# Supplementary code to reproduce the numerical results in Di Caterina and Kosmidis (2019)

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## Workspace preparation

This document provides R (R Core Team, 2017) code to reproduce the results in the manuscript 'Location-adjusted Wald statistics for scalar parameters' by Di Caterina and Kosmidis (2019).

This script assumes that the current working directory has the sub-directories code, results and lesion data as provided in the supplementary material.

```
path <- "."
code_path <- paste(path, "code", sep = "/")
results_path <- paste(path, "results", sep = "/")
lesions_path <- paste(path, "lesion data", sep = "/")</pre>
```

The contents of the directories are as follows

```
dir(code path)
# [1] "babies_simulation.R"
                                   "babies timings.R"
# [3] "brains_case_study.R"
                                   "clotting\_simulation.R"
# [5] "dyslexia_simulation.R"
                                   "loqodds_functions.R"
# [7] "meta_analysis_simulation.R" "overlay2_nifti.R"
dir(results_path)
# [1] "babies_simulation.rda"
                                     "babies_times.rda"
# [3] "brains_case_study.rda"
                                     "clotting_simulation.rda"
# [5] "dyslexia_simulation.rda"
                                     "meta_analysis_simulation.rda"
dir(lesions_path)
# [1] "data_demo.dat" "images"
```

First, make sure that you have the latest version of the waldi R package installed.

```
waldi_version <- try(packageVersion("waldi"), silent = TRUE)
if (inherits(waldi_version, "try-error")) {
    devtools::install_github("ikosmidis/waldi")
}</pre>
```

The following code chunk loads the required packages

```
library("waldi")
library("oro.nifti")
library("boot")
library("plyr")
library("plotrix")
library("dplyr")
library("survival")
library("cond")
library("lmtest")
library("betareg")
library("enrichwith")
```

```
library("brglm2")
library("ggplot2")
library("gridExtra")
library("colorspace")
```

## Pre-saved R image files

Some of the code-chunks below load objects from the pre-saved R image files in the results directory. These image files are the outputs of the scripts babies\_simulation.R, babies\_timings.R, brains\_case\_study.R, clotting\_simulation.R, dyslexia\_simulation.R, meta\_analysis\_simulation.R.

#### Table 1

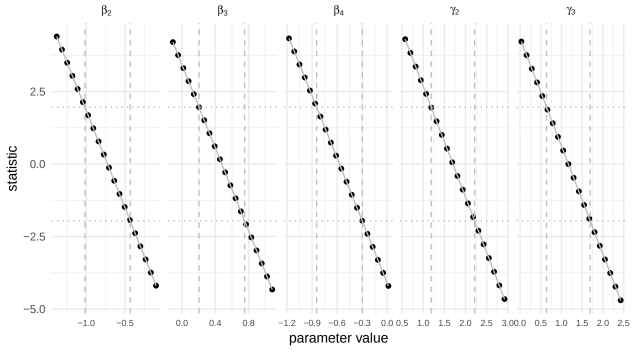
```
data("ReadingSkills", package = "betareg")
## maximum likelihood estimates and corresponding 95\% Wald confidence intervals
rs_beta_ml <- betareg(accuracy ~ dyslexia * iq | dyslexia + iq,
                     data = ReadingSkills, type = "ML", hessian = FALSE)
rs_summary_ml <- coef(summary(rs_beta_ml))</pre>
rs_ml_estimates <- do.call("rbind", lapply(rs_summary_ml,</pre>
                                         function(z) z[, c("Estimate", "Std. Error")]))
rs_ml_cis <- confint(rs_beta_ml)
## bias corrected fit and corresponding 95\% Wald confidence intervals
rs beta br <- update(rs beta ml, type = "BR")
rs_summary_br <- coef(summary(rs_beta_br))</pre>
rs_br_estimates <- do.call("rbind", lapply(rs_summary_br,</pre>
                                         function(z) z[, c("Estimate", "Std. Error")]))
rs_br_cis <- confint(rs_beta_br)</pre>
round(cbind(rs_ml_estimates, rs_br_estimates, rs_ml_cis, rs_br_cis), 3)
             Estimate Std. Error Estimate Std. Error 2.5 % 97.5 % 2.5 % 97.5 %
# (Intercept)
                1.123
                           0.143 1.114
                                              -0.742
                           0.143
                                              0.148 -1.021 -0.462 -1.024 -0.444
# dyslexia
                                  -0.734
                                              0.141 0.225 0.747 0.165 0.717
                           0.133
                                 0.441
#iq
                0.486
              -0.581
                           0.133
                                              0.140 -0.841 -0.321 -0.807 -0.257
# dyslexia:iq
                                  -0.532
# (Intercept)
             3.304
                           0.223 3.092
                                              0.225 2.868 3.741 2.652 3.533
# dyslexia
                1.747
                           0.262
                                   1.654
                                              0.264 1.232 2.261 1.138 2.171
                                              0.271 0.705 1.753 0.518 1.578
# iq
                1.229
                           0.267
                                 1.048
```

#### Table 2

dyslexia\_simulation.rda contains the outputs of dyslexia\_simulation.R in ./code, which replicates the simulation study described in Example 1.1 of Di Caterina and Kosmidis (2019)

```
"ml_cor_ses_cor", "br_cor_ses_cor")) %>%
   mutate(parameter = recode(parameter,
                             "dyslexia" = 2, "iq" = 3, "dyslexia:iq" = 4,
                             "(phi)_dyslexia" = 6, "(phi)_iq" = 7)) %>%
   mutate(level = 100 * level) %>%
   group_by(level, statistic, parameter) %>%
   summarize(coverage = round(mean(cover, na.rm = TRUE) * 100, 1)) %>%
   as.data.frame() %>%
   reshape(idvar = c("statistic", "parameter"), v.names = "coverage",
           timevar = "level",
           direction = "wide")
# Warning: package 'bindrcpp' was built under R version 3.4.4
rs_coverage %>% filter(statistic %in% c("ml", "br")) %>%
   select(statistic, parameter, coverage.90, coverage.95, coverage.99)
    statistic parameter coverage.90 coverage.95 coverage.99
# 1
           br
                      2
                              88.1
                                          93.4
# 2
           br
                      3
                               87.2
                                           92.9
                                                      98.0
# 3
                              87.3
                                          92.9
                                                      98.0
           br
                      4
                              83.8
                                          90.2
# 4
           br
                      6
                                                      96.7
                      7
# 5
                              82.7
                                          89.2
                                                      96.1
           br
# 6
           ml
                      2
                              86.9
                                         92.4
                                                      97.7
# 7
                      3
                                         91.0
           ml
                             84.8
                                                      97.1
# 8
           ml
                      4
                              85.0
                                          91.2
                                                     97.2
# 9
                                          89.1
                                                      96.1
                      6
                               82.4
           m1.
# 10
                               79.1
                                          86.0
                                                      94.4
```

```
rs_cor_ml_cis <- waldi_confint(rs_beta_ml, level = 0.95, adjust = TRUE)
interpolation <- waldi_confint(rs_beta_ml, level = 0.95,</pre>
                                which = rownames(rs cor ml cis),
                                adjust = TRUE,
                               return values = TRUE,
                               length = 20)
intervals <- data.frame(low = rs_cor_ml_cis[, 1],</pre>
                        upp = rs_cor_ml_cis[, 2],
                        parameter = rownames(rs_cor_ml_cis))
interpolation <- interpolation %>%
    filter(!(parameter %in% c("(Intercept)", "(phi)_(Intercept)"))) %>%
    mutate(parameter = recode(parameter,
                               "dyslexia" = "beta[2]",
                               "iq" = "beta[3]",
                               "dyslexia:iq" = "beta[4]",
                               "(phi)_dyslexia" = "gamma[2]",
                               "(phi)_iq" = "gamma[3]"))
intervals <- intervals %>%
    filter(!(parameter %in% c("(Intercept)", "(phi)_(Intercept)"))) %>%
    mutate(parameter = recode(parameter,
                               "dyslexia" = "beta[2]",
                               "iq" = "beta[3]",
                               "dyslexia:iq" = "beta[4]",
```



