S1: Group comparisons and behavioural analysis with brms

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Introduction

In this study, participants performed a change detection task in the fMRI scanner. Specifically, they were asked to press a button as fast as possible when the fixation cross changed orientation. While they performed this task, four faces were presented in the periphery for a short duration. These faces were coloured in pink or green and showed either a fearful or a happy expression. While the neuroimaging analysis focuses on the building and violation of expectations based on these faces, this RMarkdown focuses on the analysis of the behavioural data.

Some general settings

```
# number of simulations
nsim = 250

# set number of iterations and warmup for models
iter = 4500
warm = 1500
```

Package versions

```
## [1] "R version 4.5.0 (2025-04-11)"
## [1] "knitr version 1.50"
## [1] "ggplot2 version 3.5.2"
## [1] "brms version 2.22.0"
## [1] "designr version 0.1.13"
## [1] "bridgesampling version 1.1.2"
## [1] "tidyverse version 2.0.0"
## [1] "ggpubr version 0.6.0"
## [1] "officer version 0.6.10"
## [1] "bayesplot version 1.12.0"
## [1] "bayesplot version 0.7.2"
## [1] "BayesFactor version 0.9.12.4.7"
## [1] "bayestestR version 0.16.0"
```

General info

We planned to determine the group-level effect subjects following Barr (2013). For each model, experiment specific priors were set based on previous literature or the task (see comments in the code).

We perform prior predictive checks as proposed in Schad, Betancourt and Vasishth (2020) using the SBC package. To do so, we create simulated datasets where parameters are simulated from the priors. These parameters are used to create one fake dataset. Both the true underlying parameters and the simulated values are saved.

Then, we create graphs showing the prior predictive distribution of the simulated discrimination threshold to check whether our priors fit our general expectations about the data. Next, we perform checks of computational faithfulness and model sensitivity as proposed by Schad, Betancourt and Vasishth (2020) and implemented in the SBC package. We create models for each of the simulated datasets. Last, we calculate performance metrics for each of these models, focusing on the population-level parameters.

Preparation and group comparisons

First, we load the data and combine it with demographic information including the diagnostic status of the subjects. Then, all predictors are set to sum contrasts. We have a look at the demographics describing our three diagnostic groups: autistic adults and adults with ADHD as well as a comparison group of adults without any psychiatric diagnoses.

Since this is sensitive data, we load the anonymised version of the processed data at this point but also leave the code we used to create it.

```
# check if the data file exists, if yes load it:
if (!file.exists(file.path(model_dir, "VMM_data.RData"))) {
  # set file paths
  fl.path = '/home/emba/Documents/EMBA'
  dt.path = paste(fl.path, 'BVET', sep = "/")
  # load the preprocessed data
  load(file.path(dt.path, "VMM_preprocessed.RData"))
  # load list of participants which are included in the fMRI analysis
  df.inc = read_delim("all_use-new") %>%
   filter(diagnosis != "pilot") %>% select(subID)
  # load list of participants which are included in fMRI ET analysis
  df.ETinc = read_csv(file = file.path(dt.path, "VMM_ET_inc.csv"), show_col_types = F)
  # load demographic information
  df.sub = read_csv(file.path("/home/emba/Documents/EMBA/CentraXX", "EMBA_centraXX.csv"),
                    show_col_types = F) %>%
   mutate(
      diagnosis = recode(diagnosis, "CTR" = "COMP")
    ) %>% filter(subID %in% df.inc$subID)
  # load information on medication
  df.meds = read_csv(file.path("/home/emba/Documents/EMBA/CentraXX",
                               "VMM medication.csv"))
  # merge together and anonymise
  df.disc = merge(df.disc, df.sub %>% select(subID, diagnosis), all.y = T) %>%
   mutate(
     subID = as.factor(as.numeric(as.factor(subID)))
   )
  df.tsk = merge(df.tsk, df.sub %>% select(subID, diagnosis), all.y = T) %>%
   mutate(
```

```
subID = as.factor(as.numeric(as.factor(subID)))
 )
# set the center for the fixation cross
x.centre = 512 # 1024
y.centre = 332 # 768 but moved up 52 pixels
r = 100 # radius of the AOI in the centre
# classify fixates (centre or not), aggregate and anonymise the data
df.fix = readRDS(file.path(dt.path, "VMM_fixations.rds"))
df.fix.agg = df.fix %>%
 mutate(
    AOI.fix = if_else(dist.centre <= r, "centre", "periphery")
 ) %>%
 group_by(subID, AOI.fix) %>%
 summarise(
   fix.dur = sum(duration)
 pivot_wider(names_from = AOI.fix, values_from = fix.dur, names_prefix = "fix.") %>%
 mutate(
   fix.total = fix.centre + fix.periphery,
   fix.prop = fix.centre / fix.total
 ) %>%
 merge(., df.ETinc %>% select(subID, diagnosis) %>% distinct()) %>%
 ungroup() %>%
 mutate(
   rfix.total = rank(fix.total),
   rfix.prop = rank(fix.prop),
   diagnosis = as.factor(diagnosis),
   subID = as.factor(as.numeric(as.factor(subID)))
 )
# get gender frequencies and compare them across groups
tb.gen = xtabs(~ gender + diagnosis, data = df.sub)
ct.full = contingencyTableBF(tb.gen,
                             sampleType = "indepMulti",
                             fixedMargin = "cols")
# since only DAN in the ADHD group, we try again after excluding them
ct.mf = contingencyTableBF(tb.gen[2:3,],
                           sampleType = "indepMulti",
                           fixedMargin = "cols")
# we add this information to our demographics tablerbind(df.demo,
df.demo = data.frame(
                  measurement = "Gender",
                               = sprintf("%.0f - %.0f - %.0f",
                  ADHD
                                        tb.gen["fem","ADHD"],
                                        tb.gen["mal","ADHD"],
                                        tb.gen["dan", "ADHD"]
                  ),
                              = sprintf("%.0f - %.0f - %.0f",
                  ASD
                                        tb.gen["fem","ASD"],
                                        tb.gen["mal","ASD"],
```

```
tb.gen["dan", "ASD"]
                  ),
                  COMP
                               = sprintf("%.0f - %.0f - %.0f",
                                        tb.gen["fem","COMP"],
                                        tb.gen["mal", "COMP"],
                                        tb.gen["dan","COMP"]
                  ),
                  bf.log
                              = ct.full@bayesFactor[["bf"]]
# keep the gender descriptions of the DAN group
gen.desc = unique(df.sub[df.sub$gender == "dan",]$gender_desc)
# get education frequencies and compare them across groups
tb.edu = xtabs(~ edu + diagnosis, data = df.sub)
ct.edu = contingencyTableBF(tb.edu,
                             sampleType = "indepMulti",
                             fixedMargin = "cols")
# convert the measures to long which we include in the participant table
df.sublng = df.sub %>%
  # rename some of the variables
 rename(
    "RADS-R" = "RAADS_total",
    "BDI" = "BDI total",
   "ASRS-v1.1" = "ASRS total",
    "Age" = "age",
    "IQ estimate" = "iq"
 select(diagnosis, Age, `IQ estimate`, `ASRS-v1.1`, `RADS-R`, BDI) %>%
 pivot_longer(cols = where(is.numeric)) %>%
 mutate_if(is.character, as.factor)
# initialise the data frame for posthoc tests
df.post = data.frame()
# now we loop through our measurements to create our demographics table
for (m in unique(df.sublng$name)) {
  # select the relevant part of df.sub
 df.rel = df.sublng %>% filter(name == m)
  # check which of the group's data is not normally distributed
 df.sht = df.rel %>%
    group by(diagnosis) %>%
   shapiro_test(value) %>%
   filter(p < 0.05)
  # if more than zero is not normally distributed...
  if (nrow(df.sht) > 0) {
    # rank transform the data
   df.rel = df.rel %>% ungroup() %>% mutate(value = rank(value))
  # compute the ANOVA
 aov = anovaBF(value ~ diagnosis, data = df.rel)
  # get back the original, untransformed values
```

```
df.rel = df.sublng %>% filter(name == m)
  # put all the information into the demographics table
  df.demo = rbind(df.demo,
                  data.frame(
                    measurement = m,
                    ADHD
                                = sprintf("%.2f ±%.2f (%.0f to %.0f)",
                                          # ignore NAs because edu missing for one person
                                          mean(df.rel[df.rel$diagnosis == "ADHD",]$value, na.rm = T),
                                          sd(df.rel[df.rel$diagnosis == "ADHD",]$value, na.rm = T)/
                                            sqrt(sum(df.rel$diagnosis == "ADHD")),
                                          min(df.rel[df.rel$diagnosis == "ADHD",]$value, na.rm = T),
                                          max(df.rel[df.rel$diagnosis == "ADHD",]$value, na.rm = T)
                    ),
                    ASD
                                = sprintf("\%.2f ±\%.2f (%.0f to %.0f)",
                                          mean(df.rel[df.rel$diagnosis == "ASD",]$value),
                                          sd(df.rel[df.rel$diagnosis == "ASD",]$value)/
                                            sqrt(sum(df.rel$diagnosis == "ASD")),
                                          min(df.rel[df.rel$diagnosis == "ASD",]$value),
                                          max(df.rel[df.rel$diagnosis == "ASD",]$value)
                    ),
                    COMP
                                = sprintf("\%.2f ±\%.2f (%.0f to %.0f)",
                                          mean(df.rel[df.rel$diagnosis == "COMP",]$value),
                                          sd(df.rel[df.rel$diagnosis == "COMP",]$value)/
                                            sqrt(sum(df.rel$diagnosis == "COMP")),
                                          min(df.rel[df.rel$diagnosis == "COMP",]$value),
                                          max(df.rel[df.rel$diagnosis == "COMP",]$value)
                    ),
                    bf.log
                                = aov@bayesFactor[["bf"]]
                  ))
  # next, we want to check whether there are group differences
  if (abs(exp(aov@bayesFactor$bf)) > 3) {
    # do the group comparisons
    aov.ADHDvASD = anovaBF(value ~ diagnosis,
                            data = df.rel %>% filter(diagnosis %in% c("ADHD", "ASD")))
    aov.ADHDvCOMP = anovaBF(value ~ diagnosis,
                            data = df.rel %>% filter(diagnosis %in% c("ADHD", "COMP")))
    aov.ASDvCOMP = anovaBF(value ~ diagnosis,
                            data = df.rel %>% filter(diagnosis %in% c("ASD", "COMP")))
    # put into the posthoc data frame
    df.post = rbind(df.post,
                  data.frame(
                    measurement = m,
                    ADHDvASD = aov.ADHDvASD@bayesFactor[["bf"]],
                    ADHDvCOMP = aov.ADHDvCOMP@bayesFactor[["bf"]],
                             = aov.ASDvCOMP@bayesFactor[["bf"]]
                    ASDvCOMP
                  ))
 }
}
# save the demographics and the posthoc table as word documents
read_docx() %>%
 body_add_table(df.demo %>% arrange(measurement) %>%
```

```
mutate(bf.log =
                                  if else(
                                    bf.log > 3,
                                    sprintf("%.3f*", bf.log),
                                    sprintf("%.3f", bf.log)))) %>%
    print(target = "VMM_demo.docx")
  read_docx() %>%
    body_add_table(df.post %>%
                     mutate_if(is.numeric,
                               ~ifelse(.>3,sprintf("%.3f*", .),sprintf("%.3f", .)))) %>%
    print(target = "VMM_post.docx")
  # check how many of each group are above threshold for asrs and rads
  tb.screen = xtabs(~ diagnosis + screening,
                   data = df.sub %>%
                     select(diagnosis, ASRS_screen, RAADS_total) %>%
                     mutate(
                       screening = case_when(
                         ASRS_screen >= 4 & RAADS_total > 81 ~ "screenBOTH",
                         ASRS_screen >= 4 & RAADS_total <= 81 ~ "screenADHD",
                         ASRS_screen < 4 & RAADS_total <= 81 ~ "screenNone",
                         ASRS_screen < 4 & RAADS_total > 81 ~ "screenASD"
                       )
                     ))
  # save it all
  save(df.disc, df.tsk, df.fix.agg, tb.screen,
       ct.full, ct.mf, gen.desc, df.meds, tb.edu, ct.edu,
       file = file.path(model_dir, "VMM_data.RData"))
} else {
  load(file.path(model_dir, "VMM_data.RData"))
}
# print the group of included participants
kable(df.disc %>% select(subID, diagnosis) %>% distinct() %>% group_by(diagnosis) %>% count())
```

