Supplement 3

Primary analysis: frequency of pain

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This script addresses the primary analysis of the study, namely, what is the frequency of pain in the last week in individuals attending an urban HIV voluntary counselling and testing centre, and does the frequency of pain differ between individuals infected by HIV and those who are not.

Inspect data

Data structure

```
dim(data)
## [1] 540
head(data)
## # A tibble: 6 x 3
     PID
           pain HIV_positive
     <chr> <chr> <chr>
## 1 001
           no
## 2 003
           yes
                 no
## 3 004
           yes
                 no
## 4 005
           yes
                 no
## 5 006
           yes
                 no
## 6 007
           no
                 no
tail(data)
```

```
## # A tibble: 6 x 3
##
          pain HIV_positive
   PID
    <chr> <chr> <chr>
## 1 536
          no
                 yes
## 2 537
          no
                yes
## 3 538
         no
                no
## 4 539
          yes
                yes
## 5 540
          no
                no
## 6 541
          nο
                yes
glimpse(data)
## Observations: 540
## Variables: 3
## $ PID
                  <chr> "001", "003", "004", "005", "006", "007", "008", ...
## $ pain
                  <chr> "no", "yes", "yes", "yes", "no", "yes", "y...
## $ HIV_positive <chr> "no", "no", "no", "no", "no", "no", "no", "no", "no", "...
# HIV status data
dim(data[data$HIV_positive == 'yes' & !is.na(data$HIV_positive), ])
## [1] 70 3
dim(data[data$HIV_positive == 'no' & !is.na(data$HIV_positive), ])
## [1] 465
dim(data[is.na(data$HIV_positive), ])
## [1] 5 3
dim(data[!is.na(data$HIV_positive), ])
## [1] 535
# Remove rows with missing HIV test results (n = 5)
data %<>%
   filter(!is.na(HIV_positive))
# By pain status
dim(data[data$pain == 'yes' & !is.na(data$pain), ])
## [1] 316
dim(data[data$pain == 'no' & !is.na(data$pain), ])
## [1] 219
dim(data[is.na(data$pain), ])
## [1] 0 3
dim(data[!is.na(data$pain), ])
## [1] 535
Data summaries
# Convert character to factor
data_fct <- data %>%
   mutate(HIV_positive = factor(HIV_positive),
          pain = factor(pain)) %>%
```

```
select(-PID)
# Full cohort
data_fct %>%
   skim() %>%
   kable(caption = 'Full cohort')
## Skim summary statistics
## n obs: 535
## n variables: 2
## Variable type: factor
##
                                                    top_counts
##
              missing complete n n_unique
    variable
## ----- ---- ----- ----- ----- -----
  HIV_positive 0 535 535 2 no: 465, yes: 70, NA: 0 pain 0 535 535 2 yes: 316, no: 219, NA: 0
##
##
# By HIV status
data_fct %>%
   group_by(pain) %>%
   skim() %>%
   kable(caption = 'By HIV status')
## Skim summary statistics
## n obs: 535
## n variables: 2
##
## Variable type: factor
##
## pain
          variable missing complete n n_unique
                                                          top_counts
## ----- ----- ------
## no HIV_positive 0
## yes HIV_positive 0
                                                 2 no: 184, yes: 35, NA: 0
2 no: 281, yes: 35, NA: 0
                                 219 219
316 316
```