#### Table 3

```
rs_cor_br_stud_cis <- waldi_confint(rs_beta_br, adjust = TRUE, parallel = FALSE,
                     quantiles = quantiles_br$zstat_cor[, c("0.025", "0.975")])
round(rbind(cbind(rs_cor_ml_cis, rs_cor_br_cis),
           cbind(rs_cor_ml_stud_cis, rs_cor_br_stud_cis)), 3)
                    2.5 % 97.5 % 2.5 % 97.5 %
# (Intercept)
                    0.816 1.400 0.827 1.411
# dyslexia
                   -1.019 -0.435 -1.031 -0.446
                    0.204 0.752 0.165 0.719
#iq
                   -0.845 -0.299 -0.809 -0.257
# dyslexia:iq
# (phi)_(Intercept) 2.689 3.564 2.652 3.532
# (phi)_dyslexia
                   1.186 2.214 1.134 2.169
                    0.639 1.691 0.513 1.574
# (phi)_iq
# (Intercept)
                   0.812 1.420 0.830 1.481
# dyslexia
                   -1.059 -0.442 -1.091 -0.440
#iq
                    0.171 0.792 0.159 0.758
# dyslexia:iq
                   -0.871 -0.268 -0.853 -0.264
# (phi)_(Intercept) 2.709 3.680 2.654 3.682
# (phi)_dyslexia
                    1.112 2.303 1.040 2.241
                    0.565 1.835 0.394 1.769
# (phi)_iq
rs_coverage %>% filter(statistic %in% c("ml_cor", "br_cor")) %>%
   select(statistic, parameter, coverage.90, coverage.95, coverage.99)
    statistic parameter coverage.90 coverage.95 coverage.99
# 1
       br_cor
                      2
                               88.3
                                           93.5
                                                       98.3
# 2
       br cor
                      3
                               87.3
                                           93.0
                                                       98.0
# 3
       br cor
                      4
                               87.5
                                           93.0
                                                       98.0
                                           90.3
# 4
       br cor
                      6
                               83.9
                                                      96.8
                      7
# 5
       br\_cor
                               82.7
                                           89.2
                                                      96.2
                      2
# 6
       ml\_cor
                               88.5
                                           93.7
                                                      98.4
                      3
# 7
                                           92.8
                                                       98.0
       ml\_cor
                               87.1
                                                       98.0
# 8
       ml\_cor
                      4
                               87.2
                                           92.8
# 9
                               83.5
                                           90.0
                                                       96.6
       ml\_cor
                      6
                      7
# 10
       ml\_cor
                               81.8
                                           88.6
                                                       95.7
rs_coverage %% filter(statistic %in% c("ml_cor_stud", "br_cor_stud")) %>%
   select(statistic, parameter, coverage.90, coverage.95, coverage.99)
       statistic parameter coverage.90 coverage.95 coverage.99
# 1 br_cor_stud
                        2
                                 89.4
                                             94.6
                                                         98.6
# 2 br cor stud
                        3
                                 89.4
                                             94.5
                                                         98.5
# 3 br_cor_stud
                                                         98.6
                        4
                                 89.5
                                             94.4
# 4 br cor stud
                        6
                                 90.1
                                             94.9
                                                         98.8
                        7
# 5 br_cor_stud
                                 90.5
                                             95.1
                                                         98.8
                        2
# 6 ml_cor_stud
                                 89.5
                                             94.5
                                                         98.7
#7 ml cor stud
                        3
                                 89.2
                                             94.3
                                                         98.5
                        4
# 8 ml_cor_stud
                                 89.3
                                             94.3
                                                         98.5
# 9 ml_cor_stud
                        6
                                 89.9
                                                         98.7
                                             94.7
                        7
# 10 ml_cor_stud
                                 90.1
                                             94.9
                                                         98.7
```

The following chunk of code reproduces the times for the computation of the confidence intervals reported in Section 6.

```
## Intervals based on the location-adjusted Wald statistic
system.time({
    waldi_confint(rs_beta_ml, adjust = TRUE, parallel = FALSE, length = 5)
})
```

```
# user system elapsed
# 2.420 0.051 2.486
simu_fun <- get_simulate_function(rs_beta_ml)</pre>
generate_dyslexia <- function(data, mle) {</pre>
    simu fun(mle)
}
stat <- function(data, psi) {</pre>
    temp <- ReadingSkills</pre>
    temp$accuracy <- data</pre>
    temp_fit <- try(update(rs_beta_ml, data = temp))</pre>
    if (inherits(temp_fit, "try-error")) {
        rep(NA, 7)
    }
    else {
        waldi(temp_fit, null = psi, adjust = TRUE)
    }
## Studentized bootstrap intervals
system.time({
    stats <- boot(ReadingSkills$accuracy, statistic = stat,</pre>
                   R = 500, sim = "parametric", ran.gen = generate_dyslexia,
                   mle = coef(rs_beta_ml), psi = coef(rs_beta_ml), ncpus = 1)$t
    quant <- t(apply(stats, 2, quantile, probs = c(0.025, 0.975), na.rm = TRUE))
    waldi_confint(rs_beta_br, adjust = TRUE, parallel = FALSE,
                   quantiles = quant)
})
     user system elapsed
# 248.818 4.227 258.126
```

```
source(pasteO(code_path, "/", "logodds_functions.R"))
## Distribution of the statistic against normal
settings <- expand.grid(m = c(8, 16, 32), theta0 = c(-2, -1, 0))
plot_data <- NULL</pre>
for (j in seq.int(nrow(settings))) {
    setting <- settings[j, ]</pre>
    z \leftarrow seq(-3, 3, length = 100)
    dat <- t(sapply(z, dist_function, n = setting$m, theta0 = setting$theta0))</pre>
    dd <- stack(as.data.frame(dat))</pre>
    dd$z <- z
    names(dd) <- c("prob", "method", "z")</pre>
    dd$theta0 <- setting$theta0
    dd$m <- setting$m
    plot_data <- rbind(plot_data, dd)</pre>
plot_data$theta0 <- paste0("theta[0] == ", plot_data$theta0)</pre>
plot_data$theta0 <- factor(plot_data$theta0, levels = unique(plot_data$theta0),</pre>
                             ordered = TRUE)
plot_data$m <- paste0("n == ", plot_data$m)</pre>
plot_data$m <- factor(plot_data$m, levels = unique(plot_data$m), ordered = TRUE)</pre>
```