diagnosis	n
ADHD	23
ASD	22
COMP	22

print the group of included participants with eye tracking kable(df.fix.agg %>% select(subID, diagnosis) %>% distinct() %>% group_by(diagnosis) %>% count())

diagnosis	n
ADHD	16
ASD	11
CTR	15

```
# print the outcome of the contingency table concerning gender
ct.full@bayesFactor
##
                           bf error
                                                                       code
                                                         time
## Non-indep. (a=1) -2.932155
                                  0 Mon Feb 17 10:45:09 2025 2d6a747c15f0d
ct.mf@bayesFactor
                           bf error
                                                         time
                                                                       code
## Non-indep. (a=1) -1.088224
                                  0 Mon Feb 17 10:45:09 2025 2d6a77dfa2c24
# print the outcome of the contingency table concerning education
ct.edu@bayesFactor
##
                           bf error
                                                         time
## Non-indep. (a=1) -1.834707
                                  0 Mon Feb 17 10:45:09 2025 2d6a731fddab1
# only keep the relevant rtcs
df.rtc = df.tsk %>%
  filter(sdt == "hit" & !is.na(rtc)) %>%
  ungroup() %>%
 mutate if(is.character, as.factor)
df.disc = df.disc %>%
  ungroup() %>%
  # subtract the discrimination rate from the "perfect" number to be able to
  # use a poisson in the model
  mutate(
   negdisc = 240 - disc
  ) %>%
 mutate_if(is.character, as.factor)
# set and print the contrasts
contrasts(df.rtc$diagnosis) = contr.sum(3)
contrasts(df.rtc$diagnosis)
        [,1] [,2]
##
## ADHD
          1
## ASD
           0
                1
## COMP
               -1
        -1
contrasts(df.disc$diagnosis) = contr.sum(3)
contrasts(df.disc$diagnosis)
        [,1] [,2]
## ADHD
          1
## ASD
           0
## COMP
         -1
              -1
contrasts(df.fix.agg$diagnosis) = contr.sum(3)
contrasts(df.fix.agg$diagnosis)
        [,1] [,2]
## ADHD
          1
## ASD
           0
## CTR
         -1
               -1
# print medication for the paper
kable(df.meds)
```

diagnosis	adhd.meds	psych.meds_binary	n
ADHD	atomoxetine	TRUE	1
ADHD	lisdexamfetamine	TRUE	4
ADHD	methylphenidate	TRUE	8
ADHD	NA	FALSE	8
ADHD	NA	TRUE	2
ASD	NA	FALSE	11
ASD	NA	TRUE	11
COMP	NA	FALSE	22

Analysis of reaction times (hits-only)

Full model

Group-level: subject and trials, trials with diagnosis slope

```
code = "VMM_rtc_full"
# full model formula
f.rtc = brms::bf(rtc ~ diagnosis + (1 | subID) + (diagnosis | trl) )
# set informed priors based on previous results
priors = c(
  # general priors based on SBV
  prior(normal(6, 0.3), class = Intercept),
 prior(normal(0,
                   0.5), class = sigma),
  prior(normal(0.1, 0.1), class = sd),
  prior(lkj(2),
                            class = cor),
  # ADHD subjects being slower based on Pievsky & McGrath (2018)
  prior(normal(0.025, 0.04), class = b, coef = diagnosis1),
  # ASD subjects being slower based on Morrison et al. (2018)
  prior(normal(0.025, 0.04), class = b, coef = diagnosis2),
  prior(normal(100,
                   50), class = ndt)
# check if the SBC already exists
if (file.exists(file.path(cache_dir, sprintf("df_res_%s.rds", code)))) {
  # load in the results of the SBC
 df.results = readRDS(file.path(cache_dir, sprintf("df_res_%s.rds", code)))
 df.backend = readRDS(file.path(cache_dir, sprintf("df_div_%s.rds", code)))
            = readRDS(file.path(cache_dir, sprintf("dat_%s.rds", code)))
} else {
  # set the seed
  set.seed(2468)
  # perform the SBC
  gen = SBC_generator_brms(f.rtc, data = df.rtc, prior = priors,
                           family = shifted_lognormal,
                           thin = 50, warmup = 20000, refresh = 2000)
  bck = SBC_backend_brms_from_generator(gen, chains = 4, thin = 1,
                                     init = 0.1, warmup = warm, iter = iter)
  if (!file.exists(file.path(cache_dir, sprintf("dat_%s.rds", code)))) {
   dat = generate_datasets(gen, nsim)
```

```
saveRDS(dat, file.path(cache_dir, sprintf("dat_%s.rds", code)))
} else {
    dat = readRDS(file.path(cache_dir, sprintf("dat_%s.rds", code)))
}

# set the seed again
set.seed(2468)
res = compute_SBC(dat,
    bck,
    cache_mode = "results",
    cache_location = file.path(cache_dir, sprintf("res_%s", code)))
df.results = res$stats
df.backend = res$backend_diagnostics
saveRDS(df.results, file = file.path(cache_dir, paste0("df_res_", code, ".rds")))
saveRDS(df.backend, file = file.path(cache_dir, paste0("df_div_", code, ".rds")))
}
```

We start by investigating the rhats and the number of divergent samples. This shows that 19 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and 60 models had divergent samples (mean number of samples of the simulations with divergent samples: 190.23). This suggests serious problems with this model. We try to drop the group-level slope of diagnostics for the trials and run it again.

