Supplement 2

Baseline characteristics of RCTs for neuropathic pain

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1 Pain intensity inclusion thresholds

- Original data source: Finnerup et al 2015, Appendix 4¹.
 - All numeric values were used (after removing duplicate entries).
 - All visual analogue values were converted to a 0 to 10 scale.

¹Finnerup NB, Attal N, Haroutounian S, McNicol E, Baron R, Dworkin RH, Gilron I, Haanpää M, Hansson P, Jensen TS, Kamerman PR, Lund K, Moore A, Raja SN, Rice ASC, Rowbotham M, Sena E, Siddall P, Smith BH, Wallace M. Pharmacotherapy for neuropathic pain in adults: a systematic review and meta-analysis. Lancet Neurol 2015;14:162–173. doi:10.1016/S1474-4422(14)70251-0

```
4, 4, 4, 4, 4, 4, 4, 5, 5, 5, 5, 4, 4, 4, 3, 5, 4, 4, 4, 5, 5,
            4, 4, 5, 3, 4, 4, 4, 5, 4, 3, 3, 3, 3, 3, 3, 3, 4, 7, 5, 4, 3, 2,
            3, 4, 4, 2.5, 2.5, 4, 4, 4, 5, 4, 4, 8, 4, 4, 5, 3, 4, 4, 4)
# Generate summary statistics
## 6-number summary
summary(cut_off)
##
     Min. 1st Qu. Median
                             Mean 3rd Qu.
                                             Max.
                    4.000
##
     2.000
            4.000
                             4.015
                                     4.000
                                            8.000
## Tabular summary of inclusion thresholds
data.frame(cutoff = cut off) %>%
   group_by(cutoff) %>%
    summarise(count = n(),
             percent = round(100 * (count/length(cut_off)), 1)) %>%
   knitr::kable(., caption = 'Summary of pain inclusion thresholds')
```

Table 1: Summary of pain inclusion thresholds

cutoff	count	percent
2.0	2	1.5
2.5	2	1.5
3.0	16	12.2
4.0	93	71.0
5.0	14	10.7
6.0	2	1.5
7.0	1	0.8
8.0	1	0.8

2 Explore baseline pain intensity data

2.1 Import data

• Original data source: Finnerup et al 2015, Appendix 4¹,

```
data <- read_csv('data/baseline-data.csv')</pre>
```

2.2 Explore data

2.2.1 Median

```
Q75 = round(quantile(value, probs = 0.75, na.rm = TRUE), 1),
    max = round(max(value, na.rm = TRUE), 1)) %>%
knitr::kable(., caption = 'Summary: median values')
```

Table 2: Summary: median values

min	Q25	mean	median	Q75	max
4.7	4.9	5.8	5	5.9	8.4

2.2.2 Mean

Table 3: Summary: mean values

min	Q25	mean	median	Q75	max
3.7	5	6.2	6.4	7.2	8.7

2.2.3 SD

Table 4: Range of SD values

\min	Q25	mean	median	Q75	max
0.8	1.3	1.6	1.7	1.8	2.9

3 Weighted medians

- Three studies, four median values.
- Hahn et al., 2004 (active), Hahn et al., 2004 (placebo), Yuen et al., 2002, Low et al., 1995

4 Pooled mean

• Formula sourced from: https://www.ncbi.nlm.nih.gov/books/NBK56512/

```
# Make bootstrap function
pooled_mean <- function(d, i){</pre>
    # Select data
    data <- d[i, ]
    # Process data
    mean <- data %>%
      # Get the required data
      filter(included_in_analysis == 'yes') %>%
      filter(mean_or_median == 'mean')
    # Calculate preliminary values
    mean \leftarrow mean \%>\%
      mutate(mean_by_n = value * sample_size)
    # Calculate pooled mean
    sum(mean$mean_by_n) / sum(mean$sample_size)
}
# Perform bootstrap
set.seed(2019)
mean_pooled <- boot(data = data,</pre>
                     statistic = pooled_mean,
                     R = 999,
                     stype = 'i',
                     parallel = 'multicore',
                     ncpus = 4)
```

```
# Pooled mean
mean_pooled$t0
## [1] 6.243982
# Confidence interval of the pooled mean (percentile method)
boot.ci(mean_pooled, type = 'perc')
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 999 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = mean_pooled, type = "perc")
##
## Intervals :
            Percentile
## Level
## 95%
         (5.735, 6.652)
## Calculations and Intervals on Original Scale
```