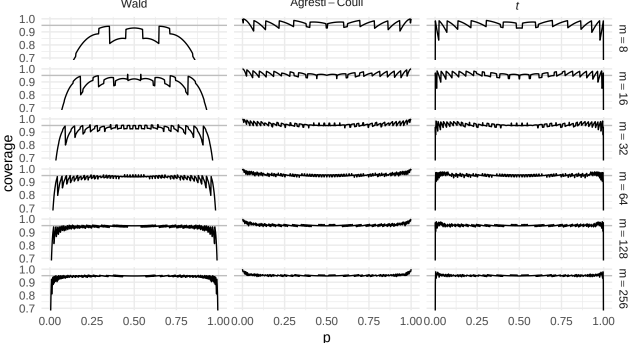
```
plot_data$method <- factor(plot_data$method, levels = c("ml", "a_ml", "br", "a_br"),</pre>
                                ordered = TRUE)
plot_data$method <- recode(plot_data$method,</pre>
                                "ml" = "italic(t)",
                                "a_ml" = "italic(t)^{\'*'}",
                                "br" = "tilde(italic(t))",
                                a_br = tilde(italic(t))^{'*'}
ggplot(plot_data) +
     geom_abline(aes(intercept = 0, slope = 0), col = "grey") +
    geom_line(aes(z, qnorm(prob) - z), alpha = 0.5) +
    facet_grid(method ~ theta0 + m, label = label_parsed) +
    theme minimal() +
    labs(y = expression(paste(Phi^list(-1),(italic(G)(italic(z))) - italic(z))),
          x = expression(italic(z))) +
    theme(text=element_text(size = 11))
          \theta_0\!=\!-2
                     \theta_0 = -2
                                \theta_0 = -2
                                                                                      \theta_0 = 0
                                                                                                 \theta_0 = 0
                                                     \theta_0 = -1
                                                                \theta_0 = -1
                                                                           \theta_0 = 0
                      n = 16
                                n = 32
                                                      n = 16
                                                                n = 32
                                                                            n = 8
                                                                                      n = 16
                                                                                                n = 32
     1.0
     0.5
                                                                 MWWWMA
     0.0
    -0.5
    -1.0
     1.0
     0.5
\Phi^{-1}(G(z))-z
     0.0
    -0.5
    -1.0
     1.0
     0.5
     0.0
    -0.5
    -1.0
     1.0
     0.5
     0.0
    -0.5
    -1.0
                     -2 0 2
                                                       0
                                                               -2 0
```

## Coverage and length of confidence intervals for a binomial proportion

This section provides evidence for the stated coverage and expected length properties of confidence intervals for a binomial proportion in Section 8 of the main text. The code chunk below computes and visualises the coverage and expected length of the 95% confindence intervals  $\bar{y} \pm z_{0.975} \sqrt{\bar{y}(1-\bar{y})/n}$  (Wald),  $\tilde{p} \pm z_{0.975} \sqrt{\tilde{p}(1-\tilde{p})/(n+4)}$ , where  $\tilde{p} = (\sum y_i + 2)/(n+4)$  (Agresti-Coull; Agresti and Coull, 1998; Agresti and Caffo, 2000), and the intervals based on the transformation of the endpoints of the confidence intervals for the log-odds based on  $\tilde{t}^*$ .

```
probs <- seq(1e-08, 1 - 1e-08, length = 500)
df <- ddply(data.frame(m = c(8, 16, 32, 64, 128, 256)), ~ m, function(x) {
    m <- x$m</pre>
```

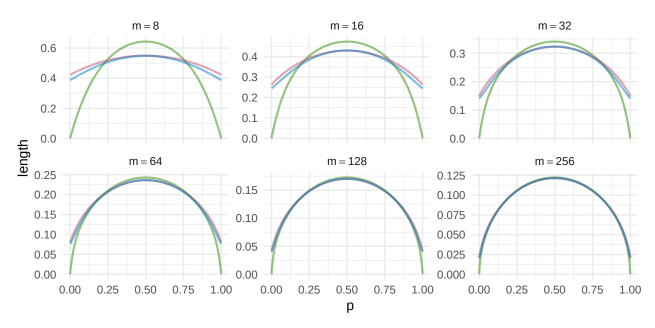
```
cis <- compute_cis(m, level = 0.95)</pre>
    cc <- lapply(probs, function(pp) cover_ci_prop(n = m, p = pp, level = 0.95, cis = cis))
    do.call("rbind", cc)
})
df$m <- factor(paste("m ==", df$m), levels = paste("m ==", sort(unique(df$m))),
               ordered = TRUE)
df$method <- factor(df$method, levels = c("wald", "ac", "a_br", "ml", "a_ml", "br"),</pre>
                     ordered = TRUE)
df$method <- recode(df$method,
                     "wald" = "Wald",
                     "ml" = "italic(t)[trans]",
                     a_ml = "italic(t)^*[trans]",
                     "br" = "tilde(italic(t))[trans]",
                     "a br" = "tilde(italic(t))^{*}",
                     "ac" = "Agresti-Coull")
## coverage
ggplot(df %>% filter(method %in% c("Wald", "Agresti-Coull", "tilde(italic(t))^{'*'}"))) +
    geom_hline(aes(yintercept = 0.95), col = "grey") +
    geom\_line(aes(x = p, y = coverage)) +
    facet_grid(m ~ method, label = label_parsed) +
    coord_cartesian(ylim = c(0.7, 1)) +
    theme_minimal()
                                           Agresti - Coull
                 Wald
  1.0
  0.9
  8.0
  0.7
  1.0
                                                                                            3
  0.9
                                                                                            Ш
  0.8
                                                                                            16
  0.7
```



```
bquote(tilde(italic(t))^{'*'}))) +
theme_minimal() +
theme(legend.position = "top")

— Wald — Agresti-Coull — t

t
```



## Hauck and Donner effect

```
sapply(28:32, t_ml, n = 32, theta0 = 0)
# [1] 3.640465 3.740749 3.708150 3.379905 0.000000
sapply(28:32, t_adjusted_ml, n = 32, theta0 = 0)
# [1] 3.770481 3.912737 3.955360 3.816022 0.0000000
sapply(28:32, t_br, n = 32, theta0 = 0)
# [1] 3.583279 3.712935 3.744298 3.587411 2.884566
sapply(28:32, t_adjusted_br, n = 32, theta0 = 0)
# [1] 3.763721 3.902155 3.935838 3.762302 2.921237
```