Group-level: subject and trials, no slopes

```
code = "VMM_rtc_full_int"
# full model formula
f.rtc = brms::bf(rtc ~ diagnosis + (1 | subID) + (1 | trl) )
# set informed priors based on previous results
priors = c(
  # general priors based on SBV
                    0.3), class = Intercept),
 prior(normal(6,
 prior(normal(0,
                     0.5), class = sigma),
  prior(normal(0.1,
                     0.1), class = sd),
  # ADHD subjects being slower based on Pievsky & McGrath (2018)
  prior(normal(0.025, 0.04), class = b, coef = diagnosis1),
  # ASD subjects being slower based on Morrison et al. (2018)
  prior(normal(0.025, 0.04), class = b, coef = diagnosis2),
  # shift
  prior(normal(100, 50),
                             class = ndt)
# check if the SBC already exists
if (file.exists(file.path(cache_dir, sprintf("df_res_%s.rds", code)))) {
  # load in the results of the SBC
  df.results = readRDS(file.path(cache_dir, sprintf("df_res_%s.rds", code)))
  df.backend = readRDS(file.path(cache_dir, sprintf("df_div_%s.rds", code)))
 dat
            = readRDS(file.path(cache_dir, sprintf("dat_%s.rds", code)))
} else {
  # set the seed
  set.seed(2468)
  # perform the SBC
  gen = SBC_generator_brms(f.rtc, data = df.rtc, prior = priors,
                           family = shifted_lognormal,
```

```
thin = 50, warmup = 20000, refresh = 2000)
  bck = SBC_backend_brms_from_generator(gen, chains = 4, thin = 1,
                                      init = 0.1, warmup = warm, iter = iter)
  if (!file.exists(file.path(cache_dir, sprintf("dat_%s.rds", code)))) {
   dat = generate_datasets(gen, nsim)
    saveRDS(dat, file.path(cache_dir, sprintf("dat_%s.rds", code)))
  } else {
   dat = readRDS(file.path(cache dir, sprintf("dat %s.rds", code)))
  }
  # set the seed again
  set.seed(2468)
  res = compute_SBC(dat,
        bck,
                       = "results".
        cache mode
        cache_location = file.path(cache_dir, sprintf("res_%s", code)))
  df.results = res$stats
  df.backend = res$backend_diagnostics
  saveRDS(df.results, file = file.path(cache_dir, paste0("df_res_", code, ".rds")))
  saveRDS(df.backend, file = file.path(cache_dir, paste0("df_div_", code, ".rds")))
}
```

Again, we start by investigating the rhats and the number of divergent samples. This shows that 24 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and 81 models had divergent samples (mean number of samples of the simulations with divergent samples: 120.36). This suggests that there are still some serious problems with this model. Therefore, we change course and try to aggregate the trials for each participant to see if we can find a better model for our simulated data before we switch to using our real data.

Aggregated model

SBC

```
code = "VMM_rtc_agg_simple"
# aggregate the data and set the contrast
df.rtc.agg = df.rtc %>%
  group_by(subID, diagnosis) %>%
  summarise(
   rtc = median(rtc)
  )
# model formula
f.rtc = brms::bf(rtc ~ diagnosis )
# set weakly informed priors
priors = c(
  # general priors based on SBV
  prior(normal(5.6,
                      0.3), class = Intercept), # median gets rid of outliers > lower
  prior(normal(0,
                      0.5), class = sigma),
  # ADHD subjects being slower based on Pievsky & McGrath (2018)
  prior(normal(0.025, 0.10), class = b, coef = diagnosis1),
  # ASD subjects being slower based on Morrison et al. (2018)
  prior(normal(0.025, 0.10), class = b, coef = diagnosis2),
  # shift > had min. 100 for reactions build in, but median is going to be higher
```

```
prior(normal(200, 100), class = ndt)
# Increased the standard deviation of b after plotting contraction.
# check if the SBC already exists
if (file.exists(file.path(cache_dir, sprintf("df_res_%s.rds", code)))) {
  # load in the results of the SBC
  df.results = readRDS(file.path(cache_dir, sprintf("df_res_%s.rds", code)))
  df.backend = readRDS(file.path(cache_dir, sprintf("df_div_%s.rds", code)))
            = readRDS(file.path(cache_dir, sprintf("dat_%s.rds", code)))
} else {
  # set the seed
  set.seed(2468)
  # perform the SBC
  gen = SBC_generator_brms(f.rtc, data = df.rtc.agg, prior = priors,
                           family = shifted_lognormal,
                           thin = 50, warmup = 20000, refresh = 2000)
  bck = SBC_backend_brms_from_generator(gen, chains = 4, thin = 1,
                                      init = 0.1, warmup = warm, iter = iter)
  dat = generate_datasets(gen, nsim)
  saveRDS(dat, file.path(cache_dir, sprintf("dat_%s.rds", code)))
  res = compute_SBC(dat,
       bck,
        cache mode
                     = "results",
        cache_location = file.path(cache_dir, sprintf("res_%s", code)))
  df.results = res$stats
  df.backend = res$backend_diagnostics
  saveRDS(df.results, file = file.path(cache_dir, paste0("df_res_", code, ".rds")))
  saveRDS(df.backend, file = file.path(cache_dir, paste0("df_div_", code, ".rds")))
}
```

Again, we start by investigating the rhats and the number of divergent samples. This shows that 0 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and 0 models had divergent samples. This suggests that this model performs well and we can continue with our checks by plotting the simulated values to perform prior predictive checks.

```
# create a matrix out of generated data
dvname = gsub(" ", "", gsub("[\\\-].*", "", f.rtc)[1])
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))
for (i in 1:length(dat[['generated']])) {
    dvfakemat[,i] = dat[['generated']][[i]][[dvname]]
}
truePars = dat$variables

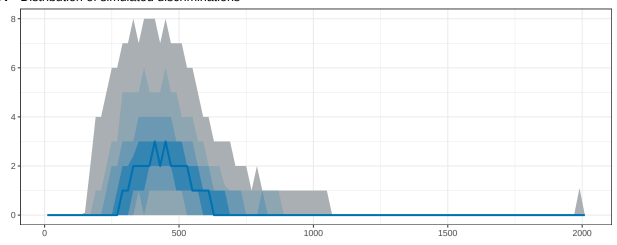
# set large values to a max
dvfakemat[dvfakemat > 2000] = 2000

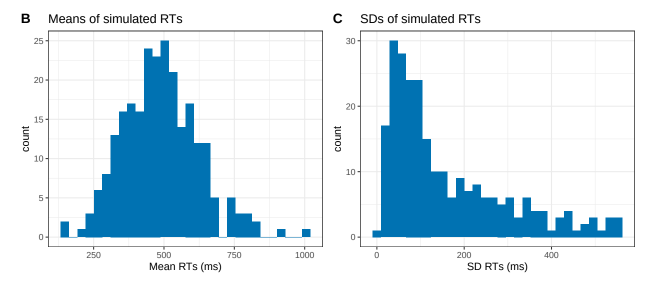
# compute one histogram per simulated data-set
binwidth = 20
breaks = seq(0, max(dvfakemat, na.rm=T) + binwidth, binwidth)
histmat = matrix(NA, ncol = length(dat), nrow = length(breaks)-1)
for (i in 1:nrow(truePars)) {
```

```
histmat[,i] = hist(dvfakemat[,i], breaks = breaks, plot = F)$counts
}
# for each bin, compute quantiles across histograms
probs = seq(0.1, 0.9, 0.1)
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))
names(quantmat) = paste0("p", probs)
for (i in 1:dim(histmat)[1]) {
  quantmat[i,] = quantile(histmat[i,], p = probs)
}
quantmat$x = breaks[2:length(breaks)] - binwidth/2 # add bin mean
p1 = ggplot(data = quantmat, aes(x = x)) +
  geom_ribbon(aes(ymax = p0.9, ymin = p0.1), fill = c_light) +
  geom_ribbon(aes(ymax = p0.8, ymin = p0.2), fill = c_light_highlight) +
  geom_ribbon(aes(ymax = p0.7, ymin = p0.3), fill = c_mid) +
  geom_ribbon(aes(ymax = p0.6, ymin = p0.4), fill = c_mid_highlight) +
  geom_line(aes(y = p0.5), colour = c_dark, linewidth = 1) +
  labs(title = "Distribution of simulated discriminations", y = "", x = "") +
  theme_bw()
tmpM = apply(dvfakemat, 2, mean) # mean
tmpSD = apply(dvfakemat, 2, sd)
p2 = ggplot() +
  stat_bin(aes(x = tmpM), fill = c_dark) +
  labs(x = "Mean RTs (ms)", title = "Means of simulated RTs") +
  theme bw()
p3 = ggplot() +
  stat_bin(aes(x = tmpSD), fill = c_dark) +
  labs(x = "SD RTs (ms)", title = "SDs of simulated RTs") +
 theme_bw()
p = ggarrange(p1,
  ggarrange(p2, p3, ncol = 2, labels = c("B", "C")),
  nrow = 2, labels = "A")
annotate_figure(p,
                top = text_grob("Prior predictive checks: reaction times",
                face = "bold", size = 14))
```

