S2: Analysis of FAB eye tracking data with brms

I. S. Plank

2025-01-31

# S2.1 Introduction

This R Markdown script analyses eye tracking data from the FAB (face attention bias) paradigm of the EMBA project. The data was preprocessed before being read into this script.

## Some general settings

# number of simulations  
nsim = 250  
  
# set the seed  
set.seed(2468)

## Package versions

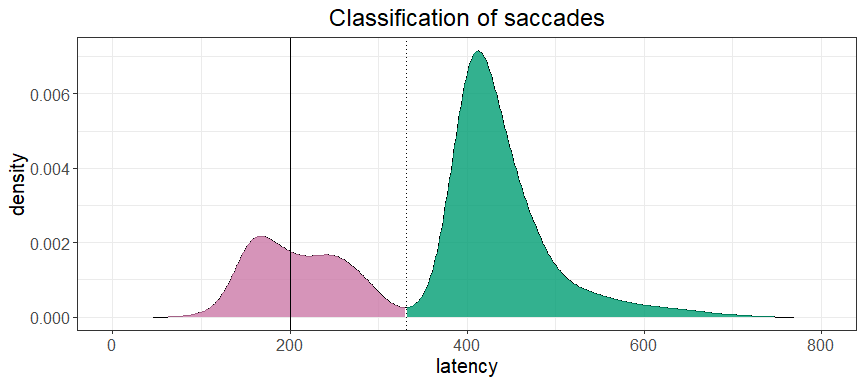
## [1] "R version 4.3.0 (2023-04-21 ucrt)"

## [1] "knitr version 1.42"  
## [1] "ggplot2 version 3.4.2"  
## [1] "brms version 2.19.0"  
## [1] "tidyverse version 2.0.0"  
## [1] "ggpubr version 0.6.0"  
## [1] "ggrain version 0.0.4"  
## [1] "bayesplot version 1.10.0"  
## [1] "SBC version 0.2.0.9000"  
## [1] "rstatix version 0.7.2"  
## [1] "logspline version 2.1.22"  
## [1] "BayesFactor version 0.9.12.4.4"  
## [1] "bayestestR version 0.15.1"

## Preparation

First, we load the eye tracking data and combine it with demographic information including the diagnostic status of the subjects. Second, we preprocess the latencies of the saccades and divide them into saccades elicited by the cues and saccades elicited by the target. To do so, we use the knowledge that latencies below 100ms are extremely unlikely and use the global minimum in the density function. We also load in the behavioural data again to be able to use reaction times for further correlational analyses.

# load the data  
load("FAB\_data.RData")  
  
# combine both behavioural datasets  
df.fab = rbind(df.fab, df.exp)  
df.fab$diagnosis = factor(df.fab$diagnosis,   
 levels = c("ADHD", "ASD", "BOTH", "COMP"))  
  
# aggregate the behavioural data disregarding stimuli and trials  
df.fab.agg = rbind(df.fab, df.exp) %>%  
 group\_by(subID, diagnosis, cue) %>%  
 summarise(  
 rt.cor = median(rt.cor, na.rm = T)  
 ) %>%  
 group\_by(subID, diagnosis) %>%  
 # calculate subject-specific difference between cues  
 summarise(  
 rt.fab = combn(rt.cor, 2, FUN = function(x) x[1] - x[2])  
 )   
  
# remove participants without any data  
df.sac = df.sac %>% filter(!is.na(lat))  
  
# remove unbelievably short and extremely long saccades  
df.sac = df.sac %>%  
 filter(lat <= quantile(df.sac$lat, probs = 0.99) &  
 lat > 100)  
  
# divide into cue and target saccades  
criticalpoints = function(density, threshold = 1){  
 up = sapply(1:threshold, function(n) c(density$y[-(seq(n))], rep(NA, n)))  
 down = sapply(-1:-threshold,   
 function(n) c(rep(NA,abs(n)),   
 density$y[-seq(length(density$y),   
 length(density$y) - abs(n) + 1)]))  
 a = cbind(density$y,up,down)  
 minima = round(density$x[which(apply(a, 1, min) == a[,1])])  
 maxima = round(density$x[which(apply(a, 1, max) == a[,1])])  
 return(list(minima = minima, maxima = maxima))  
}  
  
points = criticalpoints(density(df.sac$lat))  
  
# get the density of the latencies  
dd = with(density(df.sac$lat), data.frame(x,y))  
  
# find which point is the global minimum  
lat.points = dd$y[points$minima]  
idx = which.min(lat.points)  
  
ggplot(dd, aes(x = x, y = y)) +   
 geom\_line() +  
 geom\_vline(xintercept = points$minima[idx], linetype=3) +  
 geom\_ribbon(data = subset(dd, x <= points$minima[idx]), aes(ymax = y), ymin = 0,   
 fill = custom.col[2], colour = NA, alpha = .8) +  
 geom\_ribbon(data = subset(dd, x >= points$minima[idx]), aes(ymax = y), ymin = 0,   
 fill = custom.col[3], colour = NA, alpha = .8) +   
 geom\_vline(xintercept = 200) +   
 labs(title = "Classification of saccades", x = "latency", y = "density") +  
 xlim(0, 800) +  
 theme\_bw() +   
 theme(legend.position = "bottom",   
 plot.title = element\_text(hjust = 0.5),   
 legend.direction = "horizontal",   
 text = element\_text(size = 15))



ggsave("Fig2\_densLatency.pdf",   
 units = "mm",   
 width = 170,  
 height = 70,  
 dpi = 300)

The graph above shows the density of the latencies with zero on the x-axis being the onset of the cue. After 200ms, the cue disappears and the target is presented on the screen (solid line). We can see that there is a minimum about 130ms after the target appears (dotted line). We can assume that saccades produced before this were in response to the cue (pink) and saccades after were in response to the target (green). Therefore, we divide the saccades accordingly. Then, we aggregate the data per subject and cue. Last, we set all predictors to sum contrasts.

# preprocess face saccade counts  
df.cnt = df.sac %>%   
 group\_by(subID, diagnosis, dir\_face) %>%   
 summarise(  
 n.face = n()  
 )  
  
# preprocess cue saccade counts  
df.cnt.cue = df.sac %>%   
 filter(lat <= points$minima[idx]) %>%  
 group\_by(subID, diagnosis, dir\_face) %>%   
 summarise(  
 n.cue = n()  
 )  
  
# preprocess target saccade counts  
df.cnt.tar = df.sac %>%   
 filter(lat > points$minima[idx] & dir\_target) %>%  
 group\_by(subID, diagnosis, cue) %>%   
 summarise(  
 n.tar = n()  
 )  
  
# add a zero if no saccades were produced  
subID = rep(as.character(unique(df.sac$subID)),   
 each = length(unique(df.sac$dir\_face)))  
dir\_face = rep(as.character(unique(df.sac$dir\_face)),   
 times = length(unique(df.sac$subID)))  
df.cnt = merge(df.cnt, data.frame(subID, dir\_face), all = T) %>%  
 mutate(  
 n.face = if\_else(is.na(n.face), 0, n.face)  
 ) %>%  
 # merge with behavioural data  
 merge(., df.fab.agg) %>%  
 mutate\_if(is.character, as.factor)  
df.cnt.cue = merge(df.cnt.cue, data.frame(subID, dir\_face), all = T) %>%  
 mutate(  
 n.cue = if\_else(is.na(n.cue), 0, n.cue)  
 ) %>%  
 # merge with behavioural data  
 merge(., df.fab.agg) %>%  
 mutate\_if(is.character, as.factor)  
cue = rep(as.character(unique(df.sac$cue)), times = length(unique(df.sac$subID)))  
df.cnt.tar = merge(df.cnt.tar, data.frame(subID, cue), all = T) %>%  
 mutate(  
 n.tar = if\_else(is.na(n.tar), 0, n.tar)  
 ) %>%  
 # merge with behavioural data  
 merge(., df.fab.agg) %>%  
 mutate\_if(is.character, as.factor)  
  
# preprocess target latencies  
df.lat = df.sac %>%   
 # only keep latencies during cue  
 filter(lat > points$minima[idx]) %>%  
 # only keep the first latency of each trial  
 group\_by(subID, diagnosis, trl, cue) %>%  
 # only keep the first of the remaining saccades  
 filter(sac\_trl == min(sac\_trl)) %>%  
 merge(., df.fab) %>%  
 mutate\_if(is.character, as.factor)  
  
# add whether or not saccade to df.fab  
df.fab = df.fab %>%  
 merge(., df.lat %>% select(subID, trl, lat), all.x = T) %>%  
 mutate(  
 sac = if\_else(is.na(lat), F, T)  
 ) %>% select(-lat)  
  
# set and print the contrasts  
contrasts(df.lat$cue) = contr.sum(2)  
contrasts(df.lat$cue)

## [,1]  
## face 1  
## object -1

contrasts(df.lat$diagnosis) = contr.sum(4)  
contrasts(df.lat$diagnosis)

## [,1] [,2] [,3]  
## ADHD 1 0 0  
## ASD 0 1 0  
## BOTH 0 0 1  
## COMP -1 -1 -1

contrasts(df.cnt.tar$cue) = contr.sum(2)  
contrasts(df.cnt.tar$cue)

## [,1]  
## face 1  
## object -1

contrasts(df.cnt.tar$diagnosis) = contr.sum(4)  
contrasts(df.cnt.tar$diagnosis)

## [,1] [,2] [,3]  
## ADHD 1 0 0  
## ASD 0 1 0  
## BOTH 0 0 1  
## COMP -1 -1 -1

contrasts(df.cnt.cue$dir\_face) = contr.sum(2)  
contrasts(df.cnt.cue$dir\_face)

## [,1]  
## FALSE 1  
## TRUE -1

contrasts(df.cnt.cue$diagnosis) = contr.sum(4)  
contrasts(df.cnt.cue$diagnosis)

## [,1] [,2] [,3]  
## ADHD 1 0 0  
## ASD 0 1 0  
## BOTH 0 0 1  
## COMP -1 -1 -1

contrasts(df.cnt$dir\_face) = contr.sum(2)  
contrasts(df.cnt$dir\_face)

## [,1]  
## FALSE 1  
## TRUE -1

contrasts(df.cnt$diagnosis) = contr.sum(4)  
contrasts(df.cnt$diagnosis)

## [,1] [,2] [,3]  
## ADHD 1 0 0  
## ASD 0 1 0  
## BOTH 0 0 1  
## COMP -1 -1 -1

contrasts(df.fab$cue) = contr.sum(2)  
contrasts(df.fab$cue)

## [,1]  
## face 1  
## object -1

contrasts(df.fab$diagnosis) = contr.sum(4)  
contrasts(df.fab$diagnosis)

## [,1] [,2] [,3]  
## ADHD 1 0 0  
## ASD 0 1 0  
## BOTH 0 0 1  
## COMP -1 -1 -1

# S2.2 Number of saccades towards face during trial

First, we are going to assess the saccades that are produced in the direction of the face throughout the full trial: cue and target presentation. Based on Entzmann et al. (2021), we hypothesised that COMP participants produce more saccades towards the face than towards the object cues during the trials.