#### Table 4

```
## The clotting data set
clotting <- data.frame(
   conc = c(118,58,42,35,27,25,21,19,18,69,35,26,21,18,16,13,12,12),
   u = c(5,10,15,20,30,40,60,80,100, 5,10,15,20,30,40,60,80,100),
   lot = factor(c(rep(1, 9), rep(2, 9))))
## The maximum likelihood fit of the gamma regression model
clotting_ml <- glm(conc ~ log(u)*lot, data = clotting, family = Gamma(link = "log"))
## Maximum likelihood estimates and Wald statistics using maximum likelihood estimator
## of the dispersion parameter
dispersion_ml <- MASS::gamma.dispersion(clotting_ml)</pre>
```

```
clotting_summary_ml <- summary(clotting_ml, dispersion = dispersion_ml)</pre>
clotting_ml_estimates <- coef(clotting_summary_ml)[, c("Estimate", "z value")]</pre>
## Reduced-bias estimates and Wald statistics
clotting_summary_rb <- summary(update(clotting_ml, method = "brglmFit"))</pre>
## Maximum likelihood estimates and Wald statistics using the moment-based estimator
## of the dispersion parameter
clotting_summary_mom <- summary(clotting_ml)</pre>
dispersion mom <- clotting summary mom$dispersion
clotting mom estimates <- coef(clotting summary mom)[, c("Estimate", "t value")]
## Location-adjusted Wald statistic
clotting_waldi <- waldi(clotting_ml, null = 0, adjust = TRUE)</pre>
round(cbind(c(clotting_ml_estimates[, 1], dispersion_ml, dispersion_mom),
            c(clotting_ml_estimates[, 2], NA, NA),
            c(clotting_mom_estimates[, 2], NA, NA),
            c(clotting_waldi, NA, NA)), 3)
#
                [,1]
                       [,2]
                                [,3]
                                         [,4]
# (Intercept) 5.503 34.124 29.282 28.953
             -0.602 -12.842 -11.020 -10.896
\# loq(u)
# lot2
              -0.584 -2.563 -2.199 -2.173
# log(u):lot2 0.034 0.520
                               0.446
                                       0.441
#
               0.017
                                  NA
                                           NA
                          NA
               0.024
                          NA
                                  NA
                                           NA
```

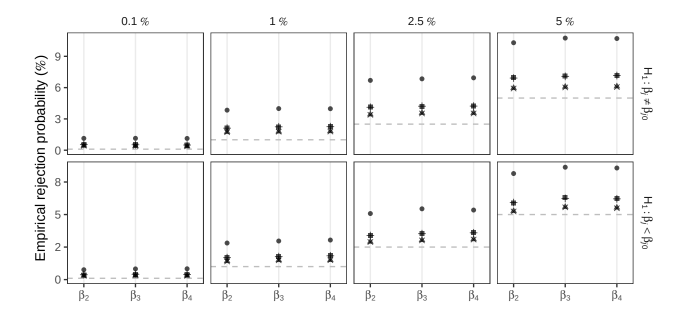
## Figure 3 including rejection probabilities based on $\tilde{t}$ and $\tilde{t}^*$

clotting\_simulation.rda below is the output of clotting\_simulation.R in ./code, which replicates the simulation study described in Section 9.3 of Di Caterina and Kosmidis (2019).

```
load(paste(results_path, "clotting_simulation.rda", sep = "/"))
## Summary of results
res statistics <- ldply(res, function(x) x$stats)
res_pvalues <- ldply(res, function(x) x$boot_pvalues)</pre>
## Type I error rates
typeI_statistics <- ddply(res_statistics, ~ name + parameter, function(x) {</pre>
  levels \langle -c(0.1, 1, 2.5, 5)/100 \rangle
  p_value_2sided <- 2 * pnorm(-abs(x$value))</pre>
  p_value_left <- pnorm(x$value)</pre>
  p_value_right <- 1 - pnorm(x$value)</pre>
  rate_2sided <- sapply(levels, function(alpha) mean(p_value_2sided < alpha))
  rate_left <- sapply(levels, function(alpha) mean(p_value_left < alpha))</pre>
  rate_right <- sapply(levels, function(alpha) mean(p_value_right < alpha))</pre>
  out <- data.frame(</pre>
    test = rep(c("2sided", "left", "right"), each = length(levels)),
    typeI = c(rate_2sided, rate_left, rate_right),
    level = rep(levels, times = 3))
  out
})
typeI_pvalues <- ddply(res_pvalues, ~ statistic + parameter, function(x) {</pre>
 levels \langle -c(0.1, 1, 2.5, 5)/100 \rangle
```

```
rate_2sided <- sapply(levels, function(alpha) mean(x$value[x$type ==
                                 'boot conv 2sided'] < alpha))</pre>
  rate_left <- sapply(levels, function(alpha) mean(x$value[x$type ==
                               'boot_conv_left'] < alpha))</pre>
  rate_right <- sapply(levels, function(alpha) mean(x$value[x$type ==</pre>
                                'boot_conv_right'] < alpha))</pre>
 out <- data.frame(</pre>
    test = rep(c("2sided", "left", "right"), each = length(levels)),
    typeI = c(rate_2sided, rate_left, rate_right),
    level = rep(levels, times = 3))
  out
})
names(typeI_statistics) <- names(typeI_pvalues)</pre>
levels(typeI_pvalues$statistic) <- c("ml_boot", "ml_cor_boot", "mom_boot")</pre>
typeI <- rbind(typeI_statistics, typeI_pvalues)</pre>
typeI <- typeI %>%
 filter(test != "right") %>%
  mutate(test = recode(test,
                       "2sided" = "H[1]: beta[italic(j)] != beta[paste(italic(j), 0)]",
                       "left" = "H[1]: beta[italic(j)] < beta[paste(italic(j), 0)]",
                       "right" = "H[1]: beta[italic(j)] > beta[paste(italic(j), 0)]"),
         level_chr = paste(level*100, "~symbol('\045')"),
         upper = typeI - qnorm(1 - 0.01/2)*sqrt(typeI*(1-typeI)/nsimu),
         lower = typeI + qnorm(1 - 0.01/2)*sqrt(typeI*(1-typeI)/nsimu))
ggplot(typeI %>% filter(parameter != 1, is.element(statistic,
                                               c("ml", "ml_cor", "mom",
                                                 "rb", "rb cor")))) +
    geom_point(aes(parameter, typeI, pch = statistic), alpha = 0.7) +
    geom_hline(aes(yintercept = level), col = "grey", lty = 2) +
    facet_grid(test ~ level_chr, labeller = label_parsed, scales = "free") +
    scale_x_continuous(name = element_blank(),
                       breaks = c(2, 3, 4),
                       limits = c(1.8, 4.2),
                       labels = c(
                           expression(beta[2]),
                           expression(beta[3]),
                           expression(beta[4]))) +
    scale_y_continuous(name = expression(paste("Empirical rejection probability (",
                                                symbol('\045'), ")")),
                       labels = function (x) {
                           if (length(x) == 0)
                                return(character())
                           x <- round_any(x, scales:::precision(x)/100)
                           scales:::comma(x * 100)
                       }) +
  scale_shape_manual(name = " ", values = c(16, 17, 15, 3, 4),
                     labels = c(expression(italic(t)), expression(italic(t)^list("*")),
            expression(paste("Wald with ", tilde(phi))), expression(tilde(italic(t))),
            expression(tilde(italic(t))^list("*")))) +
```

```
theme_bw() +
theme(legend.position = "top",
    panel.grid.major.y = element_blank(),
    panel.grid.minor.y = element_blank(),
    panel.grid.minor.x = element_blank(),
    strip.background = element_blank())
```