Prior predictive checks: reaction times

A Distribution of simulated discriminations



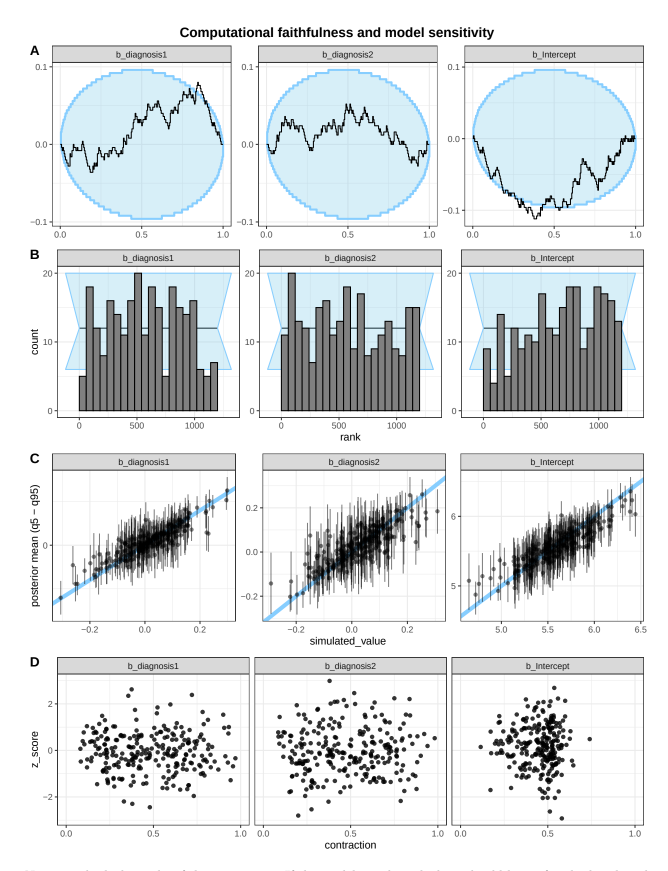


Subfigure A shows the distribution of the simulated data with bluer bands being more likely than greyer bands. It shows a distribution that fits our expectations about reaction times in a simple decision task. The same applies to the distribution of the means and standard deviations in the simulated datasets. We go ahead with these priors and check the results of the SBC. We only plot the results from the models that had no divergence issues.

```
# get simulation numbers with issues
rank = max(df.results$max_rank)
check = merge(df.results %>%
    group_by(sim_id) %>%
    summarise(
        rhat = max(rhat, na.rm = T),
        mean_rank = mean(max_rank)
        ) %>%
    filter(rhat >= 1.05 | mean_rank != rank),
    df.backend %>% filter(n_divergent > 0), all = T)

# plot SBC with functions from the SBC package focusing on population-level parameters
df.results.b = df.results %>%
```

```
filter(substr(variable, 1, 2) == "b_") %>%
  filter(!(sim_id %in% check$sim_id))
p1 = plot_ecdf_diff(df.results.b) + theme_bw() + theme(legend.position = "none") +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p2 = plot_rank_hist(df.results.b, bins = 20) + theme_bw() +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p3 = plot_sim_estimated(df.results.b, alpha = 0.5) + theme_bw() +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p4 = plot_contraction(df.results.b,
                      prior_sd = setNames(
                        c(as.numeric(
                          gsub(".*, (.+)\\).*", "\\1",
                               priors[priors$class == "Intercept",]$prior)),
                          as.numeric(
                            gsub(".*, (.+)\\).*", "\\1",
                                 priors[priors$class == "b",]$prior))),
                                          unique(df.results.b$variable))) +
  theme_bw() +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p = ggarrange(p1, p2, p3, p4, labels = "AUTO", ncol = 1, nrow = 4)
annotate_figure(p, top =
                  text_grob("Computational faithfulness and model sensitivity",
                face = "bold", size = 14))
```



Next, we check the ranks of the parameters. If the model is unbiased, these should be uniformly distributed

(Schad, Betancourt and Vasishth, 2020). The sample empirical cumulative distribution function (ECDF) lies within the theoretical distribution (95%) and the rank histogram also shows ranks within the 95% expected range, although there are some small deviations. We judge this to be acceptable.

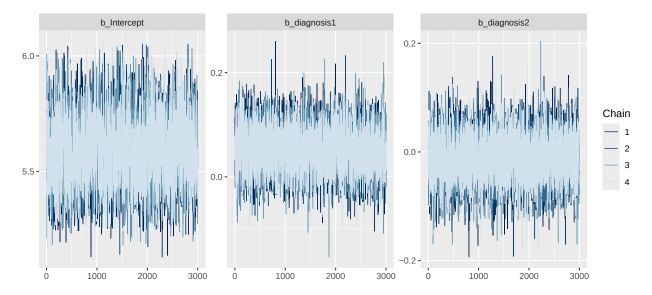
Then, we investigated the relationship between the simulated true parameters and the posterior estimates. Although there are individual values diverging from the expected pattern, most parameters were recovered successfully within an uncertainty interval of alpha = 0.05.

Last, we explore the z-score and the posterior contraction of our population-level predictors. The z-score "determines the distance of the posterior mean from the true simulating parameter", while the posterior contraction "estimates how much prior uncertainty is reduced in the posterior estimation" (Schad, Betancourt and Vasisth, 2020). All of this looks good for this model.

Posterior predictive checks

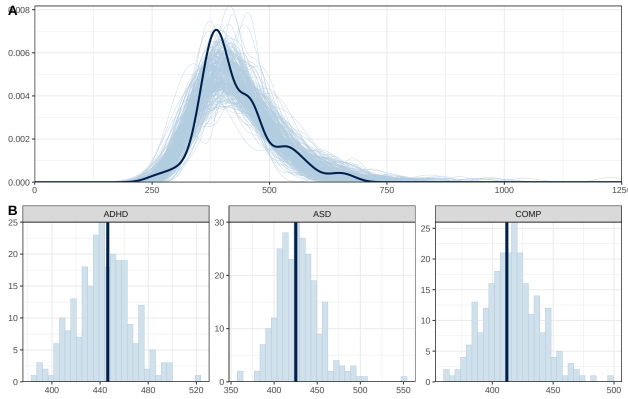
As the next step, we fit the model to the data, check whether there are divergence or rhat issues, and then check whether the chains have converged.

```
# fit the aggregated model
set.seed(2469)
m.rtc = brm(f.rtc,
            df.rtc.agg, prior = priors,
            family = shifted_lognormal,
            iter = iter, warmup = warm,
            backend = "cmdstanr", threads = threading(8),
            file = file.path(model dir, "m rtc agg")
rstan::check hmc diagnostics(m.rtc$fit)
## Divergences:
## 0 of 12000 iterations ended with a divergence.
##
## Tree depth:
## 0 of 12000 iterations saturated the maximum tree depth of 10.
## Energy:
## E-BFMI indicated no pathological behavior.
# check that rhats are below 1.01
sum(brms::rhat(m.rtc) >= 1.01, na.rm = T)
## [1] 0
# check the trace plots
post.draws = as_draws_df(m.rtc)
mcmc_trace(post.draws, regex_pars = "^b_",
           facet_args = list(ncol = 3)) +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
## Scale for x is already present.
## Adding another scale for x, which will replace the existing scale.
```



This model has no pathological behaviour with E-BFMI, no divergent samples and no rhat that is higher or equal to 1.01. Therefore, we go ahead and perform our posterior predictive checks.

Posterior predictive checks: RTs



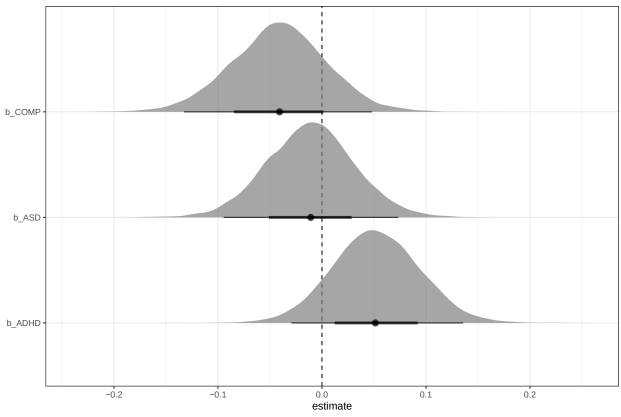
This looks much better, especially when we look at the simulated means (light blue) of the three diagnostic groups in comparison to the actual means (dark blue) which is important because we draw our inferences based on the estimates of the diagnostic groups. Therefore, we can finally move on to our inferences based on the model.