Primary questions

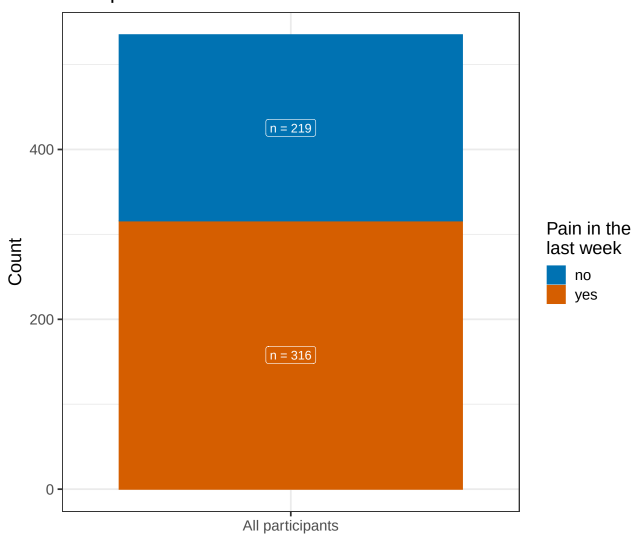
Frequency of pain

Exploratory plots

```
# Prepare data
plot_total <- data %>%
    group_by(pain) %>%
    summarise(count = n()) %>%
    mutate(prop = round(count/sum(count) * 100, 1)) %>%
    ungroup()

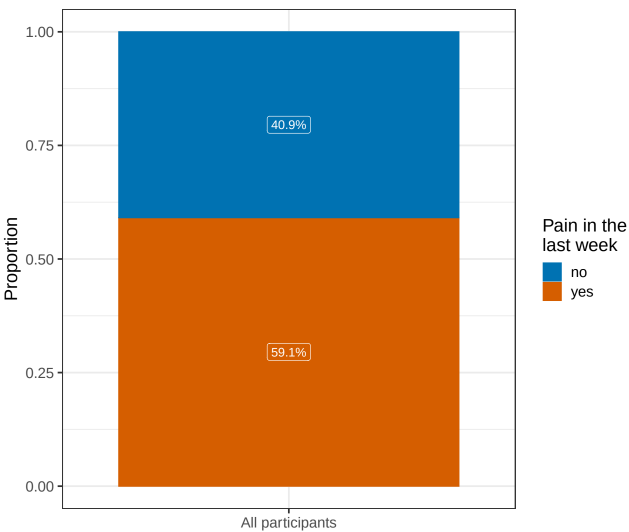
# Count plot
ggplot(data = plot_total) +
    aes(x = 'All participants',
    y = count,
```

Pain prevalence: counts



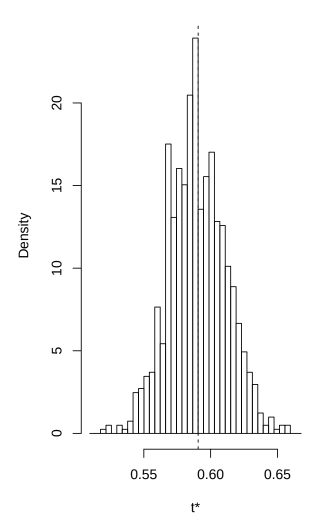
```
# Proportion plot
ggplot(data = plot_total) +
   aes(x = 'All participants',
        y = prop,
```

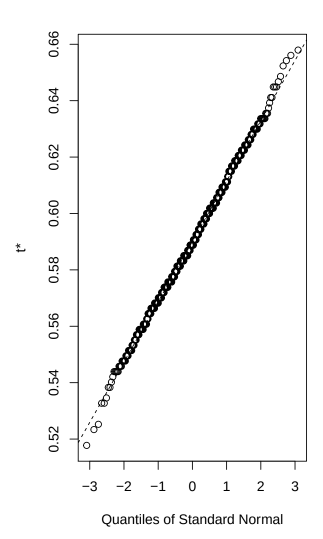
Pain prevalence: proportions



95% CI of pain frequency

```
# Define boot function
freq <- function(d, i){</pre>
    data <- d[i, ]
    mean(data$pain == 'yes')
}
# Generate bootstraps
set.seed(2019)
(boot_all <- boot(data = data,</pre>
                  statistic = freq,
                  R = 999,
                  stype = 'i'))
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = data, statistic = freq, R = 999, stype = "i")
##
##
## Bootstrap Statistics :
        original
                                 std. error
                    bias
## t1* 0.5906542 -0.0005556959 0.02138734
# Plot the distribution
plot(boot_all)
```





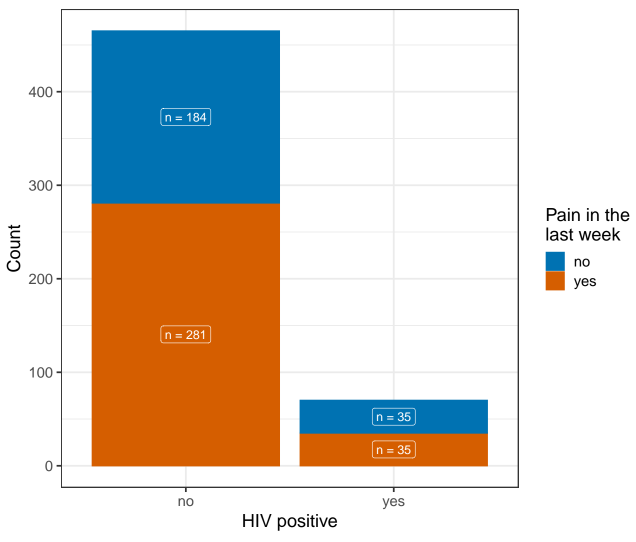
 $\mbox{\tt \#\#}$ Calculations and Intervals on Original Scale