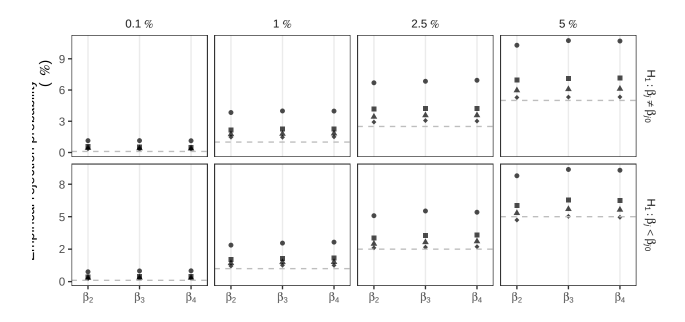


## Scale adjustment via bootstrap

The following figure shows how precise the testing accuracy of the location- and scale-adjusted statistic  $t^{**}$  obtained via parametric bootstrap is, further enhancing the performance of  $t^*$ .

```
ggplot(typeI %>% filter(parameter != 1, is.element(statistic,
                                              c("ml", "ml_cor", "mom",
                                                 "sc_ml_cor_conv")))) +
  geom_point(aes(parameter, typeI, pch = statistic), alpha = 0.7) +
  geom_hline(aes(yintercept = level), col = "grey", lty = 2) +
  facet_grid(test ~ level_chr, labeller = label_parsed, scales = "free") +
  scale_x_continuous(name = element_blank(),
                     breaks = c(2, 3, 4),
                     limits = c(1.8, 4.2),
                     labels = c(expression(beta[2]),
                       expression(beta[3]),
                       expression(beta[4]))) +
  scale_y_continuous(name = expression(paste("Empirical rejection probability
                                              (", symbol('\045'), ")")),
                     labels = function (x) {
                       if (length(x) == 0)
                         return(character())
                       x <- round_any(x, scales:::precision(x)/100)
                       scales:::comma(x * 100)
```

#### • $t t^* Wald with \widetilde{\phi} t^{**}$

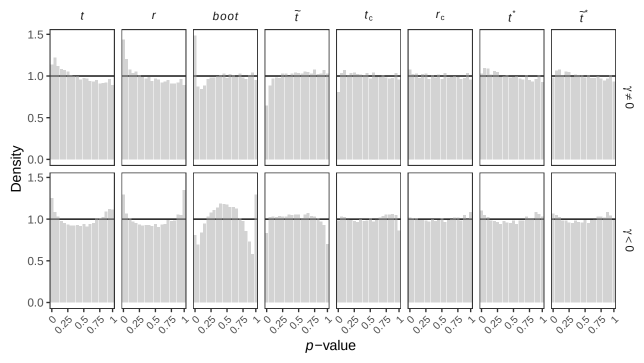


#### Table 5

```
babies_ml)
rc <- summary(babies_cond)$logtest[1]</pre>
scorec <- summary(babies_cond)$sctest[1]</pre>
out1 <- c(
   ml = unname(ml["Estimate"]),
   rb = unname(rb["Estimate"]),
   mcl = unname(mcl["coef"]),
   wald ml = unname(ml["z value"]),
   wald mcl = unname(mcl["z"]),
   wald_rb = unname(rb["z value"]),
   r = unname(sign(ml["Estimate"]) * sqrt(r$Chisq[2])),
   rc = unname(sign(mcl["coef"]) * sqrt(rc)),
   wald_ml_adjusted = unname(waldi(babies_ml, which = 19)),
   wald_rb_adjusted = unname(waldi(babies_rb, which = 19)))
out2 <- c(
   ml_se = unname(ml["Std. Error"]),
   rb_se = unname(rb["Std. Error"]),
   mcl_se = unname(mcl["se(coef)"]),
   ml_p = ml["Pr(>|z|)"],
   mcl_p = mcl["Pr(>|z|)"],
   rb_p = rb["Pr(>|z|)"],
   r_p = 2 * pnorm(-abs(out1["r"])),
   rc_p = 2 * pnorm(-abs(out1["rc"])),
   cor_ml_p = 2 * pnorm(-abs(out1["wald_ml_adjusted"])),
    cor_rb_p = 2 * pnorm(-abs(out1["wald_rb_adjusted"])))
round(matrix(c(out1, out2), ncol = 10, byrow = TRUE,
             dimnames = list(NULL,
                             c("mle", "rb", "mcle", "wald_ml", "wald_mlc",
                               "wald_rb", "r", "rc", "wald_ml_adjusted",
                               "wald_rb_adjusted"))), 4)
          mle
                  rb mcle wald_ml wald_mlc wald_rb
# [1,] 1.4324 1.1562 1.2561 1.9511 1.8307 1.7362 2.1596 2.0214
# [2,] 0.7341 0.6659 0.6861 0.0510 0.0671 0.0825 0.0308 0.0432
       wald_ml_adjusted wald_rb_adjusted
# [1,]
                1.9257
                                 1.9064
# [2,]
                 0.0541
                                  0.0566
```

babies\_simulation.rda below is the output of babies\_simulation.R in ./code, which replicates the simulation study described in Section 9.4 of Di Caterina and Kosmidis (2019).

