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S1. Supplemental Materials Part 1

Reproducibility in small-N treatment research: a tutorial using examples from aphasiology

Introduction

This document details the code needed to reproduce the analysis in section 1 of the manuscript. For batch calculations of effect sizes across participants see part 2.

Setup

Load packages and functions

set a seed for reproducibility
set.seed(42)

Read in data

Note that the current setup uses RStudio R projects (https://support.rstudio.com/hc/en-us/articles/200526207-Using-RStudio-Projects). One of the features of R projects is that the working directory is automatically set to the project root (the folder with the .Rproj). A discussion of R projects can be found at https://www.tidyverse.org/blog/2017/12/workflow-vs-script/. In this case here("study-data") refers to the /study-data folder inside the project.

Preview the data

head(df)

```
## # A tibble: 6 x 11
##
     participant condi~1 phoneme itemT~2 phase session item trials spt2017 respo~3
##
     <chr>
                 <chr>
                         <chr>
                                  <chr>
                                          <chr>
                                                  <dbl> <chr> <dbl> <chr>
                                                                                <dbl>
## 1 P1
                 blocked pr
                                 tx
                                          base~
                                                      1 pr-1
                                                                   10 pre
                                                                                    0
## 2 P1
                 blocked pr
                                          base~
                                                      1 pr-12
                                                                   10 pre
                                                                                    0
                                 tx
                                                      1 pr-4
## 3 P1
                                                                                    0
                 blocked pr
                                          base~
                                                                   10 pre
                                 \mathtt{tx}
                                                      1 pr-15
## 4 P1
                 blocked pr
                                          base~
                                 tx
                                                                   10 pre
                                                                                    0
## 5 P1
                 blocked pr
                                 tx
                                          base~
                                                      1 pr-5
                                                                   10 pre
                                                                                    0
## 6 P1
                 blocked pr
                                 tx
                                          base~
                                                      1 pr-7
                                                                   10 pre
                                                                                    0
## # ... with 1 more variable: n_baselines <dbl>, and abbreviated variable names
       1: condition, 2: itemType, 3: response
## #
## # i Use 'colnames()' to see all variable names
```

Variable	Description
participant condition phoneme	de-identified participant ID probe schedule (blocked or random) target_phoneme
itemType phase	item condition (treatment or generalization) treatment phase
session item trials spt2017 response	session number from Wambaugh 2017 item identifier number of items in the list (per phoneme) phase used to calcualte effect sizes in Wambaugh et al., 2017 accuracy of participant response
n_baselines	Number of baseline sessions

Table 1: Data variables and their description

Case example: Participant 10

Filter data for Participant 10

Starting from the entire dataset, filter for participant 10, treated items, and the blocked condition. Then to calculate session-level data (the number of correct responses per session), group by session, and use the summarize function to calculate the number of correct responses per session. The group_by function also includes phase and spt2017 because we want to keep these variables in the summary data frame, but their addition doesn't affect grouping. The .groups argument removes the grouping after summarize.

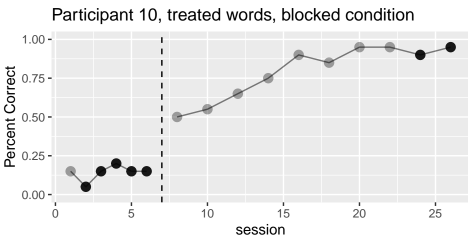
Plot performance over time

Plotting data from participant 10. First, we select only the baseline and treatment phases (ignoring the washout and maintenance phases for the purpose of this paper). The we create a dummy variable reflecting whether or not the session was included in the SMD/PMG calculations. Finally, we use the {ggplot2}

package to plot the data. A recent primer on $\{ggplot2\}$ for researchers unfamiliar with R can be found here: https://doi.org/10.1177/25152459221074654

```
p1 = P10 %>%
  # filter for baseline and treatment phases
  filter(phase == "baseline" | phase == "treatment") %>%
  # create a new variable called Measure that has a value of
  # include if spt2017 is not an NA value and exclude if it is.
  # the levels argument indicates that exclude should be
  # the first level of the factor.
  mutate(Measure = factor(
    ifelse(!is.na(spt2017), "include", "exclude"),
           levels = c("exclude", "include"))) %>%
  # create a plot with session on the x axis, percent
  # correct on the y axis, and group by phase
  ggplot(aes(x = session, y = sum_correct/20, group = phase)) +
  # add points to the graph
  geom_point(aes(alpha = Measure), size = 3) +
  # add a line to the graph
  geom_line(alpha = 0.5) +
  # add a vertical line where x = 7
  geom_vline(aes(xintercept = 7), linetype = "dashed") +
  scale_x_continuous(breaks = seq(0,30,5)) +
  vlim(0, 1) +
  scale_alpha_discrete(range = c(0.35, 0.9)) +
  labs(title = "Participant 10, treated words, blocked condition",
       caption = "Dark circles represent data points used to calculate
       the within-case standardized mean difference in Wambaugh et al.,
       (2017)",
       y="Percent Correct") +
  guides(alpha = "none")
```

```
p1
```



Dark circles represent data points used to calculate the within-case standardized mean difference in Wambaugh et al., (2017)

Within-case standardized mean difference

There are any number of ways to calculate the within case standardized mean difference using R code. In this example, we have used the SMD() function from the established package {SingleCaseES} by James Pustejovsky because it includes additional functions that may be of interest to researchers in aphasiology.