Inferences

Now that we are convinced that we can trust our model, we have a look at its estimate and use the hypothesis function to explore group differences.

```
# print a summary
summary(m.rtc)
    Family: shifted_lognormal
     Links: mu = identity; sigma = identity; ndt = identity
##
## Formula: rtc ~ diagnosis
##
      Data: df.rtc.agg (Number of observations: 67)
     Draws: 4 chains, each with iter = 4500; warmup = 1500; thin = 1;
##
##
            total post-warmup draws = 12000
##
## Regression Coefficients:
              Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
                             0.15
                  5.60
                                      5.32
                                               5.91 1.00
                                                              3304
                                                                       3077
## Intercept
  diagnosis1
                  0.05
                             0.04
                                     -0.03
                                               0.14 1.00
                                                              6186
                                                                       6527
                             0.04
##
  diagnosis2
                 -0.01
                                     -0.09
                                               0.07 1.00
                                                              5902
                                                                       5970
##
## Further Distributional Parameters:
##
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
```

```
0.28
                      0.05
                                         0.39 1.00
## sigma
                               0.19
                                                        3454
                                                                 3957
## ndt
           145.29
                      40.12
                               52.77
                                       209.99 1.00
                                                       3231
                                                                 3147
##
## Draws were sampled using sample(hmc). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
# get the estimates and compute groups
df.m.rtc = as_draws_df(m.rtc) %>%
  select(starts_with("b_")) %>%
  mutate(
   b COMP
              = - b_diagnosis1 - b_diagnosis2,
   b_ASD
                = b_diagnosis2,
   b_ADHD
                = b_diagnosis1
    )
# plot the posterior distributions
df.m.rtc %>%
  select(b_ASD, b_ADHD, b_COMP) %>%
  pivot_longer(cols = c(b_ASD, b_ADHD, b_COMP), names_to = "coef", values_to = "estimate") %>%
 group_by(coef) %>%
 mutate(
    cred = case when(
      (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |</pre>
        (mean(estimate) > 0 & quantile(estimate, probs = 0.025) > 0) ~ "credible",
     T ~ "not credible"
   )
  ) %>% ungroup() %>%
  ggplot(aes(x = estimate, y = coef, fill = cred)) +
  geom_vline(xintercept = 0, linetype = 'dashed') +
  ggdist::stat_halfeye(alpha = 0.7) + ylab(NULL) + theme_bw() +
  scale_fill_manual(values = c(credible = c_dark, c_light)) +
  theme(legend.position = "none")
```



```
# ADHD slower than COMP
e1 = hypothesis(m.rtc, "0 < 2*diagnosis1 + diagnosis2", alpha = 0.025)
## Hypothesis Tests for class b:
                  Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio
## 1 (0)-(2*diagnosis1... < 0
                                -0.09
                                          0.08
                                                    -0.25
                                                              0.06
                                                                         8.28
##
    Post.Prob Star
## 1
         0.89
## ---
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.
## '*': For one-sided hypotheses, the posterior probability exceeds 97.5%;
## for two-sided hypotheses, the value tested against lies outside the 97.5%-CI.
## Posterior probabilities of point hypotheses assume equal prior probabilities.
# ASD slower than COMP
e2 = hypothesis(m.rtc, "0 < 2*diagnosis2 + diagnosis1", alpha = 0.025)
## Hypothesis Tests for class b:
                  Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio
## 1 (0)-(2*diagnosis2... < 0 -0.03 0.08
                                                   -0.18
                                                              0.12
                                                                         1.95
    Post.Prob Star
## 1
         0.66
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.
\#\# '*': For one-sided hypotheses, the posterior probability exceeds 97.5%;
## for two-sided hypotheses, the value tested against lies outside the 97.5%-CI.
## Posterior probabilities of point hypotheses assume equal prior probabilities.
```

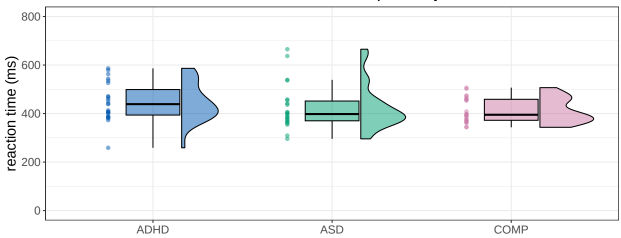
```
# extract predicted differences in ms instead of log data
df.new = df.rtc %>%
  select(diagnosis) %>%
  distinct()
df.ms = as.data.frame(
  fitted(m.rtc, summary = F,
               newdata = df.new %>% select(diagnosis),
               re_formula = NA))
colnames(df.ms) = df.new$diagnosis
# calculate our difference columns
df.ms = df.ms %>%
  mutate(
    COMP\_ADHD = COMP - ADHD,
              = COMP - ASD
   COMP_ASD
 )
```

Our Bayesian linear mixed model with the hit reaction times as the outcome and diagnostic status as a predictor showed no credible differences: COMP participants reacted similarly to the ADHD group (CI of COMP - ADHD: -68.31 to 16.15ms, posterior probability = 89.22%) and the ASD group (CI of COMP - ASD: -49.3 to 33.53ms, posterior probability = 66.11%).

Plots

```
# overall median reaction times
df.rtc.agg %>%
 ggplot(aes(diagnosis, rtc, fill = diagnosis, colour = diagnosis)) + #
  geom_rain(rain.side = 'r',
boxplot.args = list(color = "black", outlier.shape = NA, show_guide = FALSE, alpha = 0.5),
violin.args = list(color = "black", outlier.shape = NA, alpha = 0.5),
boxplot.args.pos = list(
  position = ggpp::position_dodgenudge(x = 0, width = 0.3), width = 0.3
point.args = list(show_guide = FALSE, alpha = .5),
violin.args.pos = list(
  width = 0.6, position = position_nudge(x = 0.16)),
point.args.pos = list(position = ggpp::position_dodgenudge(x = -0.25, width = 0.1))) +
 ylim(0, 800) +
  scale fill manual(values = custom.col) +
  scale_color_manual(values = custom.col) +
  labs(title = "Median reaction times per subject",
      x = "",
      y = "reaction time (ms)") +
  theme_bw() +
  theme(legend.position = "none",
        plot.title = element_text(hjust = 0.5),
       legend.direction = "horizontal",
        text = element_text(size = 15))
```

Median reaction times per subject



Analysis of discrimination rate

SBC with group-level intercept for subjects

```
code = "VMM_disc"
# model formula
f.disc = brms::bf(negdisc ~ diagnosis + (1 | subID))
# use more iterations and warmup since the sample size is smaller
warm = warm * 2
iter = iter * 2
# set weakly informed priors
priors = c(
  # expect high discrimination rates, therefore, low divergences
  prior(normal(3, 1.5), class = Intercept), # ~ 5 - 90 (+- 1 SD)
  prior(normal(0,
                  1.5), class = sd),
  # no particular expectations for effects
                   1.0), class = b)
  prior(normal(0,
# check if the SBC already exists
if (file.exists(file.path(cache_dir, sprintf("df_res_%s.rds", code)))) {
  # load in the results of the SBC
  df.results = readRDS(file.path(cache_dir, sprintf("df_res_%s.rds", code)))
  df.backend = readRDS(file.path(cache_dir, sprintf("df_div_%s.rds", code)))
  dat
             = readRDS(file.path(cache_dir, sprintf("dat_%s.rds", code)))
} else {
  # set the seed
  set.seed(2468)
  # perform the SBC
  gen = SBC_generator_brms(f.disc, data = df.disc, prior = priors,
                           family = poisson,
                           thin = 50, warmup = 20000, refresh = 2000)
```

```
dat = generate_datasets(gen, nsim)
  saveRDS(dat, file.path(cache_dir, sprintf("dat_%s.rds", code)))
  bck = SBC_backend_brms_from_generator(gen, chains = 4, thin = 1,
                                      init = 0.1, warmup = warm, iter = iter)
  # set the seed again
  set.seed(2468)
  res = compute_SBC(dat,
        bck,
        cache mode
                     = "results",
        cache_location = file.path(cache_dir, sprintf("res_%s", code)))
  df.results = res$stats
  df.backend = res$backend_diagnostics
  saveRDS(df.results, file = file.path(cache_dir, paste0("df_res_", code, ".rds")))
  saveRDS(df.backend, file = file.path(cache_dir, paste0("df_div_", code, ".rds")))
}
```

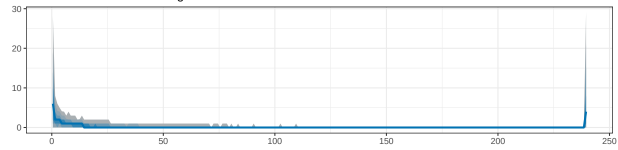
Again, we start by investigating the rhats and the number of divergent samples. This shows that 6 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and 7 models had divergent samples. This suggests that this model performs well and we can continue with our checks by plotting the simulated values to perform prior predictive checks.

```
# create a matrix out of generated data
dvname = gsub(" ", "", gsub("[\\|~].*", "", f.disc)[1])
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))
for (i in 1:length(dat[['generated']])) {
  dvfakemat[,i] = dat[['generated']][[i]][[dvname]]
}
truePars = dat$variables
dvfakemat[dvfakemat > 240] = 240
\# compute one histogram per simulated data-set
binwidth = 1
breaks = seq(0, ceiling(max(dvfakemat, na.rm=T)), binwidth)
histmat = matrix(NA, ncol = length(dat), nrow = length(breaks)-1)
for (i in 1:nrow(truePars)) {
 histmat[,i] = hist(dvfakemat[,i], breaks = breaks, plot = F)$counts
}
# for each bin, compute quantiles across histograms
probs = seq(0.1, 0.9, 0.1)
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))
names(quantmat) = paste0("p", probs)
for (i in 1:dim(histmat)[1]) {
  quantmat[i,] = quantile(histmat[i,], p = probs)
quantmat$x = breaks[2:length(breaks)] - binwidth/2 # add bin mean
p1 = ggplot(data = quantmat, aes(x = x)) +
  geom_ribbon(aes(ymax = p0.9, ymin = p0.1), fill = c_light) +
  geom_ribbon(aes(ymax = p0.8, ymin = p0.2), fill = c_light_highlight) +
  geom_ribbon(aes(ymax = p0.7, ymin = p0.3), fill = c_mid) +
  geom_ribbon(aes(ymax = p0.6, ymin = p0.4), fill = c_mid_highlight) +
  geom_line(aes(y = p0.5), colour = c_dark, linewidth = 1) +
  labs(title = "Distribution of simulated negative discriminations", y = "", x = "") +
  theme_bw()
```