## Specify the model

Since we are counting the number of saccades, we use a poisson distribution for our model. We add an group-level intercept for each subject, and assess the influence of the predictors diagnostic status and whether the saccade was directed towards the side of the face or object as well as the interaction. We set our priors very wide because we do not have strong prior assumptions apart from people producing fewer saccades than there are trials.

code = "CNT"  
  
# set the formula  
f.cnt = brms::bf(n.face ~ diagnosis \* dir\_face + (1 | subID))  
  
# set priors based on study design  
priors = c(  
 prior(normal(3, 1.5), class = Intercept),   
 prior(normal(0, 1.0), class = sd),  
 prior(normal(0, 1.0), class = b)  
)  
  
# set number of iterations and warmup for models  
iter = 4500  
warm = 1500

## Simulation-based calibration

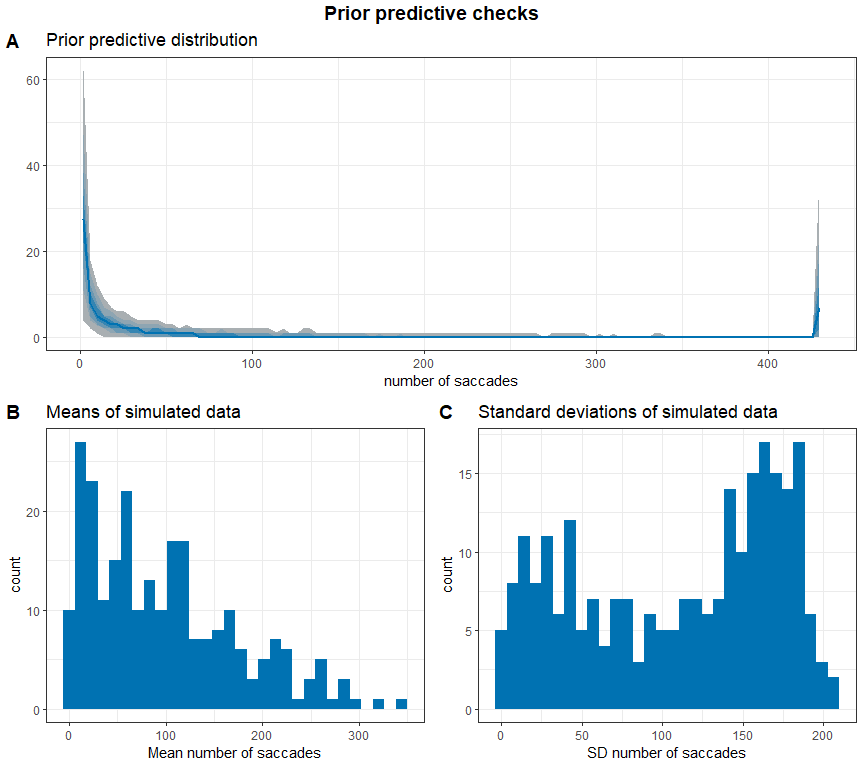
# check if the SBC already exists  
if (file.exists(file.path(cache\_dir, sprintf("df\_res\_%s.rds", code)))) {  
 # load in the results of the SBC  
 df.results = readRDS(file.path(cache\_dir, sprintf("df\_res\_%s.rds", code)))  
 df.backend = readRDS(file.path(cache\_dir, sprintf("df\_div\_%s.rds", code)))  
 dat = readRDS(file.path(cache\_dir, sprintf("dat\_%s.rds", code)))  
} else {  
 # perform the SBC  
 set.seed(2468)  
 gen = SBC\_generator\_brms(f.cnt, data = df.cnt, prior = priors,   
 thin = 50, warmup = 10000, refresh = 2000,  
 generate\_lp = TRUE, family = poisson(), init = 0.1)  
 bck = SBC\_backend\_brms\_from\_generator(gen, chains = 4, thin = 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")))  
}

We start by investigating the rhats and the number of divergent samples. This shows that 5 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and only 1 models had divergent samples. This suggests that this model performs well.

## Prior predictive checks

Next, we can plot the simulated values to perform prior predictive checks.

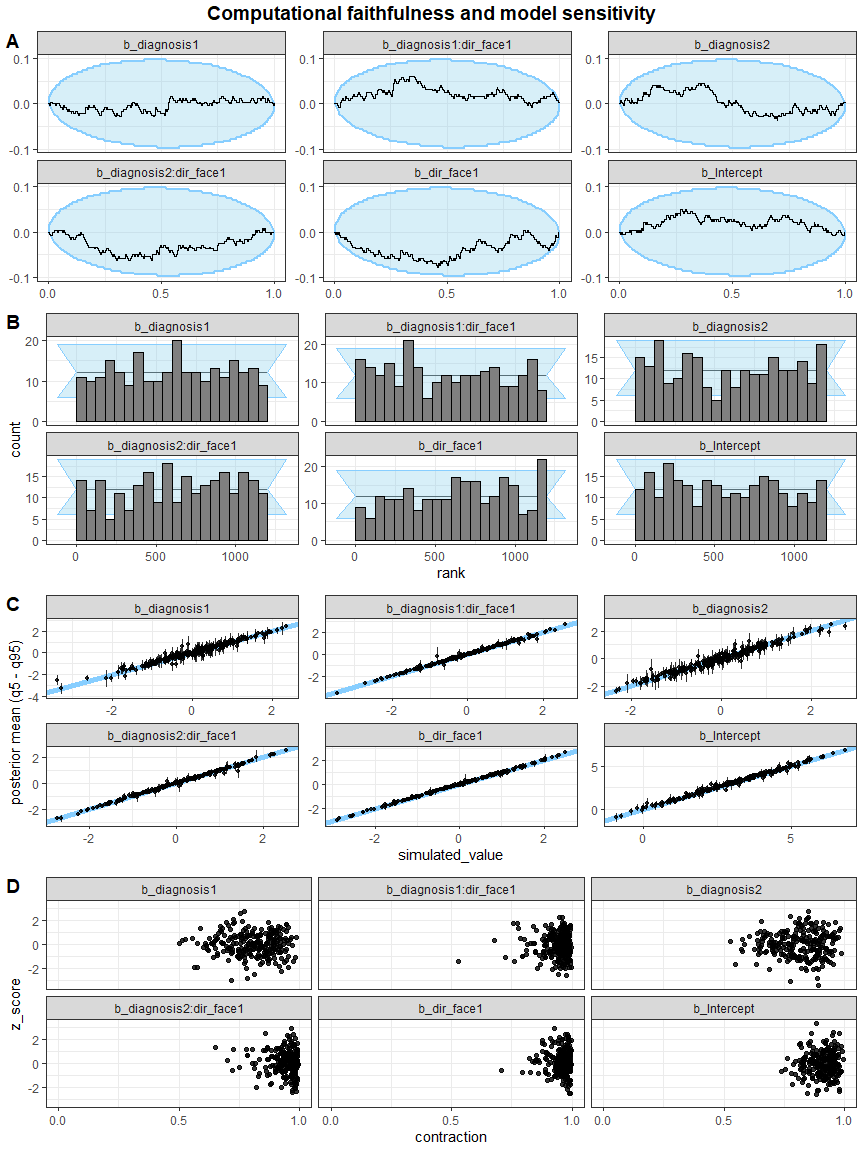
# get the true values  
truePars = dat[["variables"]]  
  
# create a matrix out of generated data  
dvname = gsub(" ", "", gsub("[\\|~].\*", "", f.cnt)[1])  
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))   
for (i in 1:length(dat[['generated']])) {  
 dvfakemat[,i] = dat[['generated']][[i]][[dvname]]  
}  
  
# set very large data points to a value of 432  
dvfakematH = dvfakemat;   
dvfakematH[dvfakematH > 432] = 432  
# compute one histogram per simulated data-set   
breaks = seq(0, max(dvfakematH, na.rm=T), length.out = 100)   
binwidth = round(breaks[2] - breaks[1])  
breaks = seq(0, max(dvfakematH, na.rm=T), binwidth)   
histmat = matrix(NA, ncol = nrow(truePars) + binwidth, nrow = length(breaks)-1)   
for (i in 1:nrow(truePars)) {  
 histmat[,i] = hist(dvfakematH[,i], breaks = breaks, plot = F)$counts   
}  
# for each bin, compute quantiles across histograms   
probs = seq(0.1, 0.9, 0.1)   
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))  
names(quantmat) = paste0("p", probs)  
for (i in 1:dim(histmat)[1]) {  
 quantmat[i,] = quantile(histmat[i,], p = probs, na.rm = T)  
}  
quantmat$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 = "Prior predictive distribution", y = "", x = "number of saccades") +  
 theme\_bw()  
  
tmpM = apply(dvfakematH, 2, mean) # mean   
tmpSD = apply(dvfakematH, 2, sd)   
p2 = ggplot() +   
 stat\_bin(aes(x = tmpM), fill = c\_dark) +   
 labs(x = "Mean number of saccades", title = "Means of simulated data") +  
 theme\_bw()  
p3 = ggplot() +   
 stat\_bin(aes(x = tmpSD), fill = c\_dark) +   
 labs(x = "SD number of saccades", title = "Standard deviations of simulated data") +  
 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", face = "bold", size = 14))



Since our priors were set very wide, we do get wide prior predictive distributions. We accept this and continue with our checks.