Additionally, we do not show all information returned by the function, which also includes a 95% confidence interval, as it is not clear that this confidence interval applies to the the $d_{\rm BR}$ modification of the original within-case standardized mean difference.

$$d_{BR} = \frac{x_B - x_A}{S_A}$$

A = P10 %>% filter(spt2017 == "pre") %>% pull(sum_correct)
B = P10 %>% filter(spt2017 == "post") %>% pull(sum_correct)
SMD(A_data = A, B_data = B)

ES Est SE CI_lower CI_upper baseline_SD
1 SMD 11.46566 3.283023 5.031052 17.90027 1.095445

Proportion of potential maximal gain

There is no R package that includes a function to calculate PMG to our knowledge. However, creating such a function is relatively straightforward. A function that calculates PMG similar to the SMD() function from the {SingleCaseES} package might take the following form, with an additional argument for the number of items treated (nitems). The function calculates the mean of the A phase and B phase, and then calculates and returns the PMG value from the same data as $d_{\rm BR}$ above.

$$PMG = \frac{x_B - x_A}{n_{items} - x_A}$$

```
# the function is named PMG and takes 3 arguments:
# vectors of the a_data and b_data, and
# a single number indicating how many items were treated
PMG <- function(a_data, b_data, nitems){
  mean_a <- mean(a_data) # calculate mean of a_data
  mean_b <- mean(b_data) # calculate mean of b_data
  pmg <- (mean_b-mean_a)/(nitems-mean_a) # calculate PMG
  return(pmg) # return the PMG value.
```

}

PMG(a_data = A, b_data = B, nitems = 20)

[1] 0.9127907

Tau-U

The Tau-U family of effect sizes (and several other non-overlap measures) can be calculated using the $\{\text{SingleCaseES}\}\ \text{package}$. In this case, we use all data summarized in the P10 data frame (and not just the data used to calculate d_{BR}).

First, we estimate the trend line during the baseline phase, which can be generated by creating a simple linear model using the lm() function. The model includes the number of correct responses as the dependent variable and the session number as the independent variable. The session coefficient reflects the slope during the baseline phase. The coef() function simple extracts the model coefficients.

```
P10 %>%
filter(phase == "baseline") %>%
lm(data = ., sum_correct~session) %>%
coef()
```

##	(Intercept)	session
##	2.133333	0.200000

The Tau() and Tau_U() functions take the same data structure as the SMD() and PMG() functions above.

Using the conservative benchmark of 0.33 recommended by Lee and Cherney (2018), we would calculate $Tau_{Avs,B}$ as the slope of the baseline phase is only 0.2 (from the result above). To calculate $Tau_{Avs,B}$, we can use the Tau() function.

```
A = P10 %>% filter(phase == "baseline") %>% pull(sum_correct)
B = P10 %>% filter(phase == "treatment") %>% pull(sum_correct)
Tau(A_data = A, B_data = B)
```

ES Est SE CI_lower CI_upper ## 1 Tau 1 0.02710291 1 1

However, if we had elected to correct for baseline trends and use $Tau_{Avs.B-trendA}$, we can use the similar Tau_U() function.

 $Tau_U(A_{data} = A, B_{data} = B)$

ES Est ## 1 Tau-U 0.95

Mixed-effects model-based effect sizes

The mixed-effects model example for participant 10 uses item-level data, so we need to create a new dataframe for this model. As discussed in the manuscript, the model formula is based on a structure from Huitema & McKean (2000).

 $Y_t = \beta_0 + \beta_1 T_t + \beta_2 D_t + \beta_3 [T_t - (n_1 + 1)] D_t + \epsilon_t$

The DV is predicted by the model intercept, baseline slope, level change, and slope change parameters.

After selecting data from participant 10, the coefficients are created by:

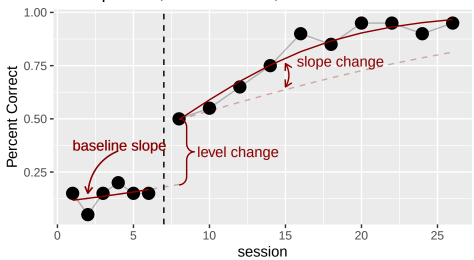
- setting baseline_slope equal to the session variable
- level_change is a dummy variable, 0 during baseline and 1 during treatment
- slope_change is created by subtracting the number of baselines plus 2 from the baseline slope value, and then multiplying the result with the level_change variable. Typically, if probing every session, the formula calls for subtracting the number of baselines plus 1. However, because Wambaugh et al., (2017) used intermittent probing schedules, and probed every other treatment session starting at the second, we need to add 2 to the number of baselines to ensure that the slope change variable starts at 0 on the first recorded treatment probe.

```
P10 <- df %>%
filter(participant == "P10",
    condition == "blocked",
    itemType == "tx",
    phase == "baseline" | phase == "treatment") %>%
mutate(baseline_slope = session,
    level_change = ifelse(phase == "baseline", 0, 1),
    slope_change = (baseline_slope - (6+2))*level_change,
    level_change = as.factor(level_change))
```

The resulting matrix looks as follows:

phase	baseline_slope	level_change	slope_change
baseline	1	0	0
baseline	2	0	0
baseline	3	0	0
baseline	4	0	0
baseline	5	0	0
baseline	6	0	0
treatment	8	1	0
treatment	10	1	2
treatment	12	1	4
treatment	14	1	6
treatment	16	1	8
treatment	18	1	10
treatment	20	1	12
treatment	22	1	14
treatment	24	1	16
treatment	26	1	18

Manuscript Figure 2. visualizes each parameter in this model structure. The code can be found in the .Rmd file, and is omitted from the pdf due to its length.



Participant 10, treated words, blocked condition

The model formula can be expressed as follows:

$$\operatorname{response}_{i} \sim \operatorname{Binomial}(n = 1, \operatorname{prob}_{\operatorname{response}=1} = \widehat{P})$$

$$\log \left[\frac{\widehat{P}}{1 - \widehat{P}}\right] = \alpha_{j[i]} + \beta_{1j[i]}T_{t} + \beta_{2j[i]}D_{t} + \beta_{3j[i]}[T_{t} - (n_{1} + 1)]D_{t}$$