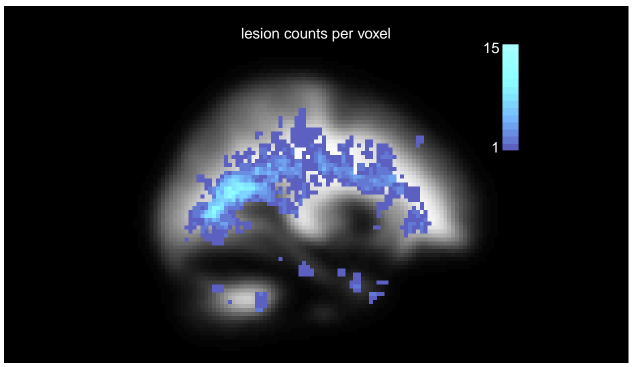
```
function(data) {
    if (all(data$type == "bootstrap_statistic")) {
        data.frame(sample = pnorm(data$value),
                   test = gsub("boot_prep_|boot_conv_", "", data$name))
   }
    else {
       p2 <- 2 * pnorm(-abs(data$value))</pre>
       pl <- pnorm(data$value)</pre>
       pr <- 1 - pl
        data.frame(sample = c(p2, p1, pr),
                   test = rep(c("2sided", "left", "right"), each = length(p2))) }
})
## Get rid of left right 2sided from statistic names
pval <- pval %>% mutate(name = gsub("_left|_right|_2sided", "", name))
pval <- pval %>%
   filter(!(name %in% c("scorec", "boot_prep")) & test != "right") %>%
    mutate(test = dplyr::recode(test,
                                "2sided" = "gamma != 0",
                                "left" = "gamma < 0",
                                "right" = "gamma > 0"),
           name = factor(name,
                         levels = c("mle", "rbe", "r", "cond", "scorec", "rc",
                                    "boot_conv", "cor", "cor_rb"),
                         ordered = TRUE)) %>%
   mutate(name = factor(name,
                         levels = c("mle", "r", "boot_conv", "rbe",
                                     "cond", "scorec", "rc",
                                     "cor", "cor_rb"),
                         ordered = TRUE)) %>%
   mutate(statistic = dplyr::recode(name,
                                     "mle" = "italic(t)",
                                     "rbe" = "italic(tilde(t))",
                                     "r" = "italic(r)",
                                     "cond" = "italic(t)[c]",
                                     "scorec" = "italic(s)[c]",
                                     "rc" = "italic(r)[c]",
                                     "cor" = "italic(t)^{'*},
                                     "cor rb" = "tilde(italic(t))^'*",
                                     "boot_conv" = "italic(boot)"))
## Bin sample
breaks <- (0:20)/20
pval <- pval %>%
    group by(statistic, test) %>%
   mutate(sample = cut(sample, breaks = breaks, include.lowest = TRUE)) %>%
    group_by(statistic, test, sample)
ggplot(pval) +
    geom_hline(aes(yintercept = 1)) +
    geom_bar(aes(x = sample, y = ..count../2500), fill = "darkgray", alpha = 0.5) +
   facet_grid(test ~ statistic, labeller = label_parsed) +
   theme bw() +
    scale_x_discrete(breaks = c("[0,0.05]", "(0.25,0.3]", "(0.5,0.55]",
                                "(0.75,0.8]", "(0.95,1]"),
                     labels = c(0, 0.25, 0.5, 0.75, 1)) +
```



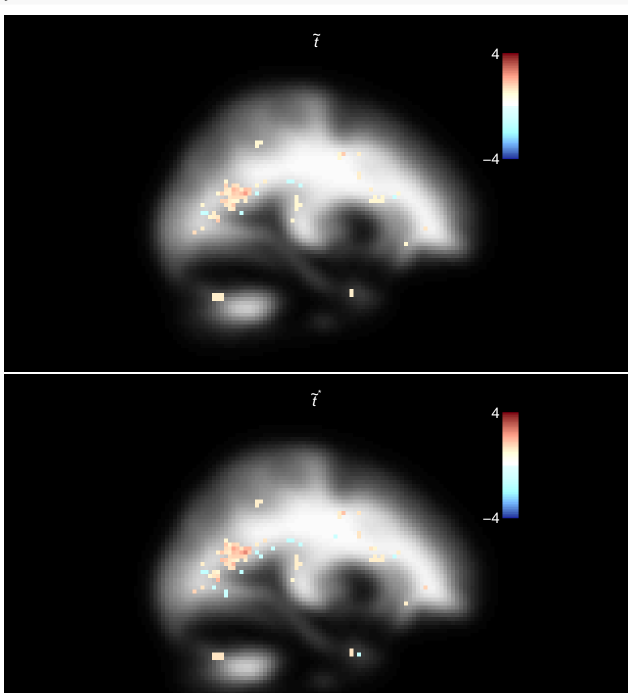
brains\_case\_study.rda below is the output of brains\_case\_study.R in ./code, which replicates the simulation study described in Section 10 of Di Caterina and Kosmidis (2019).

```
source(paste0(code_path, "/", "overlay2_nifti.R"))
load(paste(results_path, "brains_case_study.rda", sep = "/"))
## Check how many times LR failed, excluding trivial voxels, and compute probability of
## infinite estimates
fits_mat %>% filter(statistic == "r" & voxel != 1) %>% group_by(parameter) %>%
    summarize(failed = 100 * sum((value == -Inf) * count) / sum(count),
              infinite = 100 * sum(infinite * count) / sum(count))
# # A tibble: 6 x 3
   parameter failed infinite
   <fct>
              <db1>
                       <dbl>
               20.5
                        63.7
# 1 age
                        63.7
# 2 DD
               18.1
# 3 EDSS
               10.3
                        63.2
# 4 PASAT
              16.8
                        63.6
# 5 sex
               22.4
                        78.3
```

```
# 6 type2 19.2 75.5
## detections
fits mat %>%
   group_by(parameter, statistic) %>%
   filter(statistic %in% c("z br", "corz br")) %>%
   summarize(detections = mean(value < -1 | value > 1) * 100)
# # A tibble: 12 x 3
# # Groups: parameter [?]
   parameter statistic detections
    <fct> <fct>
                           <db1>
                            39.2
# 1 age
             corz\_br
# 2 age
             z\_br
                            33.0
# 3 DD
                            24.8
             corz\_br
# 4 DD
             z_br
                             18.9
# 5 EDSS
                            26.0
            corz\_br
# 6 EDSS
             z\_br
                            19.8
# 7 PASAT
             corz\_br
                            37.1
# 8 PASAT
             z\_br
                            29.9
# 9 sex
                            29.9
             corz\_br
# 10 sex
             z\_br
                            22.7
# 11 type2
                            22.1
             corz_br
# 12 type2
            z_br
                             17.1
## Empirical lesion counts
lesion_counts <- colSums(lesions)</pre>
lesion_counts[lesion_counts == 0] <- NA</pre>
nifti_counts <- nifti(img = array(lesion_counts, dim(white_matter)))</pre>
lumin <- c(45, 100)
cols_counts <-heat_hcl(n = max(lesion_counts, na.rm = TRUE),</pre>
                      h = c(265, 200),
                      c = c(80, 50),
                      1 = lumin,
                      power = c(0.7, 2))
overlay2.nifti(white_matter, y = nifti_counts, z = 32,
              plot.type = "single", plane = "sagittal",
              col.y = cols_counts, title = "lesion counts per voxel",
              col.main = "white")
```