## Computational faithfulness and model sensitivity

# get simulation numbers with issues  
des\_rank = max(df.results$max\_rank)  
check = merge(df.results %>%   
 group\_by(sim\_id) %>%   
 summarise(  
 rhat = max(rhat, na.rm = T),   
 mean\_rank = max(max\_rank)  
 ) %>%   
 filter(rhat >= 1.05 | mean\_rank < des\_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$sim\_id)) %>%  
 ungroup() %>%  
 mutate(  
 max\_rank = max(rank)  
 )  
p1 = plot\_ecdf\_diff(df.results.b) + theme\_bw() + theme(legend.position = "none") +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p2 = plot\_rank\_hist(df.results.b, bins = 20) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p3 = plot\_sim\_estimated(df.results.b, alpha = .8) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p4 = plot\_contraction(df.results.b,   
 prior\_sd = setNames(  
 c(as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "Intercept",]$prior)),   
 rep(  
 as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "b",]$prior)),  
 length(unique(df.results.b$variable))-1)),   
 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))



All of this looks good. Despite our wide priors, the contraction shows a bit of a distribution which increases our trust that the wide priors are appropriate.

## Posterior predictive checks

As the next step, we fit the model and check whether the chains have converged, which they seem to have. We then perform posterior predictive checks on the model using the bayesplot package.

# fit the model  
set.seed(2468)  
m.cnt = brm(f.cnt,  
 df.cnt, prior = priors,  
 iter = iter, warmup = warm,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt-face",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
rstan::check\_hmc\_diagnostics(m.cnt$fit)

##   
## Divergences:

## 0 of 12000 iterations ended with a divergence.

##   
## Tree depth:

## 0 of 12000 iterations saturated the maximum tree depth of 10.

##   
## Energy:

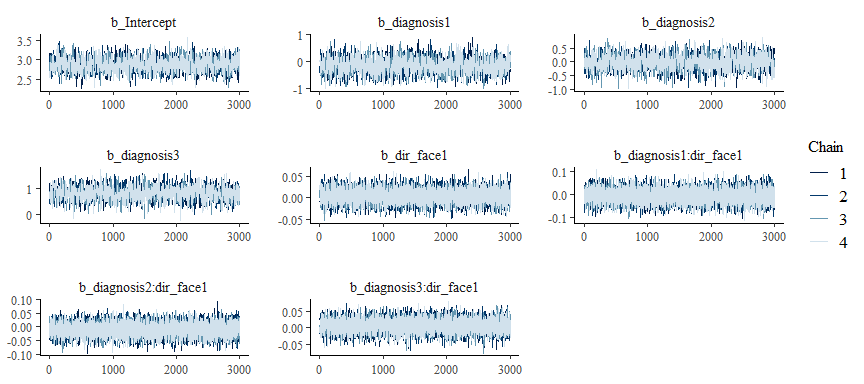
## E-BFMI indicated no pathological behavior.

# check that rhats are below 1.01  
sum(brms::rhat(m.cnt) >= 1.01, na.rm = T)

## [1] 0

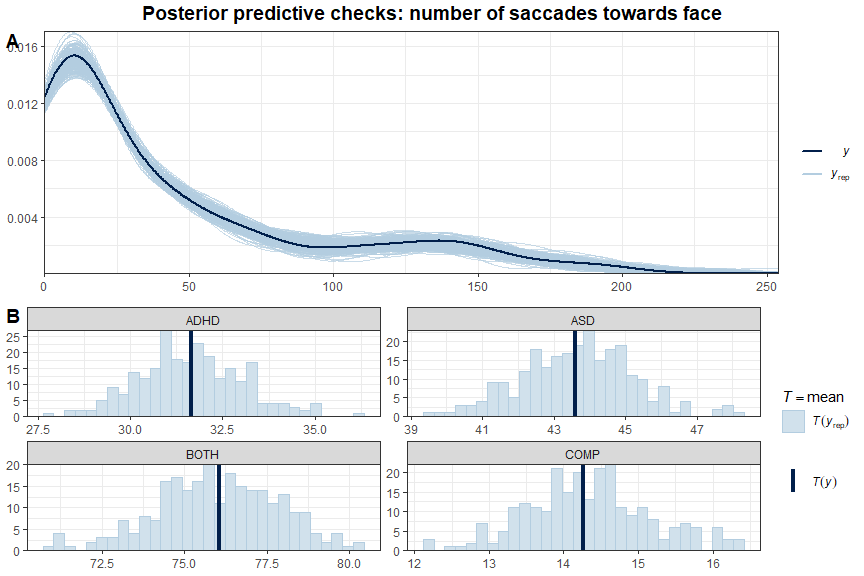
# check the trace plots  
post.draws = as\_draws\_df(m.cnt)  
mcmc\_trace(post.draws, regex\_pars = "^b\_",  
 facet\_args = list(ncol = 3)) +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))

## Scale for x is already present.  
## Adding another scale for x, which will replace the existing scale.



This model has no divergent samples and no rhats that are higher or equal to 1.01. Therefore, we go ahead and perform our posterior predictive checks.

# get the posterior predictions  
post.pred = posterior\_predict(m.cnt, ndraws = nsim)  
  
# check the fit of the predicted data compared to the real data  
p1 = pp\_check(m.cnt, ndraws = nsim) +   
 theme\_bw()  
  
# distributions of means and sds compared to the real values per group  
p2 = ppc\_stat\_grouped(df.cnt$n.face, post.pred, df.cnt$diagnosis) +   
 theme\_bw()  
  
p = ggarrange(p1, p2,  
 nrow = 2, ncol = 1, labels = "AUTO")  
annotate\_figure(p,   
 top = text\_grob("Posterior predictive checks: number of saccades towards face",   
 face = "bold", size = 14))



The predictions based on the model capture the data well. This further increases our trust in the model.

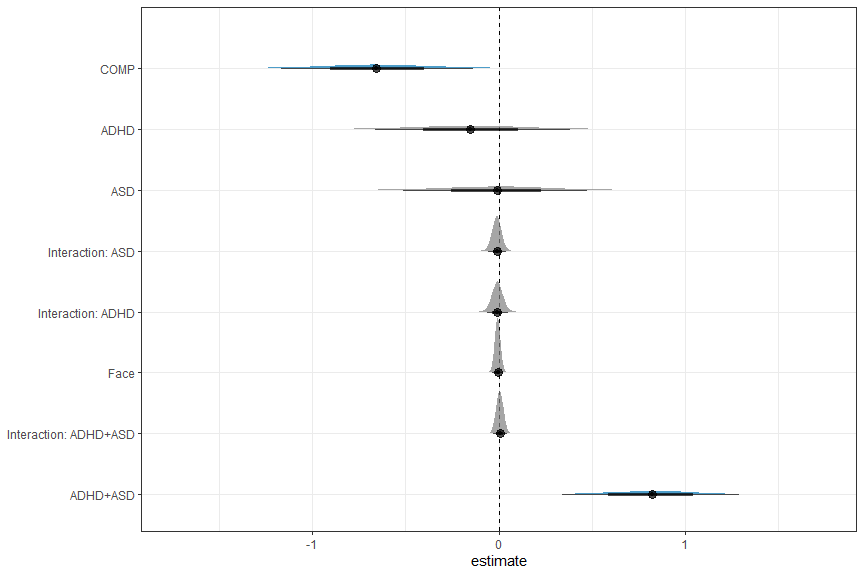
## Inferences

Now that we are convinced that we can trust our model, we have a look at the model and its estimates.

# print a summary  
summary(m.cnt)

## Family: poisson   
## Links: mu = log   
## Formula: n.face ~ diagnosis \* dir\_face + (1 | subID)   
## Data: df.cnt (Number of observations: 147)   
## Draws: 4 chains, each with iter = 4500; warmup = 1500; thin = 1;  
## total post-warmup draws = 12000  
##   
## Group-Level Effects:   
## ~subID (Number of levels: 76)   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## sd(Intercept) 1.32 0.12 1.11 1.58 1.00 1895 3363  
##   
## Population-Level Effects:   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS  
## Intercept 2.90 0.16 2.59 3.21 1.00 1372  
## diagnosis1 -0.15 0.27 -0.67 0.38 1.00 1528  
## diagnosis2 -0.01 0.25 -0.51 0.48 1.01 1312  
## diagnosis3 0.82 0.24 0.34 1.29 1.00 1458  
## dir\_face1 0.01 0.02 -0.03 0.04 1.00 8146  
## diagnosis1:dir\_face1 -0.01 0.03 -0.06 0.05 1.00 10740  
## diagnosis2:dir\_face1 -0.01 0.02 -0.06 0.04 1.00 9289  
## diagnosis3:dir\_face1 0.01 0.02 -0.03 0.04 1.00 8029  
## Tail\_ESS  
## Intercept 2549  
## diagnosis1 2591  
## diagnosis2 2437  
## diagnosis3 2562  
## dir\_face1 9054  
## diagnosis1:dir\_face1 9223  
## diagnosis2:dir\_face1 8669  
## diagnosis3:dir\_face1 8769  
##   
## 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.cnt = as\_draws\_df(m.cnt) %>%   
 select(starts\_with("b\_")) %>%  
 mutate(  
 b\_COMP = - b\_diagnosis1 - b\_diagnosis2 - b\_diagnosis3,  
 ASD = b\_Intercept + b\_diagnosis2,  
 ADHD = b\_Intercept + b\_diagnosis1,  
 BOTH = b\_Intercept + b\_diagnosis3,  
 b\_dir\_faceTRUE = - b\_dir\_face1,  
 COMP = b\_Intercept + b\_COMP  
 )  
  