$$\begin{pmatrix}\alpha_{j}\\\beta_{1j}\\\beta_{2j}\\\beta_{3j}\end{pmatrix} \sim N \left(\begin{pmatrix}\mu_{\alpha_{j}}\\\mu_{\beta_{1j}}\\\mu_{\beta_{2j}}\\\mu_{\beta_{3j}}\end{pmatrix}, \begin{pmatrix}\sigma_{\alpha_{j}}^{2} - \rho_{\alpha_{j}\beta_{1j}} - \rho_{\alpha_{j}\beta_{2j}} - \rho_{\alpha_{j}\beta_{3j}}\\\rho_{\beta_{1j}\alpha_{j}} - \sigma_{\beta_{1j}}^{2} - \rho_{\beta_{1j}\beta_{2j}} - \rho_{\beta_{1j}\beta_{3j}}\\\rho_{\beta_{2j}\alpha_{j}} - \rho_{\beta_{2j}\beta_{1j}} - \sigma_{\beta_{2j}}^{2} - \rho_{\beta_{2j}\beta_{3j}}\\\rho_{\beta_{3j}\alpha_{j}} - \rho_{\beta_{3j}\beta_{1j}} - \rho_{\beta_{3j}\beta_{2j}} - \sigma_{\beta_{3j}}^{2}\end{pmatrix}\right), \text{ for item } j = 1, \dots, J$$

The first line of the equation indicates that the dependent variable takes a binomial distribution. The second line represents the extension of the Huitema & McKean (2000) model to the hierarchical case, where each fixed effect is estimated for each item, j[i]. The third line of the equation represents the random effect structure, indicating that the effect of each fixed effect (baseline slope, level change, and slope change) are allowed to vary for each item.

The following shows how we arrived at the final model for P10

1. First, we fit the maximal random effects structure. However, the model did not converge.

```
mod1 <-
glmer(
    # fixed effects
    response ~ baseline_slope + level_change + slope_change +
    # random effects
    (1 + baseline_slope + level_change + slope_change | item),
    data = P10,
    family = binomial)</pre>
```

Warning in checkConv(attr(opt, "derivs"), opt\$par, ctrl = control\$checkConv, :
unable to evaluate scaled gradient

```
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
## Model failed to converge: degenerate Hessian with 1 negative eigenvalues
```

2. Second, we tried specifying a different optimizer, following recommendations that can be found at https://bbolker.github.io/mixedmodels-misc/glmmFAQ.html#convergence-warnings.

```
family = binomial,
# for model convergence
control = glmerControl(optimizer="bobyqa"))
```

We can now examine the model summary:

summary(mod1)

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
##
   Family: binomial (logit)
## Formula: response ~ baseline_slope + level_change + slope_change + (1 +
##
       baseline_slope + level_change + slope_change | item)
##
      Data: P10
## Control: glmerControl(optimizer = "bobyqa")
##
##
        AIC
                       logLik deviance df.resid
                 BIC
      249.5
                       -110.8
##
               302.3
                                 221.5
                                             306
##
## Scaled residuals:
                1Q Median
##
       Min
                                30
                                       Max
## -2.2995 -0.2462 0.0202 0.1563
                                   3.6534
##
## Random effects:
##
   Groups Name
                          Variance Std.Dev. Corr
   item
           (Intercept)
                          4.5354
                                   2.1296
##
                                             -0.75
##
           baseline_slope 0.3359
                                   0.5795
##
           level_change1 5.6515
                                   2.3773
                                             0.34 -0.01
##
           slope_change
                          0.8646
                                   0.9298
                                             0.66 -0.97 -0.14
## Number of obs: 320, groups: item, 20
##
## Fixed effects:
##
                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                  -2.70900
                              1.33242 -2.033
                                                  0.042 *
## baseline_slope 0.01742
                              0.33249
                                        0.052
                                                  0.958
## level_change1
                   2.50788
                              1.69433
                                        1.480
                                                  0.139
## slope change
                   0.39217
                              0.39651
                                        0.989
                                                  0.323
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
               (Intr) bsln_s lvl_c1
## baselin_slp -0.861
## level_chng1 0.535 -0.723
## slope_chang 0.761 -0.916 0.499
```

We note that in this case, further reducing the random effects structure often returns a significant result for the level_change parameter, demonstrating how our choice of random effect structure can influence the statistical significance of model parameters. For this reason, we stress that researchers are forthcoming of the model modifications made from the initial maximal random effect structure and also report what modifications result in parameters changing in significance.

Calculating an overall effect size for this participant requires contrasting performance either at the end of treatment with and without the level change and slope change parameters, or contrasting performance at the end of treatment with performance at the end of baseline. The former option assumes that any baseline trend would have continued throughout the treatment phase in the absence of treatment, is typically more conservative.

While there is a small, empirical baseline slope in this data, it may be reasonable to consider that this slope is largely driven by lower performance on the second probe session, and that performance in baseline sessions 3-6 are stable, and therefore estimate the difference in performance from the end of baseline to the end of treatment. Criteria for such decisions should ideally be made a-priori if possible, and reported in publications regardless.

1. First, we generate the marginal means for each combination of baseline slope, level change, and slope change.

marginal_means

##	<pre>baseline_slope</pre>	level_change	<pre>slope_change</pre>	emmean	SE	df	asymp.LCL	asymp.UCL
##	7	0	0	-2.5870	1.36	${\tt Inf}$	-5.252	0.0782
##	26	0	0	-2.2560	7.53	${\tt Inf}$	-17.010	12.4976
##	7	1	0	-0.0792	1.20	Inf	-2.427	2.2684
##	26	1	0	0.2519	6.39	Inf	-12.263	12.7672
##	7	0	19	4.8641	6.46	${\tt Inf}$	-7.801	17.5290
##	26	0	19	5.1952	3.06	${\tt Inf}$	-0.803	11.1928
##	7	1	19	7.3720	7.35	Inf	-7.041	21.7847
##	26	1	19	7.7030	2.49	Inf	2.819	12.5873
##								

Results are given on the logit (not the response) scale.
Confidence level used: 0.95

2. This returns a table of all possible comparisons, and we are only interested in contrasting the first row (beginning of treatment) with the last row (end of treatment). After selecting these two rows, we can then contrast their estimates.