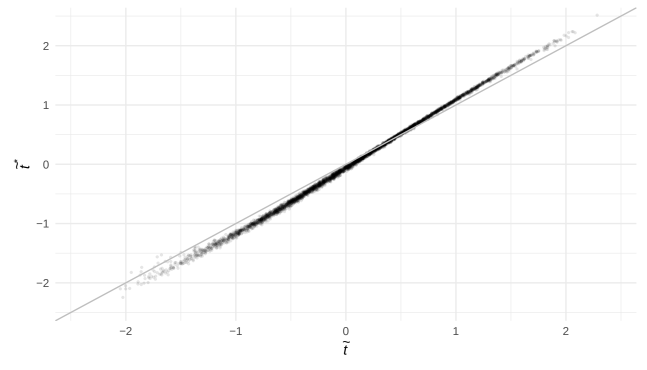


```
## Significance maps
param <- "DD"
low <- 1
upp <- 4
lumin <- c(25, 120)
cols \leftarrow c(heat_hcl(n = 32,
                     h = c(265, 200),
                     c = c(80, 50),
                     1 = lumin,
                     power = c(0.7, 2),
           rev(heat_hcl(n = 32,
                         h = c(10, 40),
                         c = c(80, 50),
                         1 = lumin,
                         power = c(0.4, 1.3)))
for (stat in c("z_br", "corz_br")) {
    zz <- (fits_mat %>% filter(statistic == stat & parameter == param))
    zz <- zz$value[array_indices]</pre>
    ## Threshold as in Ge et al (2014, AOAS)
    low_ind <- abs(zz) < low</pre>
    low_ind[is.na(low_ind)] <- FALSE</pre>
    zz[low_ind] <- NA</pre>
    upp_ind \leftarrow abs(zz) >= upp
    upp_ind[is.na(upp_ind)] <- FALSE</pre>
    zz[upp_ind] <- sign(zz[upp_ind]) * upp</pre>
    nifti_z <- nifti(img = array(zz, dim(white_matter)))</pre>
    nifti_z[1,1,1] <- -upp</pre>
    nifti_z[1,1,2] \leftarrow upp
    main <- switch(stat,</pre>
                     z_br = expression(tilde(italic(t))),
                     corz_br = expression(tilde(italic(t))^'*'))
```



```
### Plot z_br vs corz_br per parameter
v1 <- fits_mat %>%
    filter(statistic == "z_br", parameter == param) %>%
```

```
select(z_br_value = value, voxel, parameter)
v2 <- fits_mat %>%
    filter(statistic == "corz_br", parameter == param) %>%
    select(corz_br_value = value, voxel, parameter)
v <- join(v1, v2, by = c("voxel", "parameter"))
ggplot(v) +
    geom_point(aes(x = z_br_value, y = corz_br_value), alpha = 0.1, size = 0.5) +
    geom_abline(aes(intercept = 0, slope = 1), col = "grey") +
    coord_cartesian(xlim = c(-2.4, 2.4), ylim = c(-2.4, 2.4)) +
    theme_minimal() +
    labs(x = expression(tilde(italic(t))), y = expression(tilde(italic(t))^**'))</pre>
```



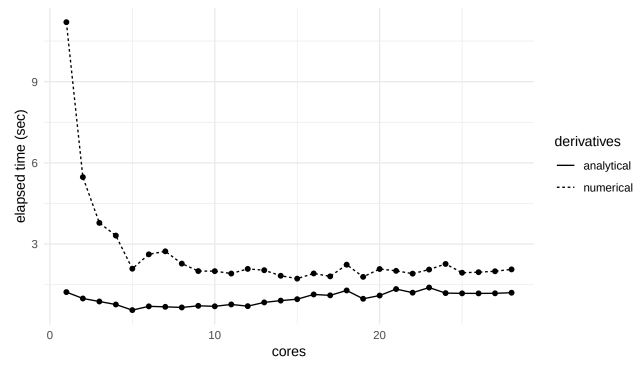
## **Execution times**

#### Timings in Concluding Remarks

```
numerical_time <- system.time(
    numerical <- waldi(babies_ml, numerical = TRUE, which = 19)
)
analytic_time <- system.time(
    analytic <- waldi(babies_ml, numerical = FALSE, which = 19)
)
(numerical_time/analytic_time)["elapsed"]
# elapsed
# 5.911111</pre>
```

#### Computational times and parallelization

babies\_times.rda below is the output of babies\_times.R in ./code. The figure illustrates the computational gain of using analytical derivatives in the calculation of the location-adjusted statistics with null value 0 for all 19 parameters in model (22) of the manuscript. When derivatives are obtained numerically, it is shown how the execution time decreases with the number of cores employed in the parallel implementation performed by the waldi R package.



## Meta-regression and meta-analysis

#### Wald statistics

The random-effects meta-regression model, i.e. the extension to random-effects meta-analysis for combining information from K studies about a common effect of interest (DerSimonian and Laird, 1986), assumes that  $Y_1, \ldots, Y_n$  are independent random variables conditionally on independent random effects  $U_1, \ldots, U_K$ . The

conditional distribution of  $Y_i$  given  $x_i$  and  $U_i = u_i$  is  $N(u_i + x_i^{\top} \beta, \hat{\sigma}_i^2)$ , where  $\hat{\sigma}_i^2$  is the known within-study variance and  $u_i$  is the realization of a  $N(0, \psi)$ . The unknown parameters are the k-vector of effects  $\beta$  and the between-study heterogeneity  $\psi$ . Note that this specification reduces to the meta-analysis model if  $x_i = 1$  for every  $i = 1, \ldots, K$ . The expected information matrix depends only on  $\psi$  and is

$$i(\psi) = \begin{bmatrix} X^{\top}W(\psi)X & 0_k \\ 0_k^{\top} & \frac{1}{2}\mathrm{tr}\{W(\psi)^2\} \end{bmatrix}, \tag{1}$$

where  $W(\psi) = \text{diag}\{w_1, \dots, w_n\}$ , with  $w_i = (\hat{\sigma}_i^2 + \psi)^{-1}$ . Thus, the Wald statistic for testing  $H_0: \beta_j = \beta_{j0}$  takes the form  $t_j = (\hat{\beta}_j - \beta_{j0})/\kappa_j(\hat{\psi})$   $(j = 1, \dots, k)$ , and can be seen as the estimate of the transformation  $T_j(\beta, \psi; \beta_{j0}) = (\beta_j - \beta_{j0})/\kappa_j(\psi)$ , where  $\kappa_j(\psi) = [\{X^\top W(\psi)X\}^{-1}]_{ij}^{1/2}$ .

Guolo and Varin (2015) provide extensive evidence that Wald and other first-order likelihood-based procedures can be highly inaccurate when the number of studies is small or moderate. In these cases, Kosmidis et al. (2017) illustrate that the downward bias of the maximum likelihood estimator  $\hat{\psi}$  affects conclusions on  $\beta$ , leading to anti-conservative tests and narrow confidence intervals, and propose asymptotic bias reduction as a means to refine first-order inference about the mean effect size.

#### Implementation

The calculation of the location-adjusted Wald statistic for the simulation studies is done analytically using the formula for the first-order bias of  $\hat{\psi}$ , namely  $b_{\psi}(\psi) = -\text{tr}\{W(\psi)H(\psi)\}/\text{tr}\{W(\psi)^2\}$  where  $H(\psi) = X\{X^{\top}W(\psi)X\}^{-1}X^{\top}W(\psi)$  (Kosmidis et al., 2017, equation (5)), the derivatives of  $T_j(\beta, \psi; \beta_{j0})$ , and the expression of the information matrix  $i(\psi)$ . The partial derivatives of the latter with respect to  $\beta$  are zero, while

$$\frac{di(\psi)}{d\psi} = \begin{bmatrix} X^{\top}W'(\psi)X & 0_k \\ 0_k^{\top} & -\sum_{i=1}^K (\hat{\sigma}_i^2 + \psi)^{-3} \end{bmatrix},$$

and

$$\frac{d^2 i(\psi)}{d\psi^2} = \begin{bmatrix} X^\top W''(\psi) X & 0_k \\ 0_k^\top & 3\sum_{i=1}^K (\hat{\sigma}_i^2 + \psi)^{-4} \end{bmatrix},$$

where  $W'(\psi) = \text{diag}\{w_1', \dots, w_n'\}$ , with  $w_i' = -(\hat{\sigma}_i^2 + \psi)^{-2}$  and  $W''(\psi) = \text{diag}\{w_1'', \dots, w_n''\}$ , with  $w_i'' = 2(\hat{\sigma}_i^2 + \psi)^{-3}$ .

