + (\#S * \sigma_I^2)}$$

- Numerator then exceeds the denominator by exactly the treatment variance as desired... except it requires complete data given that it relies on least squares
  - Not feasible in most real-world experiments

# Minimum of Quasi-F Ratio (Min F')

- Min F' was developed to be used from  $F_1$  and  $F_2$  results:

$$\min F'(\text{df}_{\text{num}}, \text{df}_{\text{den}}) = \frac{MS_T}{MS_{T \times S} + MS_I} = \frac{F_1 * F_2}{F_1 + F_2}$$

- But given that Min F' is overly conservative, having to show significance by both models is often required instead:
  - the  $F_1$  by  $F_2$  criterion... but two wrongs don't make a right
- Wouldn't it be nice if we had some way to treat subjects and items as the random effects they actually are???
  - And to assess the extent to which items are actually exchangeable?
  - And that all the extraneous item variables were adequately controlled?
  - **Multilevel models to the rescue! ... maybe?**

# Crossed Random Effects for Other Repeated Measures Designs

- Topics:
  - ANOVA for repeated measures
  - **MLM for repeated measures**

# Multilevel Models to the Rescue?

**Original Data per Person**

	B1	B2
A1	Trial 001 Trial 002 ..... Trial 100	Trial 101 Trial102 ..... Trial 200
A2	Trial 201 Trial 202 ..... Trial 300	Trial 301 Trial302 ..... Trial 400

## Pros:

- Use all original data, not summaries
- Responses can be missing at random
- Can include continuous trial predictors

## Cons:

- **Is still wrong**

$$\text{Level 1: } y_{ts} = \beta_{0s} + \beta_{1s}A_{ts} + \beta_{2s}B_{ts} + \beta_{3s}A_{ts}B_{ts} + e_{ts}$$