# plot the posterior distributions  
df.m.cnt %>%   
 select(starts\_with("b\_")) %>%  
 pivot\_longer(cols = starts\_with("b\_"), names\_to = "coef", values\_to = "estimate") %>%  
 filter(coef != "b\_Intercept" & coef != "b\_dir\_face1") %>%  
 mutate(  
 coef = case\_match(coef,  
 "b\_diagnosis1" ~ "ADHD",  
 "b\_diagnosis2" ~ "ASD",  
 "b\_diagnosis3" ~ "ADHD+ASD",  
 "b\_COMP" ~ "COMP",  
 "b\_dir\_faceTRUE" ~ "Face",  
 "b\_diagnosis1:dir\_face1" ~ "Interaction: ADHD",  
 "b\_diagnosis2:dir\_face1" ~ "Interaction: ASD",  
 "b\_diagnosis3:dir\_face1" ~ "Interaction: ADHD+ASD"  
 ),  
 coef = fct\_reorder(coef, desc(estimate))  
 ) %>%   
 group\_by(coef) %>%  
 mutate(  
 cred = case\_when(  
 (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |  
 (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")



# H2a: COMP: face > object  
h2a = hypothesis(m.cnt,   
 "0 < 2\*(dir\_face1 - diagnosis1:dir\_face1 -   
 diagnosis2:dir\_face1 - diagnosis3:dir\_face1)")  
h2a

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*(dir\_face1... < 0 -0.03 0.09 -0.18 0.11 1.77  
## Post.Prob Star  
## 1 0.64   
## ---  
## 'CI': 90%-CI for one-sided and 95%-CI for two-sided hypotheses.  
## '\*': For one-sided hypotheses, the posterior probability exceeds 95%;  
## for two-sided hypotheses, the value tested against lies outside the 95%-CI.  
## Posterior probabilities of point hypotheses assume equal prior probabilities.

# explore difference in saccades between COMP and BOTH  
e1 = hypothesis(m.cnt, "0 < 2\*(diagnosis1 + diagnosis2 + 2\*diagnosis3)",   
 alpha = 0.025)  
e1

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*(diagnosis... < 0 -2.94 0.82 -4.55 -1.28 2999  
## Post.Prob Star  
## 1 1 \*  
## ---  
## '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.

# explore difference in saccades between ADHD and BOTH  
e2 = hypothesis(m.cnt, "0 > 2\*diagnosis1 - 2\*diagnosis3",   
 alpha = 0.025)  
e2

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*diagnosis1... > 0 1.93 0.83 0.3 3.56 91.31  
## Post.Prob Star  
## 1 0.99 \*  
## ---  
## '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.

# explore difference in saccades between ASD and BOTH  
e3 = hypothesis(m.cnt, "0 < -2\*diagnosis2 + 2\*diagnosis3",   
 alpha = 0.025)  
e3

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(-2\*diagnosis... < 0 -1.66 0.79 -3.22 -0.13 59.91  
## Post.Prob Star  
## 1 0.98 \*  
## ---  
## '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.

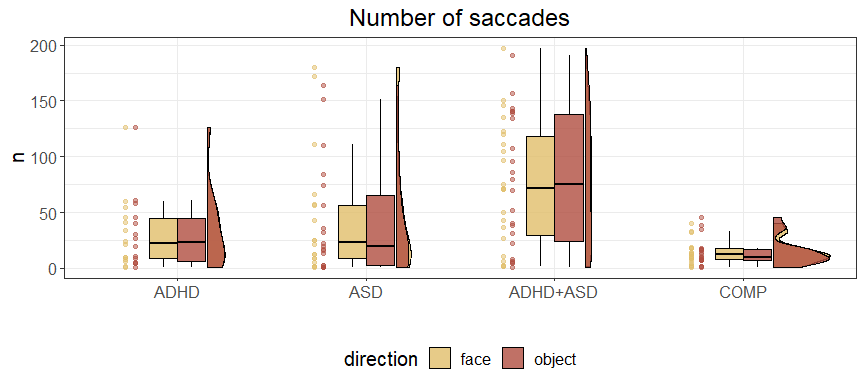
# extract predicted differences   
df.new = df.cnt %>%   
 select(diagnosis, dir\_face) %>%   
 distinct() %>%  
 mutate(  
 condition = paste(diagnosis, dir\_face, sep = "\_")  
 )  
df.est = as.data.frame(  
 fitted(m.cnt, summary = F,   
 newdata = df.new %>% select(diagnosis, dir\_face),   
 re\_formula = NA))  
colnames(df.est) = df.new$condition

Our hypothesis was not confirmed: there was no difference between the number of saccades in the direction of the face compared to the direction of the object in our COMP group (*estimate* = -0.03 [-0.18, 0.11], *posterior probability* = 63.84%). Exploration of other group differences indicates a larger number of saccades produced by the ADHD+ASD group than in any of the other groups (COMP: *estimate* = -2.94 [-4.55, -1.28], *posterior probability* = 99.97%; ADHD: *estimate* = 1.93 [0.3, 3.56], *posterior probability* = 98.92%; ASD: *estimate* = -1.66 [-3.22, -0.13], *posterior probability* = 98.36%).

## Plots

As a next step, we can now finally plot our data.

# rain cloud plot  
df.cnt %>%  
 mutate(  
 diagnosis = recode(diagnosis, "BOTH" = "ADHD+ASD")  
 ) %>%   
 mutate(direction = if\_else(dir\_face == T, "face", "object")) %>%  
 ggplot(aes(diagnosis, n.face, fill = direction, colour = direction)) + #  
 geom\_rain(rain.side = 'r',  
boxplot.args = list(color = "black", outlier.shape = NA, show\_guide = FALSE, alpha = .8),  
violin.args = list(color = "black", outlier.shape = NA, alpha = .8),  
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.col2) +  
 scale\_color\_manual(values = custom.col2) +  
 labs(title = "Number of saccades", x = "", y = "n") +  
 theme\_bw() +   
 theme(legend.position = "bottom",   
 plot.title = element\_text(hjust = 0.5),   
 legend.direction = "horizontal",   
 text = element\_text(size = 15))

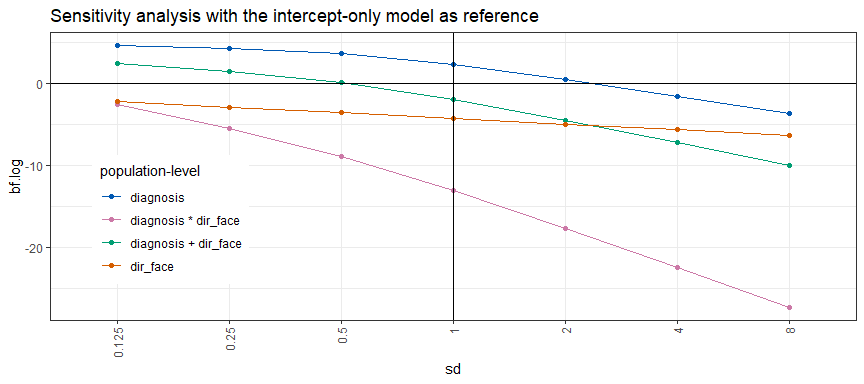


ggsave("Fig5\_nrSac.pdf",   
 units = "mm",   
 width = 170,  
 height = 100,  
 dpi = 300)

## Bayes factor analysis

To complement our hypothesis testing using brms::hypothesis(), we perform a Bayes Factor analysis with models excluding some of our population-level predictors.

# set the directory in which to save results  
sense\_dir = file.path(getwd(), "\_brms\_sens\_cache")  
log\_dir = sense\_dir  
main.code = "cnt"  
  
# describe priors  
pr.descriptions = c("chosen",  
 "sdx2", "sdx4", "sdx8",   
 "sdx0.5", "sdx0.25", "sdx0.125"  
 )  
  
# check which have been run already  
if (file.exists(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)))) {  
 pr.done = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F) %>%  
 select(priors) %>% distinct()  
 pr.descriptions = pr.descriptions[!(pr.descriptions %in% pr.done$priors)]  
}  
  
if (length(pr.descriptions) > 0) {  
 # rerun the model with more iterations for bridgesampling  
 set.seed(6633)  
 m.cnt.bf = brm(f.cnt,  
 df.cnt, prior = priors,  
 iter = 40000, warmup = 10000,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt\_bf",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
}  
  
# loop through them  
for (pr.desc in pr.descriptions) {  
 tryCatch({  
 # use the function  
 bf\_sens\_2int(m.cnt.bf, "diagnosis", "dir\_face", pr.desc,   
 main.code, # prefix for all models and MLL  
 file.path(log\_dir, "log\_FAB\_bf.txt"), # log file  
 sense\_dir, # where to save the models and MLL  
 reps = 5  
 )  
 },  
 error = function(err) {  
 message(sprintf("Error for %s: %s", pr.desc, err))  
 }  
 )  
}  
  