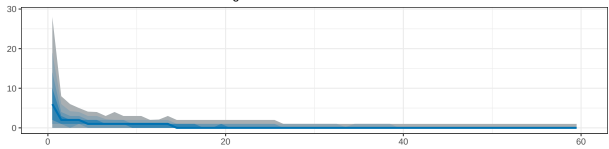
```
p2 = ggplot(data = quantmat, aes(x = x)) +
  geom_ribbon(aes(ymax = p0.9, ymin = p0.1), fill = c_light) +
  geom_ribbon(aes(ymax = p0.8, ymin = p0.2), fill = c_light_highlight) +
  geom_ribbon(aes(ymax = p0.7, ymin = p0.3), fill = c_mid) +
  geom_ribbon(aes(ymax = p0.6, ymin = p0.4), fill = c_mid_highlight) +
  geom_line(aes(y = p0.5), colour = c_dark, linewidth = 1) +
  xlim(0, 60) +
  labs(title = "Zoomed in distribution of simulated negative discriminations", y = "", x = "") +
  theme_bw()
tmpM = apply(dvfakemat, 2, mean) # mean
tmpSD = apply(dvfakemat, 2, sd)
p3 = ggplot() +
  stat_bin(aes(x = tmpM), fill = c_dark) +
 labs(x = "Mean RTs (ms)", title = "Means of simulated RTs") +
 theme_bw()
p4 = ggplot() +
  stat_bin(aes(x = tmpSD), fill = c_dark) +
  labs(x = "SD RTs (ms)", title = "SDs of simulated RTs") +
 theme_bw()
p = ggarrange(p1, p2,
  ggarrange(p3, p4, ncol = 2, labels = c("B", "C")),
  nrow = 3, labels = "A")
annotate_figure(p,
                top = text_grob("Prior predictive checks: reaction times",
                face = "bold", size = 14))
```

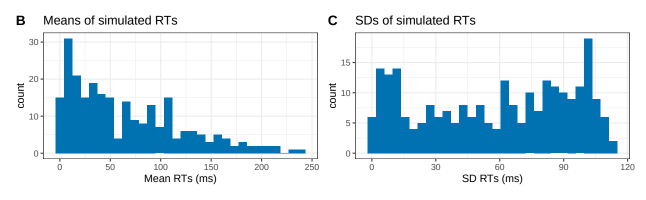
Prior predictive checks: reaction times

A Distribution of simulated negative discriminations



Zoomed in distribution of simulated negative discriminations



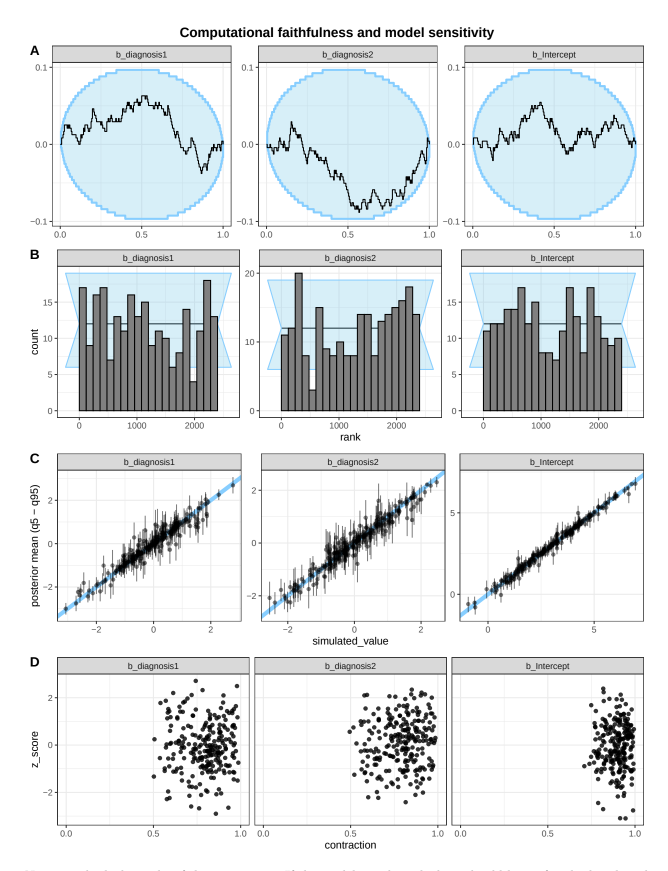


Subfigure A shows the distribution of the simulated data with bluer bands being more likely than greyer bands. Subfigure A shows a distribution that fits our expectations about the negative discrimination (perfect discrimination - discrimination rate) in a simple decision task, with most participants showing low rates of negative discrimination. If we zoom in, most of the data is indeed in the very low numbers. The distribution of the means and standard deviations in the simulated datasets also look good. We go ahead with these priors and check the results of the SBC. We only plot the results from the models that had no divergence issues.

```
# get simulation numbers with issues
rank = max(df.results$max_rank)
check = merge(df.results %>%
    group_by(sim_id) %>%
    summarise(
        rhat = max(rhat, na.rm = T),
        mean_rank = mean(max_rank)
        ) %>%
    filter(rhat >= 1.05 | mean_rank != rank),
    df.backend %>% filter(n_divergent > 0), all = T)

# plot SBC with functions from the SBC package focusing on population-level parameters
```

```
df.results.b = df.results %>%
  filter(substr(variable, 1, 2) == "b_") %>%
  filter(!(sim_id %in% check$sim_id))
p1 = plot ecdf diff(df.results.b) + theme bw() + theme(legend.position = "none") +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p2 = plot_rank_hist(df.results.b, bins = 20) + theme_bw() +
  scale x continuous(breaks=scales::pretty breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p3 = plot_sim_estimated(df.results.b, alpha = 0.5) + theme_bw() +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p4 = plot_contraction(df.results.b,
                      prior_sd = setNames(
                        c(as.numeric(
                          gsub(".*, (.+)\\).*", "\\1",
                               priors[priors$class == "Intercept",]$prior)),
                          as.numeric(
                            gsub(".*, (.+)\\).*", "\\1",
                                 priors[priors$class == "b",]$prior)),
                          as.numeric(
                            gsub(".*, (.+)\\).*", "\\1",
                                 priors[priors$class == "b",]$prior))),
                                          unique(df.results.b$variable))) +
  theme bw() +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
p = ggarrange(p1, p2, p3, p4, labels = "AUTO", ncol = 1, nrow = 4)
annotate_figure(p, top =
                  text_grob("Computational faithfulness and model sensitivity",
                face = "bold", size = 14))
```



Next, we check the ranks of the parameters. If the model is unbiased, these should be uniformly distributed

(Schad, Betancourt and Vasishth, 2020). The sample empirical cumulative distribution function (ECDF) lies within the theoretical distribution (95%) and the rank histogram also shows ranks within the 95% expected range, although there are some small deviations. We judge this to be acceptable.

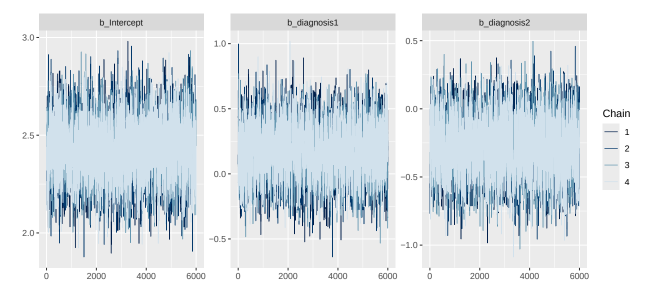
Then, we investigated the relationship between the simulated true parameters and the posterior estimates. Although there are individual values diverging from the expected pattern, most parameters were recovered successfully within an uncertainty interval of alpha = 0.05.

Last, we explore the z-score and the posterior contraction of our population-level predictors. The z-score "determines the distance of the posterior mean from the true simulating parameter", while the posterior contraction "estimates how much prior uncertainty is reduced in the posterior estimation" (Schad, Betancourt and Vasisth, 2020). All of this looks good for this model.

Posterior predictive checks

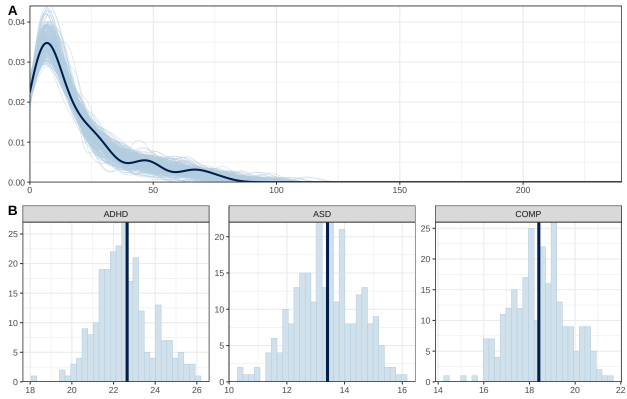
As the next step, we fit the model to the data, check whether there are divergence or rhat issues, and then check whether the chains have converged.

```
# fit the aggregated model
set.seed(2469)
m.disc = brm(f.disc,
            df.disc, prior = priors,
            family = poisson,
            iter = iter, warmup = warm,
            backend = "cmdstanr", threads = threading(8),
            file = file.path(model dir, "m disc")
rstan::check hmc diagnostics(m.disc$fit)
## Divergences:
## 0 of 24000 iterations ended with a divergence.
##
## Tree depth:
## 0 of 24000 iterations saturated the maximum tree depth of 10.
## Energy:
## E-BFMI indicated no pathological behavior.
# check that rhats are below 1.01
sum(brms::rhat(m.disc) >= 1.01, na.rm = T)
## [1] 0
# check the trace plots
post.draws = as_draws_df(m.disc)
mcmc_trace(post.draws, regex_pars = "^b_",
           facet_args = list(ncol = 3)) +
  scale_x_continuous(breaks=scales::pretty_breaks(n = 3)) +
  scale_y_continuous(breaks=scales::pretty_breaks(n = 3))
## Scale for x is already present.
## Adding another scale for x, which will replace the existing scale.
```



This model has no pathological behaviour with E-BFMI, no divergent samples and no rhat that is higher or equal to 1.01. Therefore, we go ahead and perform our posterior predictive checks.