$$\text{Level 2: } \beta_{0s} = \gamma_{00} + U_{0s}$$

$$\beta_{1s} = \gamma_{10}$$

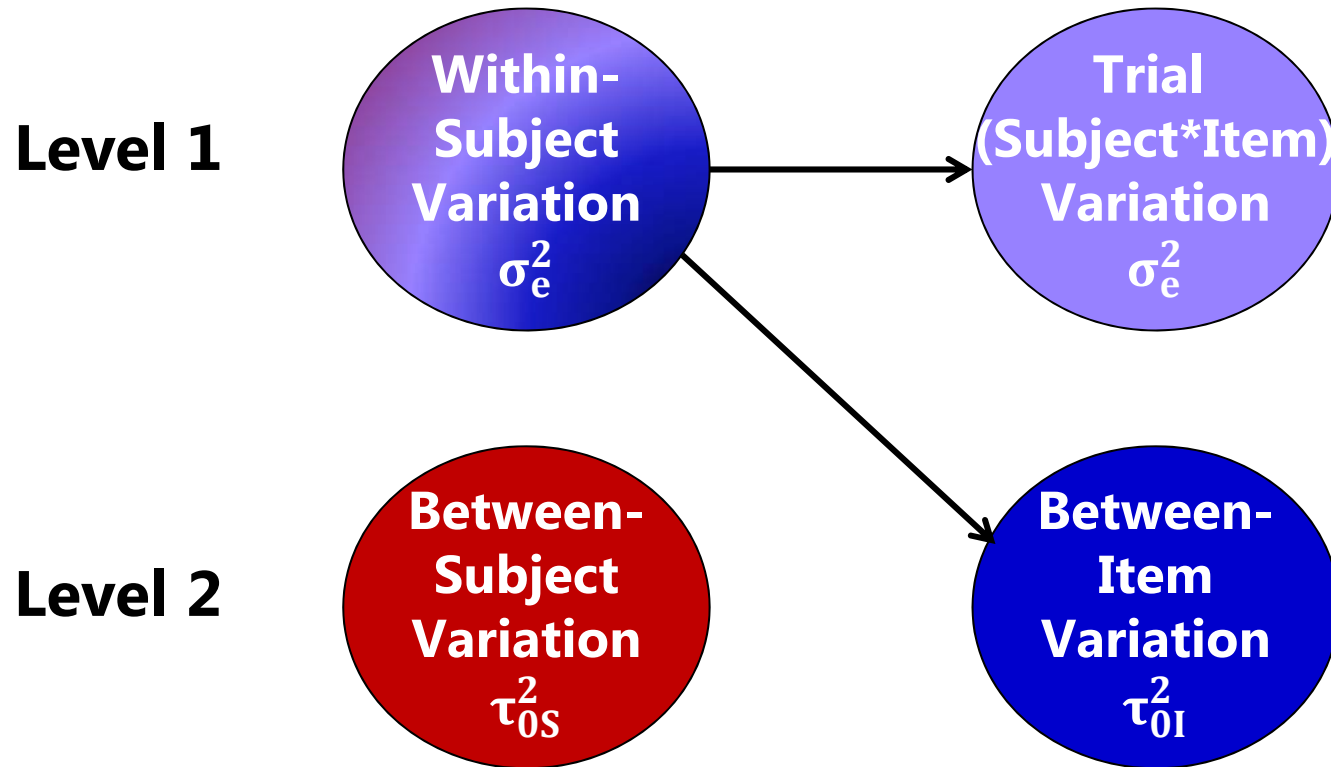
$$\beta_{2s} = \gamma_{20}$$

$$\beta_{3s} = \gamma_{30}$$

Level 1 = Within-Subject Variation  
(Across Trials)

Level 2 = Between-Subject Variation

# Multilevel Models to the Rescue?



# Empty Means, Crossed Random Effects Models

- **Residual-only model:**

- $RT_{tis} = \gamma_{000} + e_{tis}$
- Assumes no effects (dependency) of subjects or items

- **Random subjects model:**

- $RT_{tis} = \gamma_{000} + \mathbf{U}_{00s} + e_{tis}$
- Models systematic mean differences **between subjects**

- **Random subjects and items model:**

- $RT_{tis} = \gamma_{000} + U_{00s} + \mathbf{U}_{0io} + e_{tis}$
- Also models systematic mean differences **between items**

# A Better Way of (Multilevel) Life

Between-Subject Variation  
L2  $\tau_{0s}^2$

Between-Item Variation  
L2  $\tau_{0i}^2$

Trial (Subject\*Item) Variation  
 $\sigma_e^2$

Random effects over **subjects** of **item** or **trial** predictors can also be tested and predicted.

- **Multilevel Model with *Crossed* Random Effects:**

$$RT_{tis} = \gamma_{000} + \gamma_{010}A_i + \gamma_{020}B_i + \gamma_{030}A_iB_i + \mathbf{U}_{00s} + \mathbf{U}_{0i0} + \mathbf{e}_{tis}$$

$t$  trial  
 $i$  item  
 $s$  subject

- Both **subjects** and **items** as random effects:

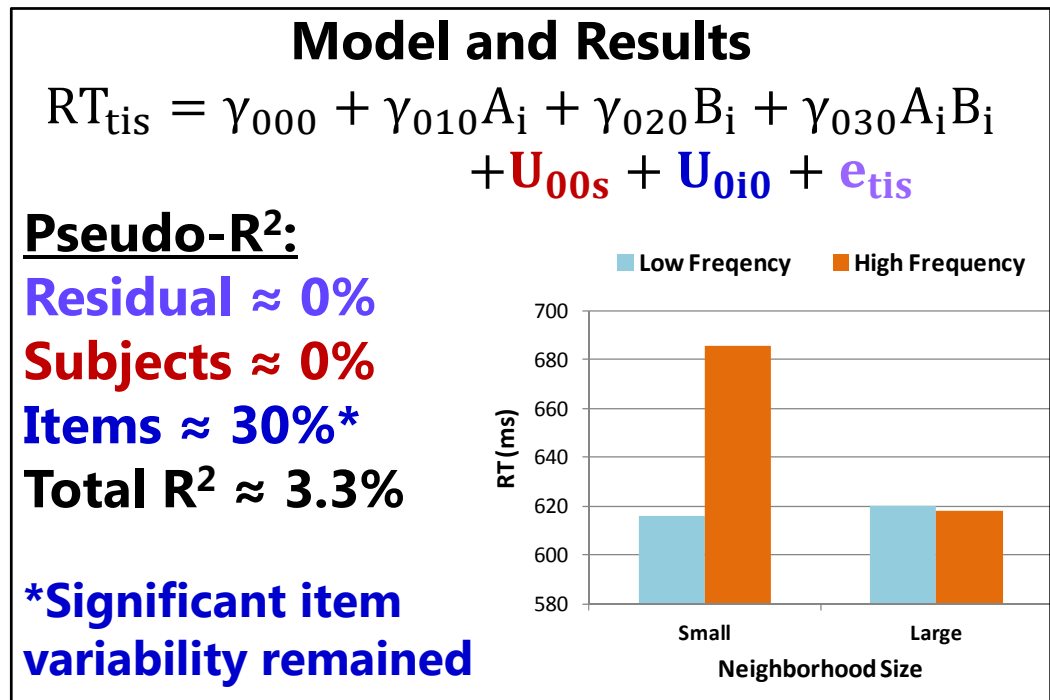
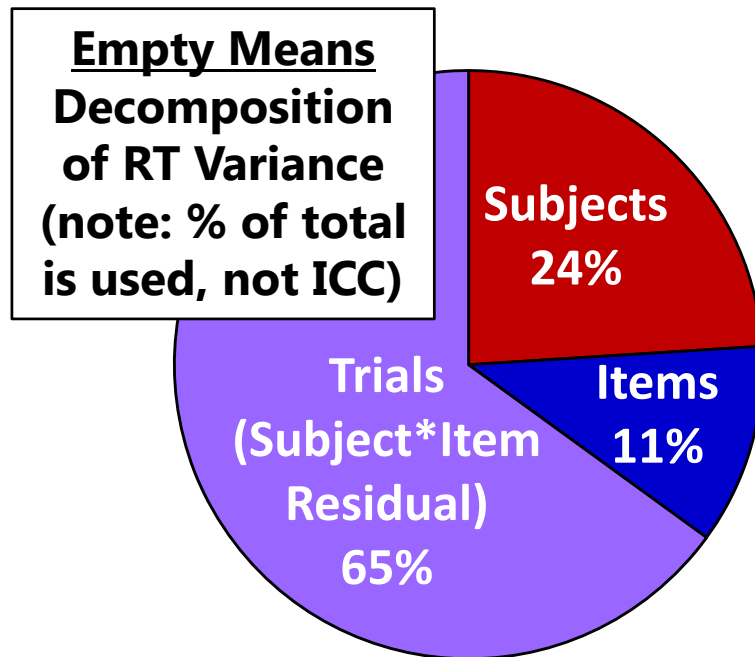
- Subject predictors explain between-subject mean variation:  $\tau_{0s}^2$
- Item predictors explain between-item mean variation:  $\tau_{0i}^2$
- Trial predictors explain trial-specific residual variation:  $\sigma_e^2$



# Example Psycholinguistic Study

(Locker, Hoffman, & Bovaird, 2007)

- Crossed design: 38 subjects by 39 items (words or nonwords)
- Lexical decision task: RT to decide if word or nonword
- 2 word-specific predictors of interest:
  - A: Low/High Phonological Neighborhood Frequency
  - B: Small/Large Semantic Neighborhood Size



# Tests of Fixed Effects by Model

	A: Frequency Marginal Main Effect	B: Size Marginal Main Effect	A*B: Interaction of Frequency by Size
<b>F<sub>1</sub> Subjects ANOVA</b>	$F(1,37) = 16.1$ $p = .0003$	$F(1,37) = 14.9$ $p = .0004$	$F(1,37) = 38.2$ $p < .0001$
<b>F<sub>2</sub> Words ANOVA</b>	$F(1,35) = 5.3$ $p = .0278$	$F(1,35) = 4.5$ $p = .0415$	$F(1,35) = 5.7$ $p = .0225$
<b>F' min (via ANOVA)</b>	$F(1,56) = 4.0$ $p = .0530$	$F(1,55) = 3.5$ $p = .0710$	$F(1,45) = 5.0$ $p = .0310$
<b>Crossed MLM (via REML)</b>	$F(1,32) = 5.4$ $p = .0272$	$F(1,32) = 4.6$ $p = .0393$	$F(1,32) = 6.0$ $p = .0199$