# read in the results  
df.cnt.bf = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F)  
  
# check the sensitivity analysis result per model  
df.cnt.bf %>%  
 filter(`population-level` != "1") %>%  
 mutate(  
 sd = as.factor(case\_when(  
 priors == "chosen" ~ "1",   
 substr(priors, 1, 3) == "sdx" ~ gsub("sdx", "", priors),  
 T ~ priors)  
 ),  
 order = case\_when(  
 priors == "chosen" ~ 1,   
 substr(priors, 1, 3) == "sdx" ~ as.numeric(gsub("sdx", "", priors)),  
 T ~ 999),  
 sd = fct\_reorder(sd, order)  
 ) %>%  
 ggplot(aes(y = bf.log,   
 x = sd,   
 group = `population-level`,   
 colour = `population-level`)) +   
 geom\_point() +  
 geom\_line() +   
 geom\_vline(xintercept = "1") +  
 geom\_hline(yintercept = 0) +  
 ggtitle("Sensitivity analysis with the intercept-only model as reference") +  
 #facet\_wrap(. ~ `population-level`, scales = "free\_y") +  
 scale\_colour\_manual(values = custom.col) +  
 theme\_bw() +  
 theme(legend.position = c(0.15,0.35),   
 axis.text.x = element\_text(angle = 90, vjust = 0.5, hjust = 1))



# print the BFs based on chosen priors  
kable(df.cnt.bf %>% filter(priors == "chosen") %>% select(-priors) %>%  
 filter(`population-level` != "1") %>% arrange(desc(bf.log)), digits = 3)

| population-level | bf.log |
| --- | --- |
| diagnosis | 2.259 |
| diagnosis + dir\_face | -2.002 |
| dir\_face | -4.266 |
| diagnosis \* dir\_face | -13.125 |

The Bayes Factor analysis revealed the model only including the main effect diagnostic status outperforming the intercept-only model. With wider priors, the Bayes Factor of this model also indicates worse fit for our data than the intercept-only model. For our chosen priors though, there is moderate evidence in favour of this model compared to the intercept-only model (diagnostic groups: log(*BF*) = 2.259). The intercept-only model consistently outperforms the other models (saccade direction: log(*BF*) = -4.266; diagnostic group and saccade direction: log(*BF*) = -2.002; full model: log(*BF*) = -13.125). These results suggest that diagnostic status influences the amount of saccades made, however, the same amount of saccades are produced towards the face and towards the object.

# S2.3 Saccade latencies

Next, we focus on the latencies of the saccades that are produced during the presentation of the targets to assess whether cue type, diagnostic group or their interaction influence latencies. We hypothesised that ASD participants show an increased latency compared to COMP participants when producing saccades towards targets appearing at the location of a face.

## Full model

We start with the model containing all latencies of saccades produced during the target presentation. We choose a shifted lognormal distribution because saccade latencies below 100ms are very unlikely. Additionally, latencies are determined based on the onset of the cue presentation (200ms).

### Setting up and assessing the model

code = "LAT"  
  
# set the formula  
f.lat = brms::bf(lat ~ diagnosis \* cue + (cue | subID) + (diagnosis \* cue | stm))  
  
# set weakly informative priors  
priors = c(  
 prior(normal(5, 0.75), class = Intercept),  
 prior(normal(0, 0.25), class = sd),  
 prior(normal(0, 0.25), class = b),  
 prior(normal(0.5, 0.50), class = sigma),  
 prior(normal(350, 50.00), class = ndt), # threshold between target and cue saccades  
 prior(lkj(2), class = cor)  
)  
  
# set number of iterations and warmup for models  
iter = 3000  
warm = 1000

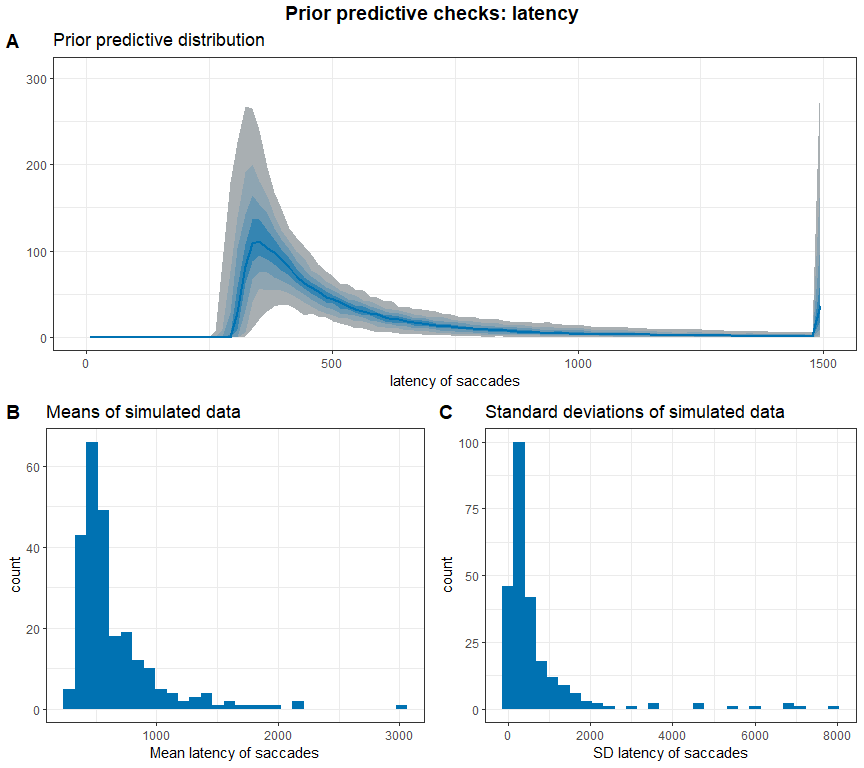
if (file.exists(file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))) {  
 # load in the results of the SBC  
 df.results = readRDS(file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 df.backend = readRDS(file = file.path(cache\_dir, paste0("df\_div\_", code, ".rds")))  
 dat = readRDS(file = file.path(cache\_dir, paste0("dat\_", code, ".rds")))  
} else {  
 # create the data and the results  
 set.seed(2468)  
 gen = SBC\_generator\_brms(f.lat, data = df.lat, prior = priors,   
 family = "shifted\_lognormal",  
 thin = 50, warmup = 10000, refresh = 2000,  
 generate\_lp = TRUE)  
 dat = generate\_datasets(gen, nsim)   
 saveRDS(dat, file = file.path(cache\_dir, paste0("dat\_", code, ".rds")))  
 backend = SBC\_backend\_brms\_from\_generator(gen, chains = 4, thin = 1,  
 warmup = 1000, iter = 3000)  
 results = compute\_SBC(dat, backend,  
 cache\_mode = "results",   
 cache\_location = file.path(cache\_dir, paste0("res\_", code)))  
 saveRDS(results$stats,   
 file = file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 saveRDS(results$backend\_diagnostics,   
 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 0 of 250 simulations had at least one parameter that had an rhat of at least 1.05 and only 1 model had divergent samples (mean number of samples of the simulations with divergent samples: 1). This suggests that this model performs well.

### Prior predictive checks

Next, we can plot the simulated values to perform prior predictive checks.

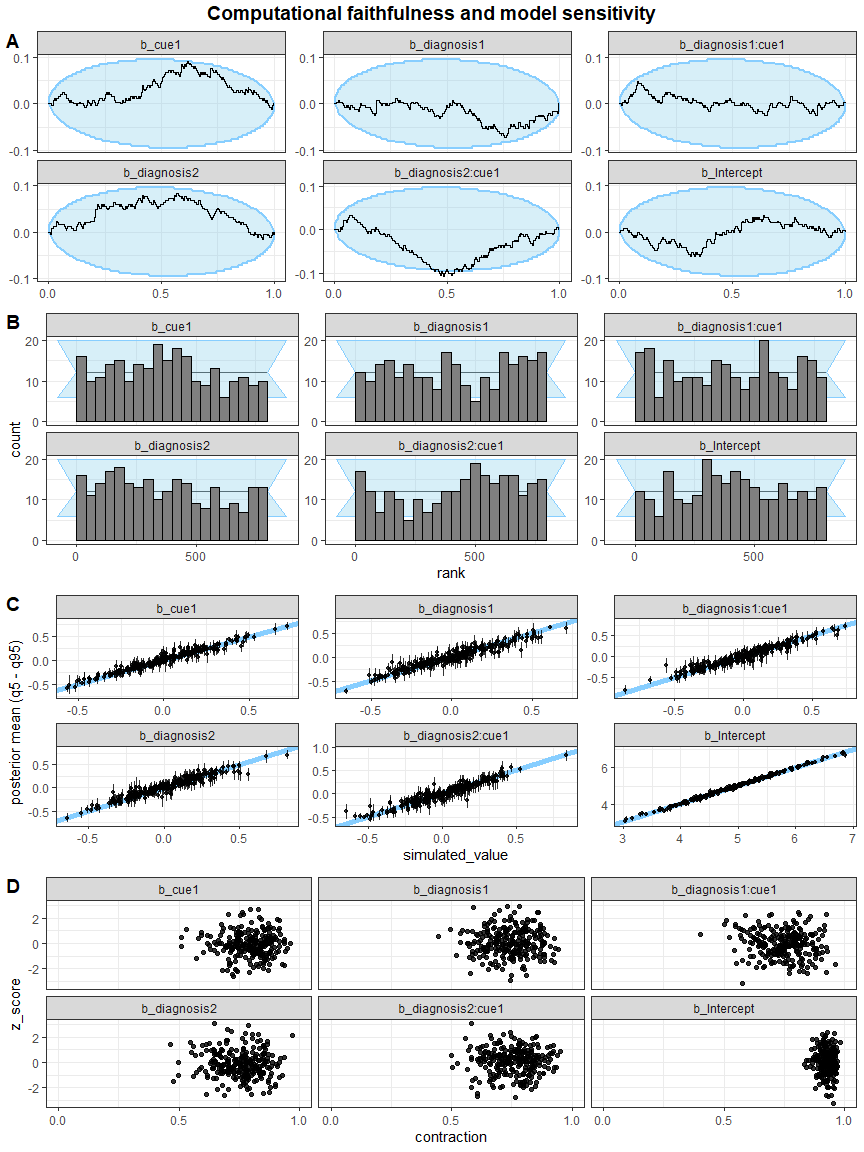
# get the true values  
truePars = dat[["variables"]]  
  