Frequency of pain by HIV status

Exploratory plot

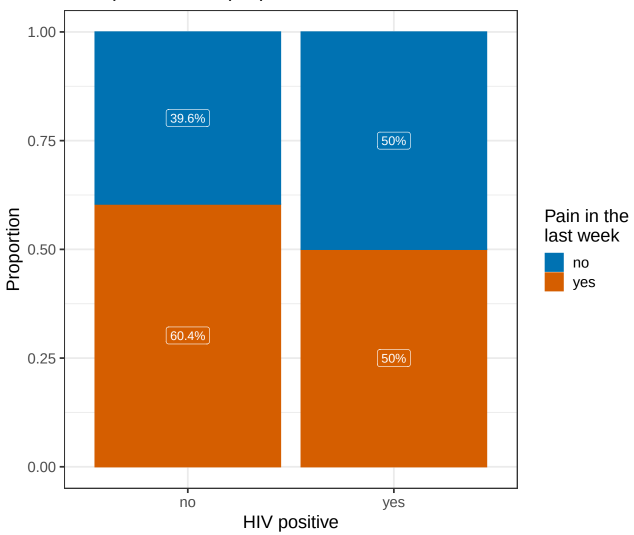
```
# Prepare data
plot_data <- data %>%
    group_by(HIV_positive, pain) %>%
    summarise(count = n()) %>%
    mutate(prop = round(count/sum(count) * 100, 1)) %>%
    ungroup()
# Count plot
ggplot(data = plot_data) +
    aes(x = HIV_positive,
       y = count,
       fill = pain,
       colour = pain) +
    geom_col() +
    geom_label(aes(label = paste0('n = ', count)),
               position = position_stack(vjust = 0.5),
               colour = 'white',
               show.legend = FALSE) +
    labs(title = 'Pain prevalence: counts',
         y = 'Count',
         x = 'HIV positive') +
    scale_fill_manual(name = 'Pain in the\nlast week',
                      values = pal) +
    scale_colour_manual(name = 'Pain in the\nlast week',
                        values = pal)
```

Pain prevalence: counts



```
# Proportion plot
ggplot(data = plot_data) +
   aes(x = HIV_positive,
       y = prop,
        fill = pain,
        colour = pain) +
   geom_col(position = position_fill()) +
   geom_label(aes(label = paste0(prop, '%')),
               position = position_fill(vjust = 0.5),
               colour = 'white',
               show.legend = FALSE) +
   labs(title = 'Pain prevalence: proportions',
         y = 'Proportion',
         x = 'HIV positive') +
   scale_fill_manual(name = 'Pain in the\nlast week',
                      values = pal) +
   scale_colour_manual(name = 'Pain in the\nlast week',
                        values = pal)
```