```
# code to select first and last rows
# The 1 indicates that the row should be selected
# There are 8 numbers corresponding to the 8 possible comparisons above
A = c(1, 0, 0, 0, 0, 0, 0, 0)
B = c(0, 0, 0, 0, 0, 0, 0, 1)
# contrast the marginal means
# infer argument returns a confidence interval and p value if
# both are set to TRUE.
contrast(marginal_means,
    method = list("Unadjusted effect size" = B-A),
    infer = c(TRUE, TRUE))
```

```
## contrast estimate SE df asymp.LCL asymp.UCL z.ratio p.value
## Unadjusted effect size 10.3 3.1 Inf 4.22 16.4 3.322 0.0009
##
## Results are given on the log odds ratio (not the response) scale.
## Confidence level used: 0.95
```

We could also make the more conservative assumption that any baseline trend continues by choosing the second row where baseline slope is set to the last treatment session.

```
# code to select first and last rows
# The 1 indicates that the row should be selected
A = c(0, 1, 0, 0, 0, 0, 0, 0)
B = c(0, 0, 0, 0, 0, 0, 0, 1)
# contrast the marginal means
# infer argument returns a confidence interval and p value if
# both are set to TRUE.
contrast(marginal_means,
    method = list("Unadjusted effect size" = B-A),
    infer = c(TRUE, TRUE))
```

```
## contrast estimate SE df asymp.LCL asymp.UCL z.ratio p.value
## Unadjusted effect size 9.96 8.51 Inf -6.71 26.6 1.171 0.2417
##
## Results are given on the log odds ratio (not the response) scale.
## Confidence level used: 0.95
```

Notice that there is much greater uncertainty in this contrast, evident by the increase in standard error, and as a result the p-value is no longer significant.

Group-level model

We can extend this individual model to all participants, still focusing on treated items in the blocked condition. First, we create a new data frame that includes all participants.

```
df_glmm <- df %>%
  # select the correct phase, condition, and itemType
  filter(phase == "baseline" | phase == "treatment",
        condition == "blocked",
        itemType == "tx") %>%
  # create the Huitema model parameters
  mutate(baseline_slope = session,
        level_change = ifelse(phase == "baseline", 0, 1),
        slope_change = (baseline_slope - (n_baselines+2))*level_change,
        level_change = as.factor(level_change))
```

Then we can start again with a relatively maximal random effect structures, noting that we could also include random slopes for items. However, it is unlikely that such a model structure could be supported by the data. In this case we have chosen to include the most theoretically important random effects (Matsucheck, 2018) that we expect to be supported by the data. A formula for the model is:

$$\begin{aligned} \operatorname{response}_{i} \sim \operatorname{Binomial}(n = 1, \operatorname{prob}_{\operatorname{response}=1} = \widehat{P}) \\ \log \left[\frac{\widehat{P}}{1 - \widehat{P}} \right] &= \alpha_{j[i],k[i]} + \beta_{1k[i]}(\operatorname{baseline_slope}) + \beta_{2k[i]}(\operatorname{level_change}_{1}) + \beta_{3k[i]}(\operatorname{slope_change}) \\ &\quad \alpha_{j} \sim N\left(\mu_{\alpha_{j}}, \sigma_{\alpha_{j}}^{2}\right), \text{ for item } j = 1, \dots, J \\ \left(\begin{array}{c} \alpha_{k} \\ \beta_{1k} \\ \beta_{2k} \\ \beta_{3k} \end{array} \right) \sim N\left(\begin{pmatrix} \mu_{\alpha_{k}} \\ \mu_{\beta_{1k}} \\ \mu_{\beta_{2k}} \\ \mu_{\beta_{3k}} \end{pmatrix}, \begin{pmatrix} \sigma_{\alpha_{k}}^{2} & \rho_{\alpha_{k}\beta_{1k}} & \rho_{\alpha_{k}\beta_{2k}} & \rho_{\alpha_{k}\beta_{3k}} \\ \rho_{\beta_{1k}\alpha_{k}} & \sigma_{\beta_{1k}}^{2} & \rho_{\beta_{1k}\beta_{2k}} & \rho_{\beta_{1k}\beta_{3k}} \\ \rho_{\beta_{2k}\alpha_{k}} & \rho_{\beta_{2k}\beta_{1k}} & \sigma_{\beta_{2k}}^{2} & \rho_{\beta_{2k}\beta_{3k}} \\ \rho_{\beta_{3k}\alpha_{k}} & \rho_{\beta_{3k}\beta_{1k}} & \rho_{\beta_{3k}\beta_{2k}} & \sigma_{\beta_{3k}}^{2} \end{pmatrix} \right) \end{aligned} \right), \text{ for participant } k = 1, \dots, K \end{aligned}$$

The model takes a little longer to run, and returns a convergence warning

```
(1|item),
    data = df_glmm,
    family = binomial)
```

```
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
## Model failed to converge with max|grad| = 0.088281 (tol = 0.002, component 1)
```

Again, we change the optimizer.

Since the model appears to have converged, we can examine the model results

summary(mod2)

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
##
     Approximation) [glmerMod]
##
   Family: binomial (logit)
## Formula: response ~ baseline_slope + level_change + slope_change + (1 +
##
       baseline_slope + level_change + slope_change | participant) +
##
       (1 | item)
      Data: df_glmm
##
## Control: glmerControl(optimizer = "bobyqa")
##
##
        AIC
                       logLik deviance df.resid
                 BIC
##
     5652.2
             5755.8 -2811.1 5622.2
                                           7385
##
## Scaled residuals:
              1Q Median
                                ЗQ
##
      Min
                                       Max
## -8.1144 -0.3551 -0.1630 0.3241 13.5509
##
## Random effects:
                               Variance Std.Dev. Corr
## Groups
                Name
##
                (Intercept)
                               1.611095 1.26929
   item
##
   participant (Intercept)
                               0.404805 0.63624
##
                                                 0.41
                baseline_slope 0.002648 0.05146
##
                level_change1 2.390619 1.54616
                                                 0.18 0.86
##
                slope_change
                              0.007304 0.08546 -0.78 -0.54 -0.52
## Number of obs: 7400, groups: item, 322; participant, 20
##
## Fixed effects:
##
                  Estimate Std. Error z value Pr(>|z|)
                 -3.44259 0.20649 -16.672 < 2e-16 ***
## (Intercept)
## baseline_slope 0.07624
                                       4.050 5.13e-05 ***
                             0.01883
```

```
0.85948
                              0.39756
                                        2.162 0.030628 *
## level_change1
                              0.02582
                                        3.534 0.000409 ***
## slope_change
                   0.09124
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
               (Intr) bsln_s lvl_c1
## baselin_slp -0.161
## level_chng1 0.096 0.310
## slope_chang -0.192 -0.666 -0.363
```