# create a matrix out of generated data  
dvname = gsub(" ", "", gsub("[\\|~].\*", "", f.lat)[1])  
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))   
for (i in 1:length(dat[['generated']])) {  
 dvfakemat[,i] = dat[['generated']][[i]][[dvname]]  
}  
  
# set very large data points to a value of 1500  
dvfakematH = dvfakemat;   
dvfakematH[dvfakematH < 0] = 0  
dvfakematH[dvfakematH > 1500] = 1500  
# compute one histogram per simulated data-set   
breaks = seq(0, 1500, length.out = 101)   
binwidth = breaks[2] - breaks[1]  
histmat = matrix(NA, ncol = nrow(truePars) + binwidth, nrow = length(breaks)-1)   
for (i in 1:nrow(truePars)) {  
 histmat[,i] = hist(dvfakematH[,i], breaks = breaks, plot = F)$counts   
}  
# for each bin, compute quantiles across histograms   
probs = seq(0.1, 0.9, 0.1)   
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))  
names(quantmat) = paste0("p", probs)  
for (i in 1:dim(histmat)[1]) {  
 quantmat[i,] = quantile(histmat[i,], p = probs, na.rm = T)  
}  
quantmat$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 = "Prior predictive distribution", y = "", x = "latency of saccades") +  
 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 latency of saccades", title = "Means of simulated data") +  
 theme\_bw()  
p3 = ggplot() +   
 stat\_bin(aes(x = tmpSD), fill = c\_dark) +   
 labs(x = "SD latency of saccades", title = "Standard deviations of simulated data") +  
 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: latency",   
 face = "bold", size = 14))



First, we assess whether the simulated values fit our expectations of the distribution of the data. Previous literature has found that saccade latencies are around 200ms with few saccades being produced faster than 100ms. If we add 200ms from the cue presentation, this means we expect most latencies to be above 300ms and centered around 400ms. Our simulated datasets seem to capture this well.

### Computational faithfulness and model sensitivity

# get simulation numbers with issues  
des\_rank = max(df.results$max\_rank)  
check = merge(df.results %>%   
 group\_by(sim\_id) %>%   
 summarise(  
 rhat = max(rhat, na.rm = T),   
 mean\_rank = max(max\_rank)  
 ) %>%   
 filter(rhat >= 1.05 | mean\_rank < des\_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$sim\_id)) %>%  
 ungroup() %>%  
 mutate(  
 max\_rank = max(rank)  
 )  
p1 = plot\_ecdf\_diff(df.results.b) + theme\_bw() + theme(legend.position = "none") +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p2 = plot\_rank\_hist(df.results.b, bins = 20) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p3 = plot\_sim\_estimated(df.results.b, alpha = .8) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p4 = plot\_contraction(df.results.b,   
 prior\_sd = setNames(  
 c(as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "Intercept",]$prior)),   
 rep(  
 as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "b",]$prior)),  
 length(unique(df.results.b$variable))-1)),   
 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))



All looks acceptable here.

### Posterior predictive checks

As the next step, we fit the model and check whether the chains have converged, which they seem to have. We then perform posterior predictive checks on the model using the bayesplot package.

# fit the maximal model  
set.seed(2587)  
m.lat = brm(f.lat,  
 df.lat, prior = priors,  
 iter = iter, warmup = warm,  
 backend = "cmdstanr", threads = threading(8),  
 family = "shifted\_lognormal",  
 file = "m\_lat",  
 save\_pars = save\_pars(all = TRUE)  
 )  
rstan::check\_hmc\_diagnostics(m.lat$fit)

##   
## Divergences:

## 0 of 8000 iterations ended with a divergence.

##   
## Tree depth:

## 0 of 8000 iterations saturated the maximum tree depth of 10.

##   
## Energy:

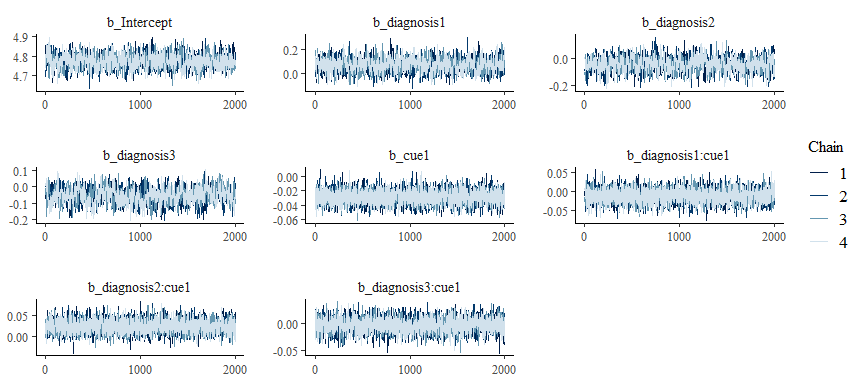
## E-BFMI indicated no pathological behavior.

# check that rhats are below 1.01  
sum(brms::rhat(m.lat) >= 1.01, na.rm = T)

## [1] 0

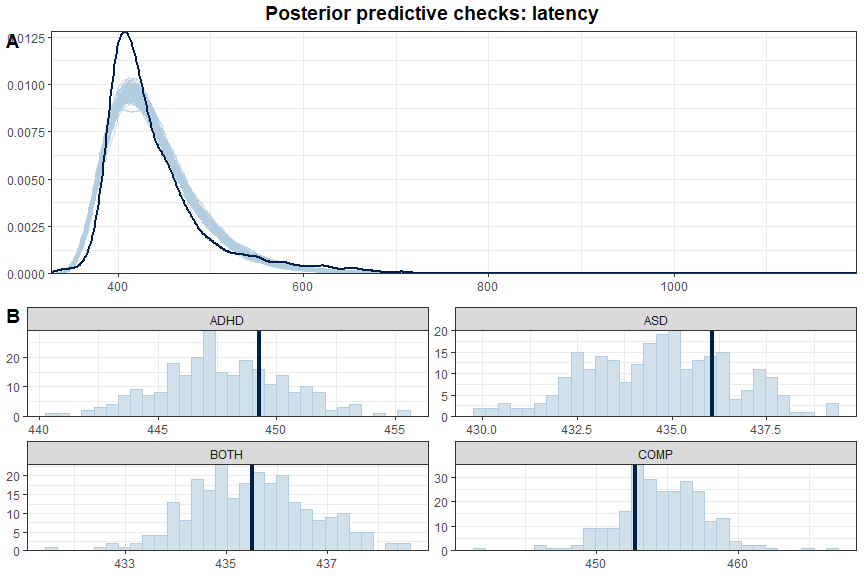
# check the trace plots  
post.draws = as\_draws\_df(m.lat)  
mcmc\_trace(post.draws, regex\_pars = "^b\_",   
 facet\_args = list(ncol = 3)) +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))

## Scale for x is already present.  
## Adding another scale for x, which will replace the existing scale.



The model does not have any divergent transitions nor high rhats. The trace plots also look good, therefore, we move on to the posterior predictive checks.

# get the posterior predictions  
post.pred = posterior\_predict(m.lat, ndraws = nsim)  
  
# check the fit of the predicted data compared to the real data  
p1 = pp\_check(m.lat, ndraws = nsim) +   
 theme\_bw() + theme(legend.position = "none")  
  
# distributions of means and sds compared to the real values per group  
p2 = ppc\_stat\_grouped(df.lat$lat, post.pred, df.lat$diagnosis) +   
 theme\_bw() + theme(legend.position = "none")  
  
p = ggarrange(p1, p2,   
 nrow = 2, ncol = 1, labels = "AUTO")  
annotate\_figure(p,   
 top = text\_grob("Posterior predictive checks: latency",   
 face = "bold", size = 14))



The simulated data based on the model does not fit our data very well: it is wider and seems to underestimate latencies for COMP while overestimating for ADHD and ASD with the dark blue line showing the mean of the actual dataset and the light blue bars showing the distribution of the predicted data.

## Aggregated model

Since we want to base our inferences on the estimates, we go back to the drawing board and aggregate our data to see whether this resolves these issues.

### Setting up and assessing the model

code = "LAT\_agg"  
  
# aggregate the data  
df.lat.agg = df.lat %>%   
 group\_by(subID, cue, diagnosis) %>%   
 summarise(lat = median(lat, na.rm = T))  
  