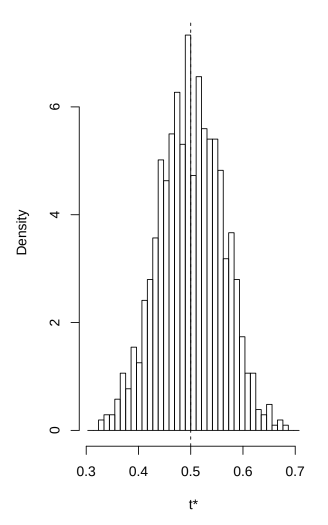
Pain prevalence: proportions

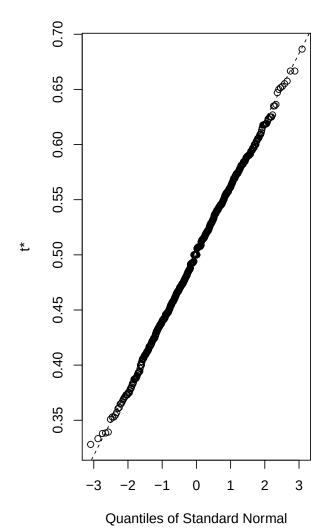


95% CI of pain frequency

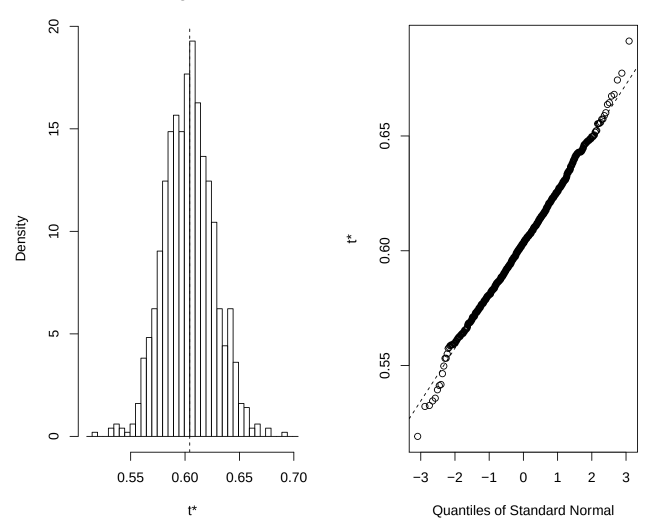
```
# Define boot function
## HIV+
freq_hiv <- function(d, i){
    data <- d[i, ]
    hiv <- data[data$HIV_positive == 'yes', ]
    pain <- mean(hiv$pain == 'yes')
    pain
}
## HIV-
freq_nohiv <- function(d, i){
    data <- d[i, ]
    hiv <- data[data$HIV_positive == 'no', ]
    pain <- mean(hiv$pain == 'yes')
    pain
}</pre>
```

```
# Generate bootstraps
## HIV+
set.seed(2019)
(boot_hiv <- boot(data = data,</pre>
                  statistic = freq_hiv,
                  R = 999,
                  stype = 'i'))
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = data, statistic = freq_hiv, R = 999, stype = "i")
##
##
## Bootstrap Statistics :
      original
                    bias std. error
         0.5 0.0003686687 0.06073863
## t1*
## HIV-
set.seed(2019)
(boot_nohiv <- boot(data = data,</pre>
                    statistic = freq_nohiv,
                    R = 999,
                    stype = 'i'))
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## boot(data = data, statistic = freq_nohiv, R = 999, stype = "i")
##
## Bootstrap Statistics :
        original
                      bias
                               std. error
## t1* 0.6043011 -0.000775804 0.02298169
# Plot the distribution
## HIV+
plot(boot_hiv)
```





HIVplot(boot_nohiv)

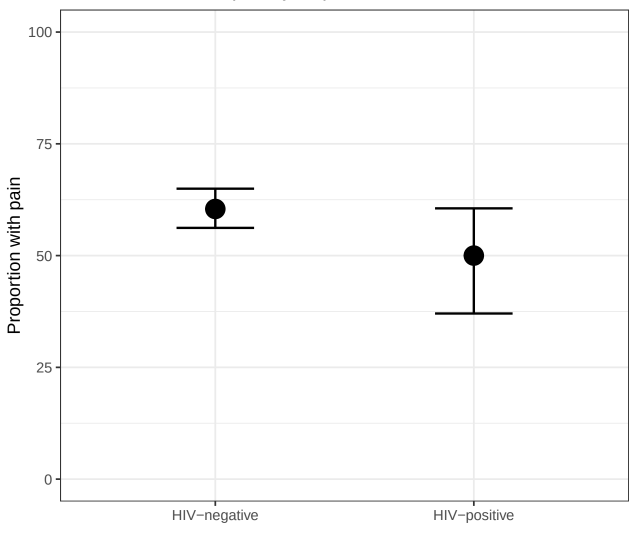


```
# Generate bias accelerated interval
## HIV+
(ci_hiv <- boot.ci(boot_hiv,</pre>
                   type = 'bca'))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 999 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot_hiv, type = "bca")
## Intervals :
               BCa
## Level
         (0.3705, 0.6056)
## Calculations and Intervals on Original Scale
## HIV-
(ci_nohiv <- boot.ci(boot_nohiv,</pre>
                      type = 'bca'))
```

BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS

```
## Based on 999 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot_nohiv, type = "bca")
## Intervals :
## Level
               BCa
## 95% ( 0.5620,  0.6497 )
## Calculations and Intervals on Original Scale
# Generate df for plotting
df_hiv <- tibble(group = c('HIV-negative',</pre>
                           'HIV-positive'),
                 t = c(100 * ci_nohiv$t0,
                       100* ci_hiv$t0),
                 lower_ci = c(100 * ci_nohiv$bca[[4]],
                              100 * ci_hiv$bca[[4]]),
                 upper_ci = c(100 * ci_nohiv$bca[[5]],
                              100* ci_hiv$bca[[5]]))
# Plot
ggplot(data = df_hiv) +
    aes(x = group,
        y = t) +
    geom_errorbar(aes(ymin = lower_ci,
                      ymax = upper_ci),
                  width = 0.3,
                  size = 1) +
    geom_point(size = 8) +
    scale_y_continuous(limits = c(0, 100)) +
    labs(title = '95% CI for the frequency of pain',
         y = 'Proportion with pain') +
    theme(axis.title.x = element_blank())
```

95% CI for the frequency of pain

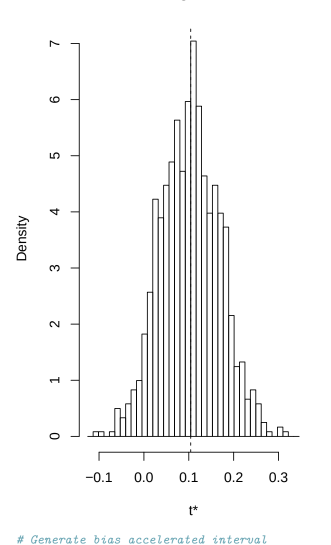


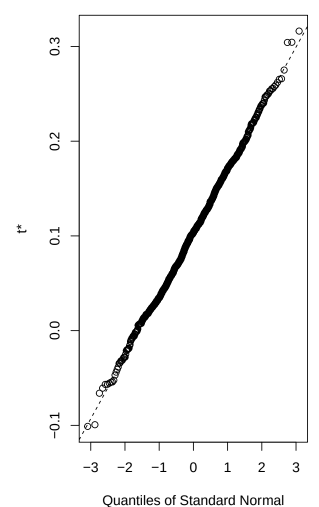
```
# Publication plot
pp <- ggplot(data = df_hiv) +</pre>
    aes(x = group,
        y = t) +
    geom_errorbar(aes(ymin = lower_ci,
                      ymax = upper_ci),
                  width = 0.3,
                  size = 1) +
    geom_point(size = 8) +
    scale_y_continuous(limits = c(0, 100)) +
    scale_x_discrete(labels = c('Negative', 'Positive')) +
    labs(y = 'Frequency of pain (%)',
         x = 'HIV test result') +
    theme_bw(base_size = 20) +
    theme(panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'))
ggsave(filename = 'figures/figure1.png',
       plot = pp,
```

```
width = 7,
height = 6)
```

95% CI for the confidence interval of the difference in proportion with pain

```
# Define boot function
freq_delta <- function(d, i){</pre>
    data <- d[i, ]
    hiv <- data[data$HIV_positive == 'yes', ]</pre>
    hivpain <- mean(hiv$pain == 'yes')</pre>
    nohiv <- data[data$HIV_positive == 'no', ]</pre>
    nohivpain <- mean(nohiv$pain == 'yes')</pre>
    nohivpain - hivpain
}
# Generate bootstraps
set.seed(2019)
(boot_delta <- boot(data = data,</pre>
                     statistic = freq_delta,
                     R = 999,
                     stype = 'i'))
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = data, statistic = freq_delta, R = 999, stype = "i")
##
## Bootstrap Statistics :
        original bias std. error
## t1* 0.1043011 -0.001144473 0.06540082
# Plot the distribution
plot(boot_delta)
```



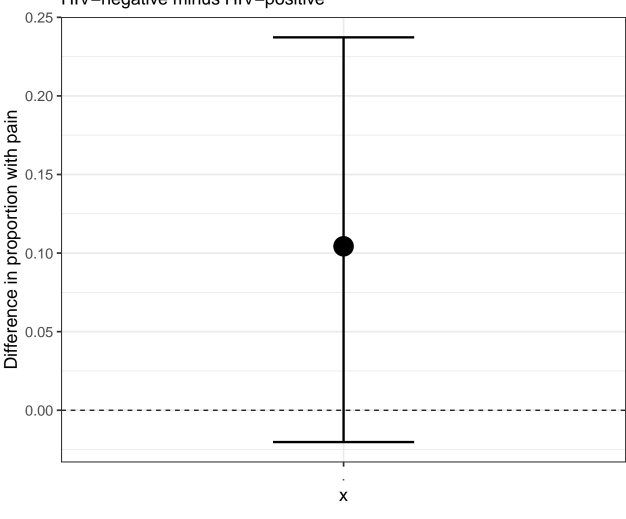


```
(ci_delta <- boot.ci(boot_delta,</pre>
                      type = 'bca'))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 999 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot_delta, type = "bca")
##
## Intervals :
## Level
                BCa
## 95%
         (-0.0202, 0.2373)
\mbox{\tt \#\#} Calculations and Intervals on Original Scale
# Generate df for plotting
df_delta <- tibble(t = ci_delta$t0,</pre>
                    lower_ci = ci_delta$bca[[4]],
```