The summary table shows us there there are statistically significant effects for all three parameters: a small but significant trend at baseline, a fairly substantial level change on average, and an increase in slope from baseline that is slightly more than double the initial average trend. We also note that there there is much more variation in the level change parameter between participants relative to the baseline slope and slope change parameters. Finally, the correlation of fixed effects shows a positive association between individuals baseline trend and their level change, but a negative association between individuals baseline trend and slope change and level change.

We can calculate an overall effect size using the same approach as the individual model. In this case, we assume the median number of baseline sessions (11) and treatment sessions (20)

marginal_means

```
baseline_slope level_change slope_change emmean
##
                                                          SE
                                                             df asymp.LCL asymp.UCL
                                            0 -2.6040 0.268 Inf
##
                                                                    -3.1289
                                                                             -2.07906
                11 0
##
                31 0
                                             0 -1.0792 0.587 Inf
                                                                    -2.2294
                                                                              0.07090
##
                                                                    -2.8120
                                                                             -0.67696
                11 1
                                            0 -1.7445 0.545 Inf
##
                31 1
                                              -0.2197 0.814 Inf
                                                                    -1.8147
                                                                              1.37518
                                             0
##
                11 0
                                           20 -0.7791 0.393 Inf
                                                                   -1.5503
                                                                             -0.00788
                31 0
                                                0.7457 0.410 Inf
                                                                    -0.0586
##
                                           20
                                                                              1.54996
                                                0.0804 0.480 Inf
##
                11 1
                                           20
                                                                    -0.8613
                                                                              1.02206
##
                31 1
                                           20
                                               1.6051 0.581 Inf
                                                                    0.4668
                                                                              2.74346
##
## Results are given on the logit (not the response) scale.
## Confidence level used: 0.95
```

Because the baseline trend, on average, was statistically reliable, we calculated an overall effect size assuming that it would have continued in the absence of treatment.

```
# code to select first and last rows
# The 1 indicates that the row should be selected
A = c(0, 1, 0, 0, 0, 0, 0, 0)
B = c(0, 0, 0, 0, 0, 0, 0, 1)
# contrast the marginal means
# infer argument returns a confidence interval and p value if
# both are set to TRUE.
contrast(marginal_means,
    method = list("Unadjusted effect size" = B-A),
    infer = c(TRUE, TRUE))
```

```
## contrast estimate SE df asymp.LCL asymp.UCL z.ratio p.value
## Unadjusted effect size 2.68 0.525 Inf 1.66 3.71 5.112 <.0001
##
## Results are given on the log odds ratio (not the response) scale.
## Confidence level used: 0.95</pre>
```

This results in a statistically reliable group effect size of 2.7 logits. Given that the group model suggests a starting place of only around 3% (found by converting logits to percentage plogis(-3.44)), this indicates a gain of about 35 percentage points on average can be attributed to the level and slope changes. We can estimate change in percentage points by adding the intercept, the baseline slope times an average number of baseline sessions, and the predicted change, converting to the predicted percent correct, and then subtracting the percent correct at baseline plus the baseline change: plogis(-3.44 + 0.07*5 + 2.68)-plogis(-3.44 + 0.07*5). However, we're not aware of a straightforward method of estimating individual effect sizes and confidence intervals using the frequentist approach.

Bayesian Mixed effects models

Bayesian mixed-effects models can be used in the same fashion as model 2 above to obtain both group and individual effect size estimates. First, a group-level model is estimated.

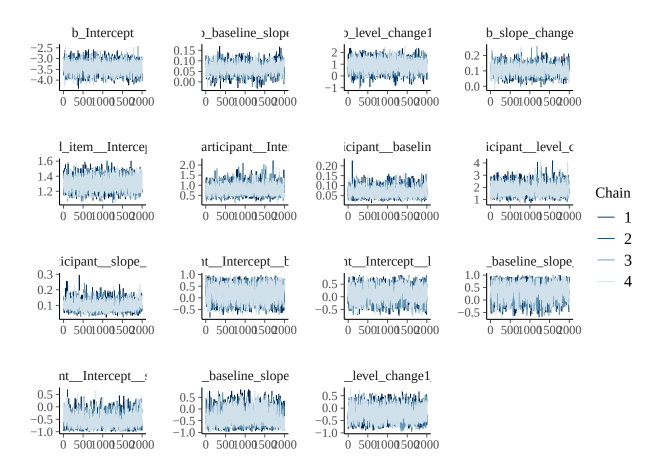
```
family = bernoulli(), # special case of binomial with n=1 trials
iter = 3000, # number of draws per chain
warmup = 1000, # number of draws to toss on "burn in"
chains = 4, # total number of chains
seed = 42, # set a seed
prior = c( # prior distributions
prior(normal(-1, 2), class = b, coef = Intercept),
prior(normal(0, 2.5), class = b)
),
# extra arguments, see rmd file
cores = 4,
file = here("output", "group_brm"),
file_refit = "on_change")
```

We can check several aspects of model fit and convergence.

1. Traceplots: Note that there are four lines within each plot, representing all the samples from each Markov Chain Monte Carlo (MCMC) simulation from the above model. There should be no noticeable variations in the patterns of the estimates across each model. Visual inspection of convergence is achieved by affirming that the MCMC estimates were consistently sampled from a range of values. Graphically, this is presented by the horizontal structure in each trace plot. For example, all estimates of model 3s intercept, b_Intercept fall within a range of about -4 and -2.5 logits. "Spikes" observed in traceplots are normal, as can be seen in the participant_slope plot. These spikes represent estimates at the tail end of a probability distribution. Traceplots with positive or negative slopes or sinusoidal wave shapes do not reflect convergence.

brms::mcmc_plot(mod3, type = "trace")

No divergences to plot.