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# Simulation: Type 1 Error Rates

Condition		Models					
Item Variance	Subject Variance	1: Both Random Effects	2: Random Subjects Only	3: Random Items Only	4: No Random Effects	5: F1 Subjects ANOVA	6: F2 Item ANOVA
<b>Item Effect:</b>							
2	2	0.03	0.09	0.03	0.09	0.09	0.03
2	10	0.05	0.14	0.05	0.12	0.15	0.05
10	2	0.04	0.32	0.04	0.31	0.32	0.04
10	10	0.05	0.31	0.05	0.29	0.33	0.05
<b>Subject Effect:</b>							
2	2	0.04	0.04	0.12	0.11	0.04	0.12
2	10	0.05	0.05	0.34	0.34	0.05	0.36
10	2	0.04	0.03	0.12	0.09	0.03	0.12
10	10	0.06	0.06	0.34	0.31	0.05	0.37

# Model Items as Fixed → Wrong Item Effect

Condition		Models					
Item Variance	Subject Variance	1: Both Random Effects	2: <b>Random Subjects Only</b>	3: Random Items Only	4: No Random Effects	5: <b>F1 Subjects ANOVA</b>	6: F2 Item ANOVA
<b>Item Effect:</b>							
2	2	0.03	<b>0.09</b>	0.03	0.09	<b>0.09</b>	0.03
2	10	0.05	<b>0.14</b>	0.05	0.12	<b>0.15</b>	0.05
10	2	0.04	<b>0.32</b>	0.04	0.31	<b>0.32</b>	0.04
10	10	0.05	<b>0.31</b>	0.05	0.29	<b>0.33</b>	0.05
<b>Subject Effect:</b>							
2	2	0.04	0.04	0.12	0.11	0.04	0.12
2	10	0.05	0.05	0.34	0.34	0.05	0.36
10	2	0.04	0.03	0.12	0.09	0.03	0.12
10	10	0.06	0.06	0.34	0.31	0.05	0.37

# Model Subjects as Fixed → Wrong Subject Effect

Condition		Models					
Item Variance	Subject Variance	1: Both Random Effects	2: Random Subjects Only	3: <b>Random Items Only</b>	4: No Random Effects	5: F1 Subjects ANOVA	6: <b>F2 Item ANOVA</b>
<b>Item Effect:</b>							
2	2	0.03	0.09	0.03	0.09	0.09	0.03
2	10	0.05	0.14	0.05	0.12	0.15	0.05
10	2	0.04	0.32	0.04	0.31	0.32	0.04
10	10	0.05	0.31	0.05	0.29	0.33	0.05
<b>Subject Effect:</b>							
2	2	0.04	0.04	<b>0.12</b>	0.11	0.04	<b>0.12</b>
2	10	0.05	0.05	<b>0.34</b>	0.34	0.05	<b>0.36</b>
10	2	0.04	0.03	<b>0.12</b>	0.09	0.03	<b>0.12</b>
10	10	0.06	0.06	<b>0.34</b>	0.31	0.05	<b>0.37</b>

# Conclusions

- An ANOVA model may be less than ideal when:
  - Stimuli are not completely controlled or exchangeable
  - Experimental conditions are not strictly discrete
  - Missing data may result in bias, a loss of power, or both
- ANOVA is a special case of a more general family of multilevel models (with nested or crossed effects as needed) that can offer additional flexibility:
  - Useful in addressing statistical problems →
    - Dependency, heterogeneity of variance, unbalanced or missing data
    - Examine predictor effects pertaining to each source of variation more accurately given that all variation is properly represented in the model
  - Useful in addressing substantive hypotheses →
    - Examining individual differences in effects of experimental manipulations