# set the formula  
f.lat = brms::bf(lat ~ diagnosis \* cue + (1 | subID) )  
  
# set weakly informative priors  
priors = priors %>% filter(class != "cor")  
  
# set number of iterations and warmup for models  
iter = 3000  
warm = 1000

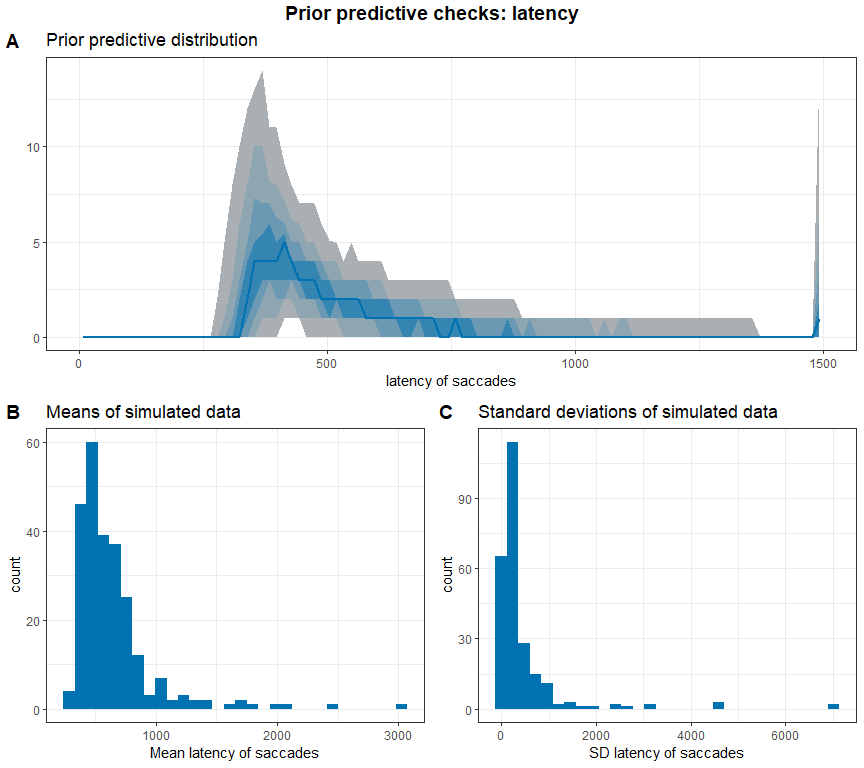
if (file.exists(file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))) {  
 # load in the results of the SBC  
 df.results = readRDS(file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 df.backend = readRDS(file = file.path(cache\_dir, paste0("df\_div\_", code, ".rds")))  
 dat = readRDS(file = file.path(cache\_dir, paste0("dat\_", code, ".rds")))  
} else {  
 # create the data and the results  
 set.seed(2468)  
 gen = SBC\_generator\_brms(f.lat, data = df.lat.agg, prior = priors,   
 family = "shifted\_lognormal",  
 thin = 50, warmup = 10000, refresh = 2000,  
 generate\_lp = TRUE)  
 dat = generate\_datasets(gen, nsim)   
 saveRDS(dat, file = file.path(cache\_dir, paste0("dat\_", code, ".rds")))  
 backend = SBC\_backend\_brms\_from\_generator(gen, chains = 4, thin = 1,  
 warmup = 1000, iter = 3000)  
 results = compute\_SBC(dat, backend,  
 cache\_mode = "results",   
 cache\_location = file.path(cache\_dir, paste0("res\_", code)))  
 saveRDS(results$stats,   
 file = file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 saveRDS(results$backend\_diagnostics,   
 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 1 of 250 simulations had at least one parameter that had an rhat of at least 1.05, but 105 models had divergent samples (mean number of samples of the simulations with divergent samples: 7.21). This is something to look out for in the final model.

### Prior predictive checks

Next, we can plot the simulated values to perform prior predictive checks.

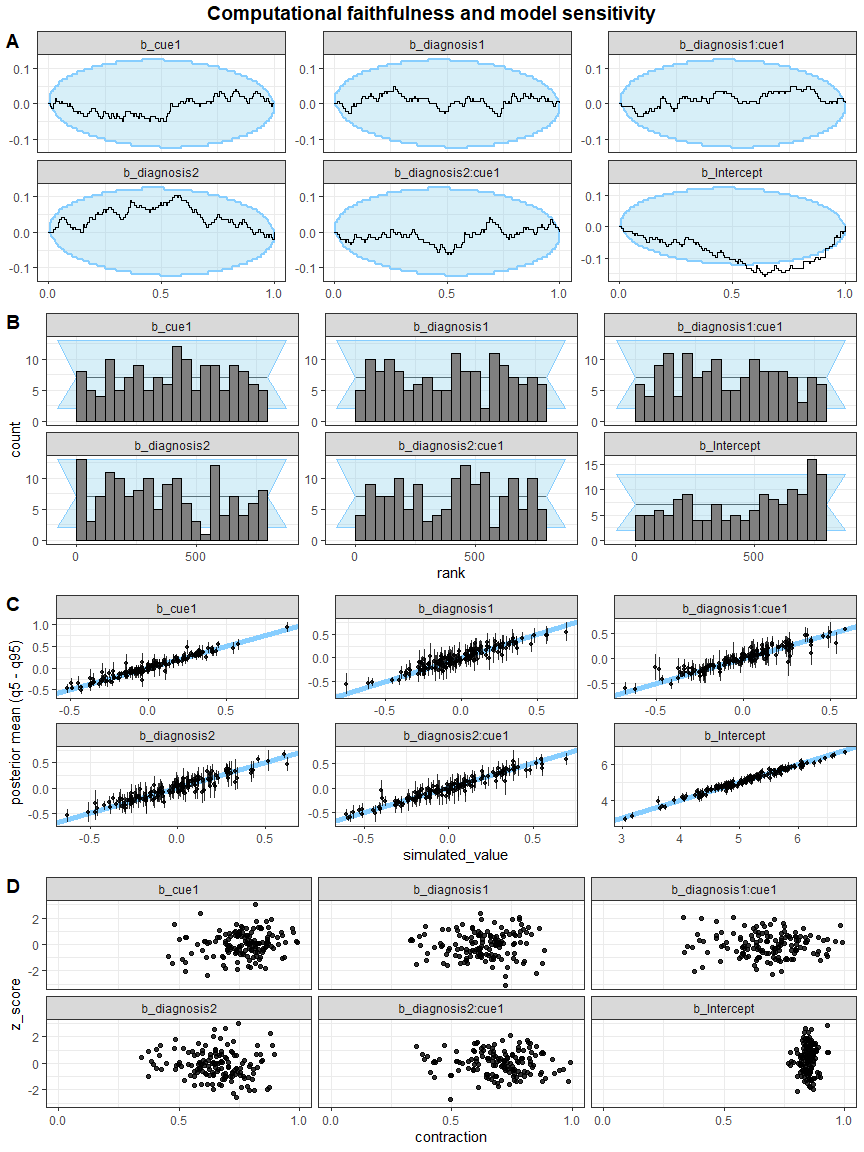
# get the true values  
truePars = dat[["variables"]]  
  
# create a matrix out of generated data  
dvname = gsub(" ", "", gsub("[\\|~].\*", "", f.lat)[1])  
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))   
for (i in 1:length(dat[['generated']])) {  
 dvfakemat[,i] = dat[['generated']][[i]][[dvname]]  
}  
  
# set very large data points to a value of 1500  
dvfakematH = dvfakemat;   
dvfakematH[dvfakematH < 0] = 0  
dvfakematH[dvfakematH > 1500] = 1500  
# compute one histogram per simulated data-set   
breaks = seq(0, 1500, length.out = 101)   
binwidth = breaks[2] - breaks[1]  
histmat = matrix(NA, ncol = nrow(truePars) + binwidth, nrow = length(breaks)-1)   
for (i in 1:nrow(truePars)) {  
 histmat[,i] = hist(dvfakematH[,i], breaks = breaks, plot = F)$counts   
}  
# for each bin, compute quantiles across histograms   
probs = seq(0.1, 0.9, 0.1)   
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))  
names(quantmat) = paste0("p", probs)  
for (i in 1:dim(histmat)[1]) {  
 quantmat[i,] = quantile(histmat[i,], p = probs, na.rm = T)  
}  
quantmat$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 = "Prior predictive distribution", y = "", x = "latency of saccades") +  
 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 latency of saccades", title = "Means of simulated data") +  
 theme\_bw()  
p3 = ggplot() +   
 stat\_bin(aes(x = tmpSD), fill = c\_dark) +   
 labs(x = "SD latency of saccades", title = "Standard deviations of simulated data") +  
 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: latency",   
 face = "bold", size = 14))



Again, our simulated datasets seem to capture well what we know about saccade latencies.

### Computational faithfulness and model sensitivity

# get simulation numbers with issues  
des\_rank = max(df.results$max\_rank)  
check = merge(df.results %>%   
 group\_by(sim\_id) %>%   
 summarise(  
 rhat = max(rhat, na.rm = T),   
 mean\_rank = max(max\_rank)  
 ) %>%   
 filter(rhat >= 1.05 | mean\_rank < des\_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$sim\_id)) %>%  
 ungroup() %>%  
 mutate(  
 max\_rank = max(rank)  
 )  
p1 = plot\_ecdf\_diff(df.results.b) + theme\_bw() + theme(legend.position = "none") +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p2 = plot\_rank\_hist(df.results.b, bins = 20) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p3 = plot\_sim\_estimated(df.results.b, alpha = .8) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p4 = plot\_contraction(df.results.b,   
 prior\_sd = setNames(  
 c(as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "Intercept",]$prior)),   
 rep(  
 as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "b",]$prior)),  
 length(unique(df.results.b$variable))-1)),   
 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))



The intercept looks slightly off here, the model could have a slight tendency to underestimate it.

### Posterior predictive checks

As the next step, we fit the model and check whether the chains have converged, which they seem to have. We then perform posterior predictive checks on the model using the bayesplot package.

# fit the maximal model  
set.seed(7799)  
m.lat = brm(f.lat,  
 df.lat.agg, prior = priors,  
 iter = iter, warmup = warm,  
 backend = "cmdstanr", threads = threading(8),  
 family = "shifted\_lognormal",  
 file = "m\_lat\_agg",  
 save\_pars = save\_pars(all = TRUE)  
 )  
rstan::check\_hmc\_diagnostics(m.lat$fit)

##   
## Divergences:

## 0 of 8000 iterations ended with a divergence.

##   
## Tree depth:

## 0 of 8000 iterations saturated the maximum tree depth of 10.

##   
## Energy:

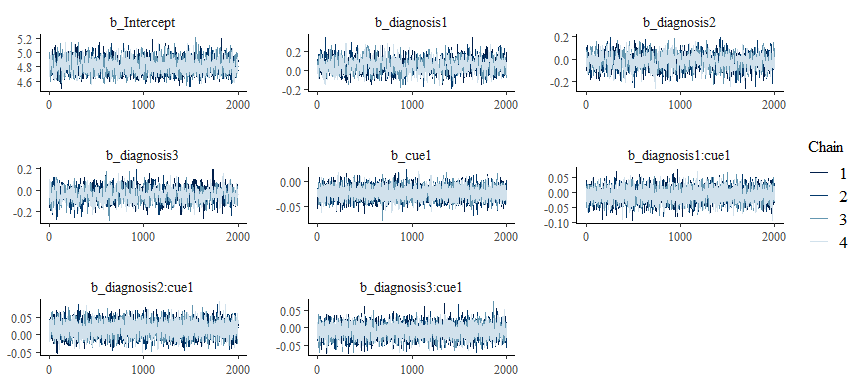
## E-BFMI indicated no pathological behavior.