2. Check that the model can successfully re-estimate the data. In this case, we conducted posterior predictive checks on three statistics that can be estimated from the model. The posterior_predict function creates n number of posterior distributions, where n is the total number of post warmup iterations for one MCMC chain. We label these distributions **yrep** in the code below.

Then, we estimate posterior predictive p-values for the mean, standard deviation, and kurtosis of the n number of posteriors included in yrep. We note, that this is only one type of method of posterior checking, and was chosen because graphical posterior predictive checks of the predicted number outcome estimates from logistic regression models involving a large number of 0s can be misleading.

```
y = mod3$data$response
yrep = posterior_predict(mod3)
mean_ppc = mean(apply(yrep, 1, mean) > mean(y))
sd_ppc = mean(apply(yrep, 1, sd) > sd(y))
kurtosis_ppc = mean(apply(yrep, 1, e1071::kurtosis) > e1071::kurtosis(y))
print(c(mean_ppc, sd_ppc, kurtosis_ppc))
```

[1] 0.506625 0.510625 0.489125

mean_ppc, sd_ppc, and kurtosis_ppc represent the proportion of mean, standard deviation, and kurtosis estimates less than the observed estimates of these parameters. The p-value is the proportion of posterior samples with greater prediction error than the actual data. Low p-values mean worse model fit than expected if the model were correct. Values near 0.5 are ideal.

3. Rhat statistic should be < 1.05, ideally < 1.01

The rhat stastistic represents the potential scale reduction factor across the split chains, the case of this model across the four chains. A value of 1.0 represents perfect convergence and estimates between 1 and 1.05, and ideally between 1 and 1.01 represent adequate convergence. Model fitting should be reevaluated with rhat statistics greater than 1.05.

max(rhat(mod3))

[1] 1.003422

We can preview the model results using summary() again. Notably, the model estimates are largely similar to the frequentist model.

```
summary(mod3)
```

```
##
    Family: bernoulli
##
    Links: mu = logit
## Formula: response ~ 0 + Intercept + baseline_slope + level_change + slope_change + (1 + baseline_slop
      Data: df_glmm (Number of observations: 7400)
##
     Draws: 4 chains, each with iter = 3000; warmup = 1000; thin = 1;
##
##
            total post-warmup draws = 8000
##
## Group-Level Effects:
## ~item (Number of levels: 322)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                     1.30
                                0.08
                                          1.15
                                                   1.46 1.00
                                                                  2366
                                                                           4243
##
## ~participant (Number of levels: 20)
##
                                      Estimate Est.Error 1-95% CI u-95% CI Rhat
## sd(Intercept)
                                          0.77
                                                     0.23
                                                              0.40
                                                                        1.28 1.00
                                          0.06
                                                     0.02
## sd(baseline_slope)
                                                              0.03
                                                                        0.11 1.00
## sd(level_change1)
                                          1.80
                                                     0.40
                                                              1.12
                                                                        2.70 1.00
## sd(slope_change)
                                          0.10
                                                     0.03
                                                              0.05
                                                                        0.16 1.00
## cor(Intercept,baseline_slope)
                                          0.21
                                                     0.33
                                                             -0.45
                                                                        0.79 1.00
## cor(Intercept,level_change1)
                                                             -0.43
                                          0.15
                                                     0.28
                                                                        0.65 1.00
## cor(baseline_slope,level_change1)
                                          0.36
                                                     0.32
                                                             -0.32
                                                                        0.87 1.00
## cor(Intercept,slope_change)
                                          -0.56
                                                     0.25
                                                             -0.93
                                                                        0.02 1.00
## cor(baseline slope,slope change)
                                          -0.37
                                                             -0.85
                                                     0.31
                                                                        0.33 1.00
## cor(level_change1,slope_change)
                                          -0.22
                                                     0.28
                                                             -0.72
                                                                        0.35 1.00
```

##				Bulk_ESS	Tail_ESS			
##	<pre>sd(Intercept)</pre>	2463	3834					
##	<pre>sd(baseline_slc</pre>	pe)		2559	3731			
##	sd(level_change	1)		3090	4799			
##	sd(slope_change)		1180	2898			
##	cor(Intercept, b	aseline_slope))	1883	3701			
##	cor(Intercept,1	evel_change1)		1310	2675			
##	<pre>cor(baseline_sl</pre>	ope,level_char	ige1)	915	1814			
##	cor(Intercept,s	lope_change)		1053	2559			
##	cor(baseline_slope,slope_change)				1958			
##	<pre>t cor(level_change1,slope_change)</pre>				2921			
##								
##	Population-Leve	l Effects:						
##		Estimate Est.H	Error	1-95% CI	u-95% CI	Rhat	$Bulk_ESS$	Tail_ESS
##	Intercept	-3.41	0.24	-3.89	-2.96	1.00	2118	3857
##	<pre>baseline_slope</pre>	0.06	0.02	0.02	0.11	1.00	1309	2171
##	level_change1	0.90	0.45	-0.01	1.78	1.00	2587	3970
##	<pre>slope_change</pre>	0.11	0.03	0.05	0.17	1.00	1582	2628
##								
##	Draws were samp	led using sam	oling	(NUTS). F	or each pa	aramet	ter, Bulk ₋	_ESS
##	and Tail_ESS ar	e effective sa	ample	size mea	sures, and	l Rhat	t is the p	potential
##	scale reduction	factor on spl	Lit c	hains (at	convergen	nce, H	Rhat = 1)	

Finally, we can calculate effect sizes from the model for each individual

To do this efficiently, we wrote a function which takes arguments for the model object, and an argument we called "adjust" that can be TRUE if we would like to extrapolate the baseline slope through the end of treatment and FALSE otherwise. The default is FALSE.

The function works be selecting rows for each participant and item in the data and estimating the posterior distribution for the values in each row.