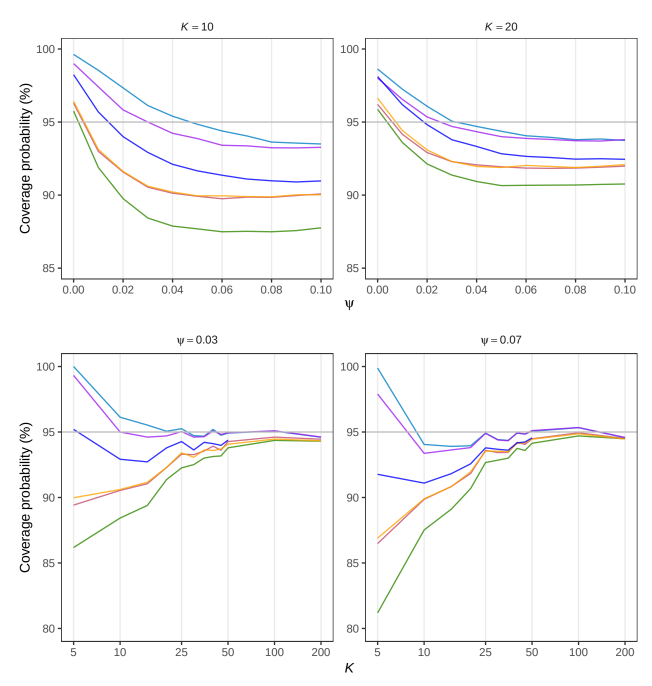
#### Simulation studies

This section illustrates results of the simulations performed under the random-effects meta-analysis model using the same design as in Brockwell and Gordon (2001). In particular, the realizations  $y_i$  are simulated from a meta-analysis model with  $\beta=0.5$  and variance  $\hat{\sigma}_i+\psi$ , where  $\hat{\sigma}_i$  are independently generated from a  $\chi^2_1$  distribution multiplied by 0.25 and then restricted to the interval (0.009, 0.6). The parameter  $\psi$  ranges from 0 to 0.1, and the number of studies K from 5 to 200. Due to long execution times, we have ran the experiment based on 10 000 Monte Carlo iterations for only those combinations of  $\psi$  and K which are needed to reproduce below Figure 1 of Kosmidis et al. (2017).

Note that meta\_analysis\_simulation.rda below is the output of meta\_analysis\_simulation.R in ./code. load(paste(results\_path, "meta\_analysis\_simulation.rda", sep = "/"))
### plot K = 10, 20

```
fig1 <- ggplot(cov_df_K %>% filter(test == "2sided")) +
  geom_line(aes(psi, cov, group = statistic, col = statistic), size = 0.5, alpha = 0.8) +
  geom_hline(aes(yintercept = 95), col = "grey") +
  facet_wrap( ~ Klab, labeller = label_parsed, scales = "free") +
  labs(y = "Coverage probability (%)", x = expression(psi)) +
  lims(y = c(85, 100)) +
  scale_x_continuous(name = expression(psi), breaks = seq(0, 0.1, length = 6),
                     labels = c("0.00", "0.02", "0.04", "0.06", "0.08", "0.10")) +
  scale_colour_manual(name = "", values = c("#328900", "#0080C5", "#C54E6D", "purple",
        "orange", 4), labels = c(expression(italic(t)), expression(italic(t)^list("*")),
        expression(tilde(italic(t))),
        expression(tilde(italic(t))^list("*")), "DL", "ZL")) +
  theme bw() +
  theme(legend.position = "top", panel.grid.major.y = element_blank(),
        panel.grid.minor.y = element_blank(), panel.grid.minor.x = element_blank(),
        strip.background = element_blank()) +
  guides(colour = guide_legend(nrow = 1))
## plot psi = 0.03, 0.07
fig2 <- ggplot(cov_df_psi %>% filter(test == "2sided")) +
  geom_line(aes(log(K), cov, group = statistic, col = statistic), size = 0.5,
            alpha = 0.8) +
  geom_hline(aes(yintercept = 95), col = "grey") +
  facet_wrap( ~ psilab, labeller = label_parsed, scales = "free") +
  labs(y = "Coverage probability (%)", x = expression(italic(K))) +
  lims(y = c(80, 100)) +
  scale x continuous(name = expression(italic(K)), breaks = c(log(5), log(10),
                    log(25), log(50), log(100), log(200)), labels =
                      c("5", "10", "25", "50", "100", "200")) +
  scale_colour_manual(name = "", values = c("#328900", "#0080C5", "#C54E6D",
                      "purple", "orange", 4), labels = c(expression(italic(t)),
                      expression(italic(t)^list("*")), expression(tilde(italic(t))),
                      expression(tilde(italic(t))^list("*")), "DL", "ZL")) +
  theme_bw() +
  theme(legend.position = "none", panel.grid.major.y = element_blank(),
        panel.grid.minor.y = element_blank(), panel.grid.minor.x = element_blank(),
        strip.background = element_blank())
```





The last plots show the empirical coverage probabilities of individual confidence intervals for  $\beta$ , as  $\psi$  varies when  $K \in \{10, 20\}$  (top row) and as K increases (in log scale) when  $\psi \in \{0.03, 0.07\}$  (bottom row). The curves correspond to  $t, t^*, \tilde{t}, \tilde{t}^*$ , the Wald statistic based on the DerSimonian & Laird estimator (DL) and, for  $K \leq 50$ , the Zeng & Lin double-resampling method (ZL). The grey horizontal line is the target 95% nominal level.

The inversion of the location-adjusted Wald statistic, both that based on maximum likelihood and that based on reduced-bias estimates, results in intervals with empirical coverage that is remarkably close to the nominal level in all scenarios. Whereas the adoption of  $\tilde{t}$  produces an improvement over t which is similar to that implied by using the DerSimonian & Laird estimator of  $\psi$  (DerSimonian and Laird, 1986) in the standard

Wald statistic, the adjustment in location performed through  $t^*$  and  $\tilde{t}^*$  proves to be even more effective than the double resampling, despite the significantly higher computational intensity of the approach proposed by Zeng & Lin (2015).

#### References

Agresti, A., and B. Caffo. 2000. "Simple and Effective Confidence Intervals for Proportions and Differences of Proportions Result from Adding Two Successes and Two Failures." *The American Statistician* 54: 280–88.

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