Plot

upper_ci = ci_delta\$bca[[5]])

95% CI for the difference in proportion with pain HIV-negative minus HIV-positive



Session information

```
sessionInfo()
## R version 3.6.0 (2019-04-26)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Mojave 10.14.6
##
## Matrix products: default
          /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.6/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] car_3.0-3
                           carData_3.0-2
                                               sjPlot_2.7.0
  [4] DataExplorer_0.8.0 boot_1.3-23
                                               skimr_1.0.7
## [7] forcats_0.4.0
                           stringr_1.4.0
                                               dplyr_0.8.3
## [10] purrr_0.3.2
                           readr_1.3.1
                                               tidyr_0.8.99.9000
## [13] tibble_2.1.3
                           ggplot2_3.2.1
                                               tidyverse_1.2.1
## [16] magrittr_1.5
##
## loaded via a namespace (and not attached):
## [1] nlme_3.1-141
                          lubridate_1.7.4
                                             insight_0.4.1
   [4] httr_1.4.1
                          tools_3.6.0
                                             TMB 1.7.15
## [7] backports_1.1.4
                          utf8_1.1.4
                                            R6_2.4.0
## [10] sjlabelled 1.1.0
                          lazyeval 0.2.2
                                             colorspace 1.4-1
## [13] withr 2.1.2.9000
                          tidyselect 0.2.5
                                            gridExtra 2.3
                          emmeans_1.4
## [16] mnormt_1.5-5
                                             curl_4.0
## [19] compiler_3.6.0
                          performance_0.3.0 cli_1.1.0
## [22] rvest_0.3.4
                          xm12_1.2.2
                                             sandwich_2.5-1
## [25] labeling_0.3
                          bayestestR_0.2.5
                                            scales_1.0.0
## [28] mvtnorm_1.0-11
                          psych_1.8.12
                                             digest_0.6.20
                                             rmarkdown_1.14
## [31] foreign_0.8-72
                          minqa_1.2.4
## [34] rio_0.5.16
                          pkgconfig_2.0.2
                                             htmltools_0.3.6
## [37] lme4_1.1-21
                          highr_0.8
                                             htmlwidgets_1.3
## [40] rlang_0.4.0
                          readxl_1.3.1
                                             rstudioapi_0.10
## [43] generics_0.0.2
                          zoo_1.8-6
                                             jsonlite_1.6
                          Matrix_1.2-17
                                             fansi_0.4.0
## [46] zip_2.0.3
## [49] Rcpp 1.0.2
                          munsell_0.5.0
                                             abind 1.4-5
## [52] lifecycle_0.1.0
                          stringi_1.4.3
                                             multcomp_1.4-10
## [55] yaml_2.2.0
                          MASS_7.3-51.4
                                             grid_3.6.0
## [58] parallel_3.6.0
                                             sjmisc_2.8.1
                          ggrepel_0.8.1
## [61] crayon_1.3.4
                          lattice_0.20-38
                                             ggeffects_0.11.0
## [64] haven_2.1.1
                          splines_3.6.0
                                             sjstats_0.17.5
## [67] hms_0.5.0
                          zeallot_0.1.0
                                             knitr_1.24
## [70] pillar_1.4.2
                          igraph_1.2.4.1
                                             estimability_1.3
                                             evaluate_0.14
## [73] codetools_0.2-16
                          glue_1.3.1
```

networkD3_0.4

vctrs_0.2.0

cellranger_1.1.0

[76] data.table_1.12.2 modelr_0.1.5

[79] nloptr_1.2.1

```
## [82] gtable_0.3.0 assertthat_0.2.1 openxlsx_4.1.0.1
## [85] xfun_0.8 xtable_1.8-4 broom_0.5.2
## [88] coda_0.19-3 survival_2.44-1.1 glmmTMB_0.2.3
## [91] TH.data_1.0-10
```