Then the data is transformed and the posterior distribution at the beginning of treatment (or at the end of treatment without the level change and slope change parameters) is subtracted from the posterior distribution at the end of treatment.

The resulting posterior distribution characterized the magnitude of change, the mean or median can be used as a point estimate and the middle 95% of the distribution is the 95% credible interval.

```
glmmES = function(fit, adjust = FALSE){
```

```
# start with the data that went into the model,
# for each participant and phase (here we just used level_change
# because they are equivalent) make a new variable called last_session
# which is the highest value of the baseline slope coefficient
# then filter for only rows where the baseline slope is
# equal to the highest value in the phase (in other words
# this selects the last baseline and last treatment session).
```

```
# Remove the response column, reduce the data frame to only the
# unique rows
data = fit$data %>%
    group_by(level_change, participant) %>%
    mutate(last session = max(baseline slope)) %>%
    filter(baseline_slope == last_session) %>%
    select(-response) %>%
    distinct()
# If adjust is TRUE, then for each participant,
# set the baseline slope to always equal the highest value
# of baseline slope.
# in other words, we end up with a row where baseline slope
# represents the last treatment session, but level and slope
# change are still zero.
if(adjust){
 data = data %>%
    group_by(participant) %>%
    mutate(baseline_slope = max(baseline_slope))
}
# calculate an effect size in logits or log-odds
# start with the data frame we just created and add
# model draws from the linear/link-level predictor
# change timepoint to entry if level change is
# 0 and exit if 1. Select only the needed columns
# and take the data from long to wide. using pivot wider
# Then subtract the value for the draw for entry from
# the draw for exit.
# then for each participant, calculate the point
# estimate and interval using point_interval
# the last line just adds a column indicating this effect size is in logits
linepred = data %>%
  add_linpred_draws(fit) %>%
  ungroup() %>%
  mutate(timepoint = ifelse(level change == 0, "entry", "exit")) %>%
  select(timepoint, item, .draw, .linpred, participant) %>%
  pivot_wider(names_from = "timepoint", values_from = .linpred) %>%
  mutate(ES = exit-entry) %>%
  group_by(participant) %>%
  point_interval(ES) %>%
  mutate(unit = "logit")
# This block does the same thing, except using the expectation of the
# posterior (i.e., in percent correct terms)
epred = data %>%
  add_epred_draws(fit) %>%
  ungroup() %>%
  mutate(timepoint = ifelse(level_change == 0, "entry", "exit")) %>%
  select(timepoint, .draw, item, .epred, participant) %>%
  pivot_wider(names_from = "timepoint", values_from = .epred) %>%
  mutate(ES = exit-entry) %>%
  group_by(participant) %>%
  point_interval(ES) %>%
```

participant	ES	.lower	.upper	.width	.point	.interval	unit
P1	2.1895495	-0.3676083	4.988229	0.95	median	qi	logit
P10	4.0086213	1.5187068	6.748027	0.95	median	qi	logit
P11	2.9714045	1.5204160	4.609804	0.95	median	qi	logit
P12	3.0598628	1.9864749	4.130898	0.95	median	qi	logit
P13	5.8589987	4.3140540	7.506057	0.95	median	qi	logit
P14	3.9626795	2.6624822	5.335825	0.95	median	qi	logit
P15	2.2477880	0.6808346	3.828825	0.95	median	qi	logit
P16	7.2926630	4.4957225	11.309594	0.95	median	qi	logit
P17	6.2582404	3.5445388	10.266857	0.95	median	qi	logit
P18	1.5851814	-0.0863041	3.492992	0.95	median	qi	logit
P19	5.4483097	2.8257044	8.691806	0.95	median	qi	logit
P2	5.7103733	2.8251632	9.211314	0.95	median	qi	logit
P20	3.4349465	2.2776165	4.745630	0.95	median	qi	logit
P3	5.0591621	2.4579499	8.537965	0.95	median	qi	logit
P4	4.6388253	2.4688143	7.444193	0.95	median	qi	logit
P5	2.8040056	1.1536990	4.869518	0.95	median	qi	logit
P6	3.7047279	1.2918042	6.876803	0.95	median	qi	logit
$\mathbf{P7}$	0.2205605	-2.6562366	2.434720	0.95	median	qi	logit
P8	2.2802549	-0.1926789	4.786389	0.95	median	qi	logit
P9	1.1980474	-1.1147377	3.580816	0.95	median	qi	$\log it$

```
mutate(unit = "percent")
```

```
return(bind_rows(linepred, epred))
```

Here's how we might use the function, adjusting for baseline slope

bayesian_es = glmmES(mod3, adjust = TRUE)

Examine the results for logits

}

head(bayesian_es %>% filter(unit == "logit"), 20) %>% kable(format = "latex", booktabs = TRUE) %>%
kable_styling(position = "center")

Examine the results for percent

```
head(bayesian_es %>% filter(unit == "percent"), 20) %>% kable(format = "latex", booktabs = TRUE) %>%
kable_styling(position = "center")
```

To convert to odds ratios, exponentiate the logit effect sizes. To convert to number of items correct, multiply the percent gain by the number of items treated