5 Pooled SD

• Formula sourced from: https://www.ncbi.nlm.nih.gov/books/NBK56512/

```
# Make bootstrap function
pooled_sd <- function(d, i){</pre>
    # Select data
    data <- d[i, ]</pre>
    # Process the data
    sd <- data %>%
      filter(included_in_analysis == 'yes') %>%
      filter(mean_or_median == 'mean') %>%
      filter(!is.na(sd)) %>%
      # Square the SD
      mutate(SD_squared = sd^2) %>%
      # Calculate sample size -1
      mutate(n_minus_1 = sample_size - 1) %>%
      # Get the number of groups
      mutate(k = length(unique(.$authors))) %>%
      # Calculate (n-1)SD^2
      mutate(numerator = SD_squared * n_minus_1)
    # Pooled SD
    sqrt(sum(sd$numerator) / (sum(sd$sample_size) - sd$k[[1]]))
}
# Perform bootstrap
set.seed(2019)
sd_pooled <- boot(data = data,</pre>
                    statistic = pooled_sd,
                    R = 999,
```

```
stype = 'i',
                    parallel = 'multicore',
                    ncpus = 4)
# Pooled mean
sd_pooled$t0
## [1] 1.665597
# Confidence interval of the pooled mean (percentile method)
boot.ci(sd_pooled, type = 'perc')
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 999 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = sd_pooled, type = "perc")
##
## Intervals :
## Level
            Percentile
## 95%
         (1.362, 1.982)
## Calculations and Intervals on Original Scale
```

6 Mean placebo reponse

6.1 DPN/PHN

• Freeman R, Emir B, Parsons B. Predictors of placebo response in peripheral neuropathic pain: insights from pregabalin clinical trials. *J Pain Res* 8:257–268, 2015. DOI:10.2147/JPR.S78303.

6.2 chronic OA/LBP

• Vase L, Vollert J, Finnerup NB, Miao X, Atkinson G, Marshall S, Nemeth R, Lange B, Liss C, Price DD, Maier C, Jensen TS, Segerdahl M. Predictors of the placebo analgesia response in randomized controlled trials of chronic pain: a meta-analysis of the individual data from nine industrially sponsored trials. *Pain* 156:1795–1802, 2015. DOI:10.1097/j.pain.0000000000000217.

```
# Taken from Figure 1
scale_placebo = 5/42
vase_placebo <- c(89, 146, 62, 97.5, 48, 52.5, 131, 155.5, 141.5) + 42
vase_placebo <- round(scale_placebo * vase_placebo)</pre>
vase_placebo <- vase_placebo / 10 # Conversion from 100mm VAS to 10 unit NRS
scale_size = 50/15
vase_size <- c(35, 37, 37, 65, 65, 72.5, 95, 100, 100)</pre>
vase_size <- round(vase_size * scale_size)</pre>
length(vase_placebo)
## [1] 9
df_placebo2 <- data.frame(value = vase_placebo,</pre>
                           size = vase_size) %>%
 mutate(value_size = value * size)
# Weighted mean
sum(df_placebo2$value_size) / sum(df_placebo2$size)
## [1] 1.818348
```

7 Session information

```
sessionInfo()
## R version 4.0.4 (2021-02-15)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Catalina 10.15.7
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.0/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:
                graphics grDevices utils
## [1] stats
                                              datasets methods
##
## other attached packages:
## [1] skimr_2.1.3
                          boot_1.3-27
                                             matrixStats_0.58.0 forcats_0.5.1
## [5] stringr_1.4.0
                          dplyr_1.0.5
                                             purrr_0.3.4
                                                                readr_1.4.0
## [9] tidyr_1.1.3
                          tibble_3.1.0
                                             ggplot2_3.3.3
                                                                tidyverse_1.3.0
```

```
##
## loaded via a namespace (and not attached):
   [1] tidyselect_1.1.0 xfun_0.22
                                             repr_1.1.3
                                                               haven_2.3.1
   [5] colorspace_2.0-0
                          vctrs_0.3.6
                                                               htmltools_0.5.1.1
                                             generics_0.1.0
   [9] base64enc_0.1-3
                          yaml_2.2.1
##
                                             utf8_1.2.1
                                                               rlang_0.4.10
## [13] pillar_1.5.1
                          glue_1.4.2
                                             withr_2.4.1
                                                               DBI_1.1.1
## [17] dbplyr_2.1.0
                          modelr_0.1.8
                                             readxl_1.3.1
                                                               lifecycle_1.0.0
## [21] munsell_0.5.0
                                                               rvest_1.0.0
                          gtable_0.3.0
                                             cellranger_1.1.0
## [25] evaluate_0.14
                          knitr_1.31
                                             parallel_4.0.4
                                                               fansi_0.4.2
## [29] highr_0.8
                          broom_0.7.5
                                             Rcpp_1.0.6
                                                               scales_1.1.1
## [33] backports_1.2.1
                          jsonlite_1.7.2
                                             fs_1.5.0
                                                               hms_1.0.0
## [37] digest_0.6.27
                          stringi_1.5.3
                                             grid_4.0.4
                                                               cli_2.3.1
## [41] tools_4.0.4
                          magrittr_2.0.1
                                             crayon_1.4.1
                                                               pkgconfig_2.0.3
## [45] ellipsis_0.3.1
                          xm12_1.3.2
                                             reprex_1.0.0
                                                               lubridate_1.7.10
## [49] assertthat_0.2.1
                          rmarkdown_2.7
                                             httr_1.4.2
                                                               rstudioapi_0.13
## [53] R6_2.5.0
                          compiler_4.0.4
```