Posterior predictive checks: Discrimination rate



The predicted values based on the model in light blue capture the actual data in dark blue quite well, both in terms of overall data (A) and aggregated values for the diagnostic groups (B). Therefore, we can finally move on to our inferences based on the model.

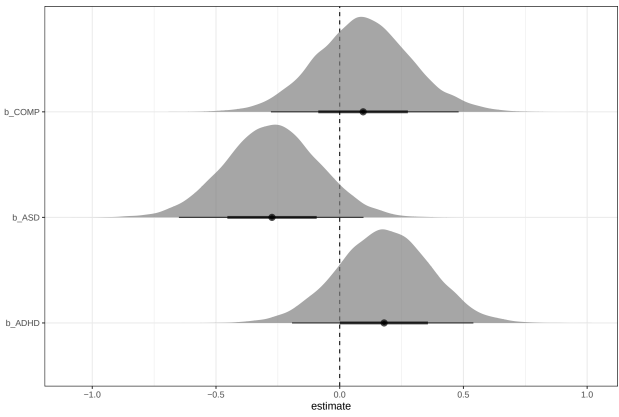
Inferences

Now that we are convinced that we can trust our model, we have a look at its estimate and use the hypothesis function to explore group differences.

```
# print a summary
summary(m.disc)
```

```
Family: poisson
##
     Links: mu = log
##
## Formula: negdisc ~ diagnosis + (1 | subID)
      Data: df.disc (Number of observations: 67)
##
##
     Draws: 4 chains, each with iter = 9000; warmup = 3000; thin = 1;
            total post-warmup draws = 24000
##
##
## Multilevel Hyperparameters:
  ~subID (Number of levels: 67)
##
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                                0.11
                                         0.85
                                                   1.28 1.00
                                                                 4753
                                                                           8889
## sd(Intercept)
                      1.04
##
## Regression Coefficients:
              Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                                                              4101
## Intercept
                  2.43
                             0.14
                                      2.16
                                                2.69 1.00
                                                                        7578
## diagnosis1
                  0.18
                             0.19
                                     -0.19
                                               0.54 1.00
                                                              3261
                                                                        6317
```

```
0.19
                                    -0.65
                                              0.10 1.00
                                                            3529
## diagnosis2
                -0.27
                                                                      6414
##
## Draws were sampled using sample(hmc). For each parameter, Bulk ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
# get the estimates and compute groups
df.m.disc = as_draws_df(m.disc) %>%
 select(starts_with("b_")) %>%
 mutate(
   b_COMP
              = - b_diagnosis1 - b_diagnosis2,
               = b_diagnosis2,
   b ASD
   b_ADHD
                = b_diagnosis1
# plot the posterior distributions
df.m.disc %>%
  select(b_ASD, b_ADHD, b_COMP) %>%
  pivot_longer(cols = c(b_ASD, b_ADHD, b_COMP), names_to = "coef", values_to = "estimate") %>%
  group_by(coef) %>%
  mutate(
   cred = case_when(
      (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |</pre>
        (mean(estimate) > 0 & quantile(estimate, probs = 0.025) > 0) ~ "credible",
     T ~ "not credible"
   )
  ) %>% ungroup() %>%
  ggplot(aes(x = estimate, y = coef, fill = cred)) +
  geom_vline(xintercept = 0, linetype = 'dashed') +
  ggdist::stat_halfeye(alpha = 0.7) + ylab(NULL) + theme_bw() +
  scale_fill_manual(values = c(credible = c_dark, c_light)) +
  theme(legend.position = "none")
```



```
# ADHD worse discrimination than COMP
e1 = hypothesis(m.disc, "0 < 2*diagnosis1 + diagnosis2", alpha = 0.025)
e1
## Hypothesis Tests for class b:
                  Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio
## 1 (0)-(2*diagnosis1... < 0
                                -0.08
                                           0.33
                                                    -0.72
                                                              0.58
                                                                         1.53
##
    Post.Prob Star
## 1
          0.6
## ---
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.
## '*': For one-sided hypotheses, the posterior probability exceeds 97.5%;
## for two-sided hypotheses, the value tested against lies outside the 97.5%-CI.
## Posterior probabilities of point hypotheses assume equal prior probabilities.
# ASD worse discrimination than COMP
e2 = hypothesis(m.disc, "0 > 2*diagnosis2 + diagnosis1", alpha = 0.025)
e2
## Hypothesis Tests for class b:
                  Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio
## 1 (0)-(2*diagnosis2... > 0
                                                    -0.28
                                                              1.03
                                                                         6.62
                                0.37
                                       0.33
    Post.Prob Star
## 1
         0.87
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.
\#\# '*': For one-sided hypotheses, the posterior probability exceeds 97.5%;
## for two-sided hypotheses, the value tested against lies outside the 97.5%-CI.
## Posterior probabilities of point hypotheses assume equal prior probabilities.
```

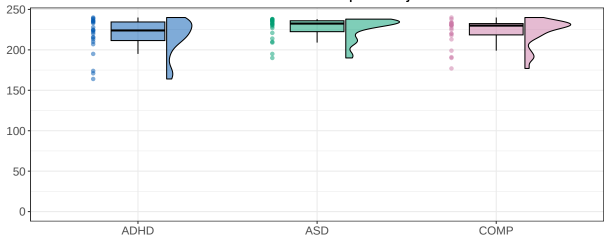
```
# extract predicted differences in ms instead of log data
df.new = df.disc %>%
  select(diagnosis) %>%
  distinct()
df.ms = as.data.frame(
  fitted(m.disc, summary = F,
               newdata = df.new %>% select(diagnosis),
               re_formula = NA))
colnames(df.ms) = df.new$diagnosis
# calculate our difference columns
df.ms = df.ms %>%
  mutate(
    COMP\_ADHD = COMP - ADHD,
   COMP_ASD
              = COMP - ASD
 )
```

Our Bayesian linear mixed model with the negative discrimination rate (perfect discrimination rate - actual discrimination rate) as the outcome and diagnostic status as a predictor showed no credible differences: COMP participants reacted similarly to the ADHD group (CI of COMP - ADHD: -9.72 to 7.92ms, posterior probability = 60.45%) and the ASD group (CI of COMP - ASD: -2.97 to 11.87ms, posterior probability = 86.88%).

Plots

```
# overall median reaction times
df.disc %>%
  ggplot(aes(diagnosis, disc, fill = diagnosis, colour = diagnosis)) + #
  geom_rain(rain.side = 'r',
boxplot.args = list(color = "black", outlier.shape = NA, show_guide = FALSE, alpha = 0.5),
violin.args = list(color = "black", outlier.shape = NA, alpha = 0.5),
boxplot.args.pos = list(
  position = ggpp::position_dodgenudge(x = 0, width = 0.3), width = 0.3
point.args = list(show_guide = FALSE, alpha = .5),
violin.args.pos = list(
  width = 0.6, position = position_nudge(x = 0.16)),
point.args.pos = list(position = ggpp::position_dodgenudge(x = -0.25, width = 0.1))) +
 ylim(0, 240) +
  scale_fill_manual(values = custom.col) +
  scale_color_manual(values = custom.col) +
  labs(title = "Discrimination rate per subject",
      x = ""
      y = "") +
  theme bw() +
  theme(legend.position = "none",
       plot.title = element_text(hjust = 0.5),
        legend.direction = "horizontal",
       text = element_text(size = 15))
```





Analysis of fixation proportions to centre AOI

Bayesian ANOVAs

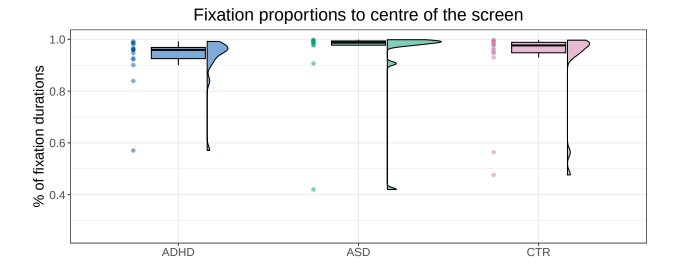
```
# check which outcomes of interest are normally distributed
df.fix.agg %>%
  group_by(diagnosis) %>%
  rstatix::shapiro_test(fix.total, fix.prop, rfix.total, rfix.prop) %>%
  mutate(
    sig = if_else(p < 0.05, "*", "")
 ) %>% arrange(variable)
## # A tibble: 12 x 5
##
      diagnosis variable
                           statistic
                                                p sig
##
      <fct>
                <chr>
                               <dbl>
                                            <dbl> <chr>
##
   1 ADHD
                fix.prop
                               0.580 0.0000107
                                                  "*"
                               0.443 0.000000317 "*"
  2 ASD
##
                fix.prop
  3 CTR
                fix.prop
                               0.538 0.00000725
                                                  "*"
## 4 ADHD
                fix.total
                               0.905 0.0954
## 5 ASD
                fix.total
                               0.784 0.00586
                                                  "*"
                               0.656 0.0000864
##
  6 CTR
                fix.total
                               0.947 0.449
                                                  11 11
  7 ADHD
                rfix.prop
## 8 ASD
                rfix.prop
                               0.880 0.104
                                                  11 11
## 9 CTR
                rfix.prop
                               0.949 0.510
## 10 ADHD
                rfix.total
                               0.898 0.0738
                               0.902 0.195
                                                  11 11
## 11 ASD
                rfix.total
## 12 CTR
                rfix.total
                               0.948 0.491
# ANOVA for the ranked proportional fixation durations
aov.fix = BayesFactor::anovaBF(rfix.prop ~ diagnosis, data = df.fix.agg)
aov.fix@bayesFactor[["bf"]]
## [1] -0.04049587
effectsize::interpret_bf(aov.fix@bayesFactor[["bf"]], log = T)
```