participant	ES	.lower	.upper	.width	.point	.interval	unit
P1	0.3181775	-0.0483361	0.7443294	0.95	median	qi	percent
P10	0.4250740	0.0199313	0.8940472	0.95	median	qi	percent
P11	0.0869928	0.0082954	0.6206214	0.95	median	qi	percent
P12	0.5262244	0.1291868	0.7416138	0.95	median	qi	percent
P13	0.8543259	0.5419068	0.9424602	0.95	median	qi	percent
P14	0.6685233	0.2632803	0.8466362	0.95	median	qi	percent
P15	0.3431008	0.0433472	0.6799142	0.95	median	qi	percent
P16	0.8872033	0.5491569	0.9820621	0.95	median	$\mathbf{q}\mathbf{i}$	percent
P17	0.5681732	0.0907430	0.9129990	0.95	median	$\mathbf{q}\mathbf{i}$	percent
P18	0.0834094	-0.0056740	0.3588588	0.95	median	qi	percent
P19	0.7742871	0.1480239	0.9677798	0.95	median	qi	percent
P2	0.5197516	0.0352241	0.9583102	0.95	median	$\mathbf{q}\mathbf{i}$	percent
P20	0.4809946	0.0834361	0.7640616	0.95	median	qi	percent
P3	0.7847071	0.2275039	0.9540422	0.95	median	qi	percent
P4	0.7240730	0.1448200	0.9362476	0.95	median	qi	percent
P5	0.4817848	0.0932917	0.7780038	0.95	median	qi	percent
P6	0.6157490	0.2034815	0.8813546	0.95	median	qi	percent
$\mathbf{P7}$	0.0327783	-0.2965442	0.5060892	0.95	median	qi	percent
$\mathbf{P8}$	0.3184706	-0.0086866	0.7888893	0.95	median	qi	percent
P9	0.1379460	-0.1858118	0.6055480	0.95	median	qi	percent

```
sessionInfo()
```

```
## R version 4.2.1 (2022-06-23)
## Platform: aarch64-apple-darwin20 (64-bit)
## Running under: macOS Monterey 12.5
##
## Matrix products: default
## BLAS:
           /Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    base
##
## other attached packages:
## [1] officer_0.4.3
                                              forcats_0.5.1
                           flextable_0.7.2
                                                                  stringr_1.4.0
## [5] dplyr_1.0.9
                           purrr_0.3.4
                                              readr_2.1.2
                                                                  tidyr_1.2.0
## [9] tibble_3.1.8
                           ggplot2_3.3.6
                                              tidyverse_1.3.2
                                                                  ggdist_3.2.0
## [13] tidybayes_3.0.2
                           brms_2.17.0
                                              Rcpp_1.0.9
                                                                  emmeans_1.7.5
## [17] lme4_1.1-30
                           Matrix_1.4-1
                                              SingleCaseES_0.6.1 here_1.0.1
## [21] kableExtra_1.3.4
##
## loaded via a namespace (and not attached):
##
     [1] uuid_1.1-0
                              readxl_1.4.0
                                                   backports_1.4.1
##
     [4] systemfonts_1.0.4
                              plyr_1.8.7
                                                   igraph_1.3.4
     [7] splines_4.2.1
                              svUnit_1.0.6
                                                   crosstalk_1.2.0
##
  [10] rstantools_2.2.0
                                                   digest_0.6.29
                              inline_0.3.19
##
```

##	[13]	htmltools_0.5.3	fansi_1.0.3	magrittr_2.0.3
##		checkmate_2.1.0	ggbrace_0.1.0	googlesheets4_1.0.0
##		tzdb_0.3.0	modelr_0.1.8	RcppParallel_5.1.5
##		_ matrixStats_0.62.0	_ vroom_1.5.7	xts_0.12.1
##			_ prettyunits_1.1.1	_ colorspace_2.0-3
##		rvest_1.0.2	haven_2.5.0	xfun_0.31
##		jsonlite_1.8.0	callr_3.7.1	 crayon_1.5.1
##		zoo_1.8-10		gtable_0.3.0
##		gargle_1.2.0	webshot_0.5.3	distributional_0.3.0
##	[40]	pkgbuild_1.3.1	rstan_2.21.5	abind_1.4-5
##	[43]	scales_1.2.0	mvtnorm_1.1-3	DBI_1.1.3
##	[46]	miniUI_0.1.1.1	viridisLite_0.4.0	xtable_1.8-4
##	[49]	proxy_0.4-27	bit_4.0.4	stats4_4.2.1
##	[52]	StanHeaders_2.21.0-7	DT_0.23	htmlwidgets_1.5.4
##	[55]	httr_1.4.3	threejs_0.3.3	arrayhelpers_1.1-0
##	[58]	posterior_1.2.2	ellipsis_0.3.2	pkgconfig_2.0.3
##	[61]	100_2.5.1	farver_2.1.1	dbplyr_2.2.1
##	[64]	utf8_1.2.2	labeling_0.4.2	tidyselect_1.1.2
##	[67]	rlang_1.0.2	reshape2_1.4.4	later_1.3.0
##	[70]	munsell_0.5.0	cellranger_1.1.0	tools_4.2.1
##		cli_3.3.0	generics_0.1.3	broom_1.0.0
##		ggridges_0.5.3	evaluate_0.15	fastmap_1.1.0
##		yam1_2.3.5	bit64_4.0.5	processx_3.7.0
##		knitr_1.39	fs_1.5.2	zip_2.2.0
##		nlme_3.1-157	mime_0.12	projpred_2.1.2
##		xml2_1.3.3	compiler_4.2.1	bayesplot_1.9.0
##		<pre>shinythemes_1.2.0</pre>	rstudioapi_0.13	gamm4_0.2-6
##		e1071_1.7-11	reprex_2.0.1	stringi_1.7.8
##		ps_1.7.1	Brobdingnag_1.2-7	gdtools_0.2.4
##		lattice_0.20-45	nloptr_2.0.3	markdown_1.1
##		shinyjs_2.1.0	$tensorA_0.36.2$	vctrs_0.4.1
##		pillar_1.8.0	lifecycle_1.0.1	bridgesampling_1.1-2
		estimability_1.4	data.table_1.14.2	httpuv_1.6.5
		R6_2.5.1	promises_1.2.0.1	gridExtra_2.3
		codetools_0.2-18	boot_1.3-28	<pre>colourpicker_1.1.1</pre>
		MASS_7.3-57	gtools_3.9.3	assertthat_0.2.1
	[121]	rprojroot_2.0.3	withr_2.5.0	shinystan_2.6.0
		mgcv_1.8-40	parallel_4.2.1	hms_1.1.1
		grid_4.2.1	class_7.3-20	coda_0.19-4
		minqa_1.2.4	rmarkdown_2.14	<pre>googledrive_2.0.0</pre>
	[133]	lubridate_1.8.0	shiny_1.7.2	base64enc_0.1-3
##	[136]	dygraphs_1.1.1.6		