Table 4: Summary Statistics

diagnosis	ADHD		ASD		CTR				
Variable	N	Mean	SD	N	Mean	SD	N	Mean	SD
fix.centre	16	1040452	515022	11	1293957	467256	15	1308761	516135
fix.periphery	16	68752	74349	11	73310	172166	15	73488	132065
fix.total	16	1109204	513172	11	1367266	376425	15	1382249	495608
fix.prop	16	0.93	0.1	11	0.93	0.17	15	0.91	0.16

Plots

```
# overall
df.fix.agg %>%
  ggplot(aes(diagnosis, fix.prop, fill = diagnosis, colour = diagnosis)) + #
  geom_rain(rain.side = 'r',
boxplot.args = list(color = "black", outlier.shape = NA, show_guide = FALSE, alpha = 0.5),
violin.args = list(color = "black", outlier.shape = NA, alpha = 0.5),
boxplot.args.pos = list(
  position = ggpp::position_dodgenudge(x = 0, width = 0.3), width = 0.3
point.args = list(show_guide = FALSE, alpha = .5),
violin.args.pos = list(
  width = 0.6, position = position_nudge(x = 0.16)),
point.args.pos = list(position = ggpp::position_dodgenudge(x = -0.25, width = 0.1))) +
  ylim(0.25, 1) +
  scale_fill_manual(values = custom.col) +
  scale_color_manual(values = custom.col) +
  labs(title = "Fixation proportions to centre of the screen",
       y = "% of fixation durations") +
  theme bw() +
  theme(legend.position = "none",
        plot.title = element_text(hjust = 0.5),
        legend.direction = "horizontal",
```



${\bf Heatmaps}$

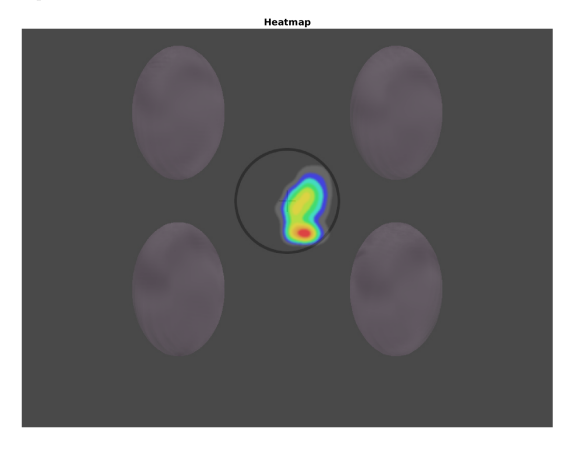


Figure 1: Heatmap of gaze pattern of the COMP group

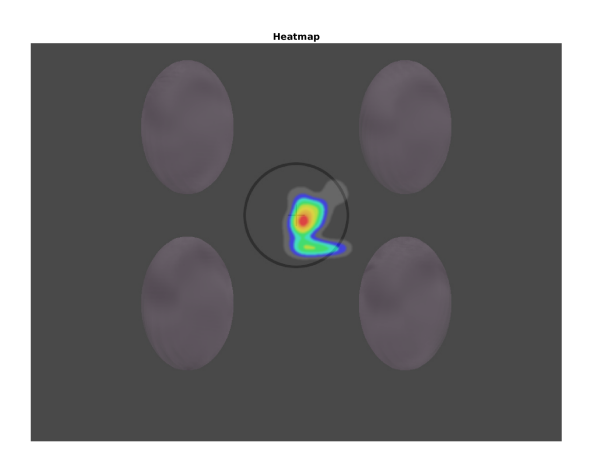


Figure 2: Heatmap of gaze pattern of the ADHD group

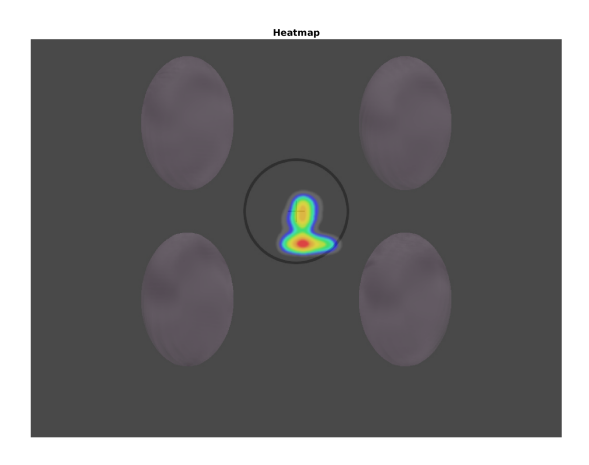


Figure 3: Heatmap of gaze pattern of the ASD group

Explorative analysis of RT costs of changes

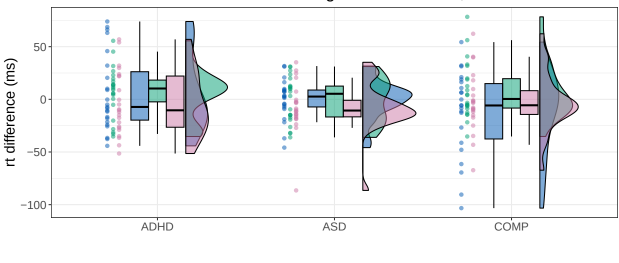
```
# calculate rt difference from early to late fixation cross switches within a block
df.rtc.change = df.rtc %>%
 mutate(
   switch = case when(
     time_since_c < 2*750 ~ "early",</pre>
     time_since_c \geq= 2*750 & time_since_c < 5*750 ~ "middle",
     T ~ "late"
  ) %>% filter(switch != "late") %>%
  group_by(subID, diagnosis, switch, which_change) %>%
  summarise(
   rtc.avg = mean(rtc, na.rm = T)
  ) %>%
  pivot_wider(values_from = rtc.avg, names_from = switch) %>%
  mutate(
   rt.diff = middle - early
df.rtc.change %>%
  group_by(which_change) %>%
  shapiro test(rt.diff)
## # A tibble: 3 x 4
    which_change variable statistic
##
     <fct>
                 <chr> <dbl> <dbl>
                 rt.diff
## 1 both
                               0.969 0.0876
## 2 colour
                               0.973 0.162
                 rt.diff
                               0.969 0.0954
## 3 emo
                  rt.diff
# run an ANOVA
aov.diff = anovaBF(rt.diff ~ diagnosis*which_change, data = df.rtc.change)
aov.diff
## Bayes factor analysis
## [1] diagnosis
                                                          : 0.1039953
                                                                       ±0.02%
## [2] which_change
                                                          : 0.2900511
                                                                       ±0.01%
## [3] diagnosis + which_change
                                                         : 0.02973831 ±1.07%
## [4] diagnosis + which_change + diagnosis:which_change : 0.005065659 ±1.93%
## Against denominator:
    Intercept only
## ---
## Bayes factor type: BFlinearModel, JZS
aov.diff@bayesFactor$bf
```

```
## [1] -2.263410 -1.237698 -3.515319 -5.285271
```

None of the models outperform the intercept model. This suggests that the difference in reaction time between early and middle switches were not dependent on the diagnosis, the type of the change (emotion, colour or both) or their interaction. Let's still plot the differences.

```
df.rtc.change %>%
  ggplot(aes(diagnosis, rt.diff, fill = which_change, colour = which_change)) + #
  geom_rain(rain.side = 'r',
boxplot.args = list(color = "black", outlier.shape = NA, show_guide = FALSE, alpha = 0.5),
violin.args = list(color = "black", outlier.shape = NA, alpha = 0.5),
boxplot.args.pos = list(
 position = ggpp::position_dodgenudge(x = 0, width = 0.3), width = 0.3
point.args = list(show_guide = FALSE, alpha = .5),
violin.args.pos = list(
  width = 0.6, position = position_nudge(x = 0.16)),
point.args.pos = list(position = ggpp::position_dodgenudge(x = -0.25, width = 0.1))) +
  scale_fill_manual(values = custom.col) +
  scale_color_manual(values = custom.col) +
 labs(title = "Reaction time costs due to changes in face colour, emotion or both",
      x = ""
       y = "rt difference (ms)") +
  theme_bw() +
  theme(legend.position = "bottom",
       plot.title = element_text(hjust = 0.5),
        legend.direction = "horizontal",
       text = element_text(size = 15))
```

Reaction time costs due to changes in face colour, emotion or both



which change both colour emo