# check that rhats are below 1.01  
sum(brms::rhat(m.lat) >= 1.01, na.rm = T)

## [1] 0

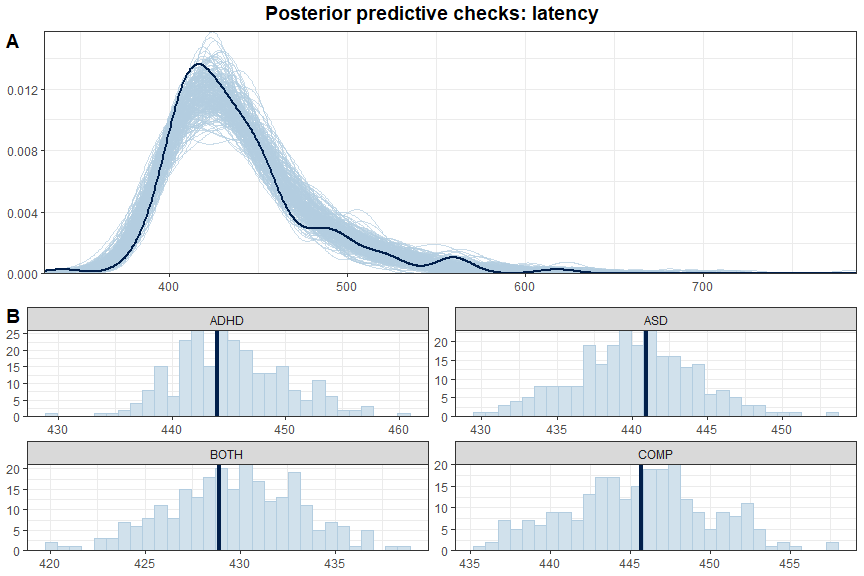
# check the trace plots  
post.draws = as\_draws\_df(m.lat)  
mcmc\_trace(post.draws, regex\_pars = "^b\_",   
 facet\_args = list(ncol = 3)) +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))

## Scale for x is already present.  
## Adding another scale for x, which will replace the existing scale.



The final model does not exhibit any divergence issues or suboptimal rhats.

# get the posterior predictions  
post.pred = posterior\_predict(m.lat, ndraws = nsim)  
  
# check the fit of the predicted data compared to the real data  
p1 = pp\_check(m.lat, ndraws = nsim) +   
 theme\_bw() + theme(legend.position = "none")  
  
# distributions of means and sds compared to the real values per group  
p2 = ppc\_stat\_grouped(df.lat.agg$lat, post.pred, df.lat.agg$diagnosis) +   
 theme\_bw() + theme(legend.position = "none")  
  
p = ggarrange(p1, p2,   
 nrow = 2, ncol = 1, labels = "AUTO")  
annotate\_figure(p,   
 top = text\_grob("Posterior predictive checks: latency",   
 face = "bold", size = 14))



This looks much better with the simulated data based on the model capturing our actual data well.

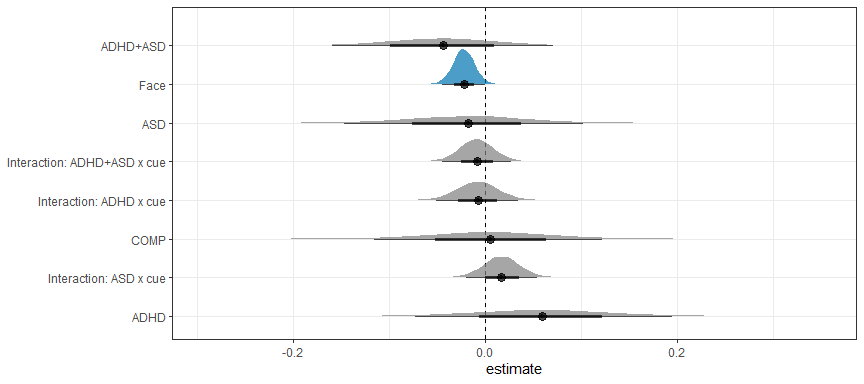
### Inferences

Now that we are convinced that we can trust our model, we have a look at the model and its estimates.

# print a summary  
summary(m.lat)

## Family: shifted\_lognormal   
## Links: mu = identity; sigma = identity; ndt = identity   
## Formula: lat ~ diagnosis \* cue + (1 | subID)   
## Data: df.lat.agg (Number of observations: 144)   
## Draws: 4 chains, each with iter = 3000; warmup = 1000; thin = 1;  
## total post-warmup draws = 8000  
##   
## Group-Level Effects:   
## ~subID (Number of levels: 76)   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## sd(Intercept) 0.30 0.04 0.23 0.39 1.00 1606 2660  
##   
## Population-Level Effects:   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## Intercept 4.82 0.09 4.65 5.01 1.00 1883 2993  
## diagnosis1 0.06 0.07 -0.07 0.19 1.00 1046 1940  
## diagnosis2 -0.02 0.06 -0.15 0.10 1.00 1153 1948  
## diagnosis3 -0.04 0.06 -0.16 0.07 1.00 1042 2027  
## cue1 -0.02 0.01 -0.04 -0.00 1.00 7366 5787  
## diagnosis1:cue1 -0.01 0.02 -0.05 0.03 1.00 5985 5124  
## diagnosis2:cue1 0.02 0.02 -0.02 0.05 1.00 7074 5703  
## diagnosis3:cue1 -0.01 0.02 -0.04 0.03 1.00 6622 5519  
##   
## Family Specific Parameters:   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## sigma 0.13 0.02 0.10 0.16 1.00 2514 3608  
## ndt 309.30 10.65 284.58 325.59 1.00 2173 2914  
##   
## 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).

# plot the posterior distributions  
as\_draws\_df(m.lat) %>%   
 select(starts\_with("b\_")) %>%  
 mutate(  
 b\_COMP = - b\_diagnosis1 - b\_diagnosis2 - b\_diagnosis3  
 ) %>%  
 pivot\_longer(cols = starts\_with("b\_"), names\_to = "coef", values\_to = "estimate") %>%  
 filter(coef != "b\_Intercept") %>%  
 mutate(  
 coef = case\_match(coef,  
 "b\_cue1" ~ "Face",  
 "b\_diagnosis1" ~ "ADHD",  
 "b\_diagnosis2" ~ "ASD",  
 "b\_diagnosis3" ~ "ADHD+ASD",  
 "b\_COMP" ~ "COMP",  
 "b\_diagnosis1:cue1" ~ "Interaction: ADHD x cue",  
 "b\_diagnosis2:cue1" ~ "Interaction: ASD x cue",  
 "b\_diagnosis3:cue1" ~ "Interaction: ADHD+ASD x cue"  
 ),  
 coef = fct\_reorder(coef, desc(estimate))  
 ) %>%   
 group\_by(coef) %>%  
 mutate(  
 cred = case\_when(  
 (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |  
 (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")



# H2b: ASD(face) > COMP(face)  
h2b = hypothesis(m.lat,   
 "0 < diagnosis1 + diagnosis3 + 2\*diagnosis2 +  
 diagnosis1:cue1 + diagnosis3:cue1 + 2\*diagnosis2:cue1")  
h2b

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(diagnosis1+d... < 0 0.01 0.1 -0.17 0.18 0.9  
## Post.Prob Star  
## 1 0.47   
## ---  
## 'CI': 90%-CI for one-sided and 95%-CI for two-sided hypotheses.  
## '\*': For one-sided hypotheses, the posterior probability exceeds 95%;  
## for two-sided hypotheses, the value tested against lies outside the 95%-CI.  
## Posterior probabilities of point hypotheses assume equal prior probabilities.

# explore: faster towards faces  
e = hypothesis(m.lat, "0 > 2\*cue1", alpha = 0.025)  
e

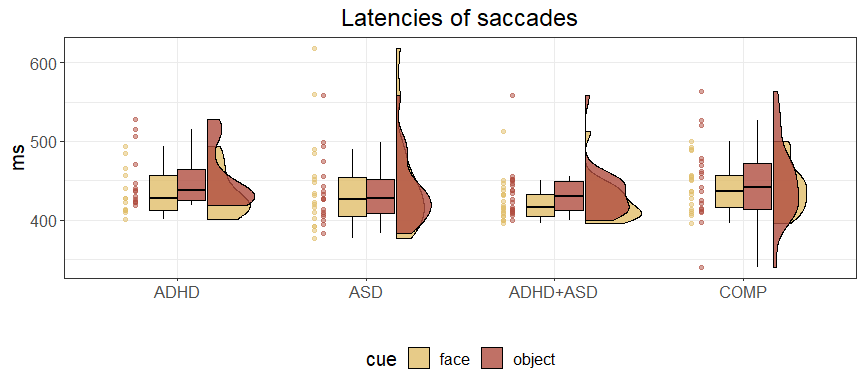
## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio Post.Prob  
## 1 (0)-(2\*cue1) > 0 0.04 0.02 0 0.09 39.4 0.98  
## Star  
## 1 \*  
## ---  
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.  
## '\*': For one-sided hypotheses, the posterior 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   
df.new = df.lat %>%   
 select(diagnosis, cue) %>%   
 distinct() %>%  
 mutate(  
 condition = paste(diagnosis, cue, sep = "\_")  
 )  
df.ms = as.data.frame(  
 fitted(m.lat, summary = F,   
 newdata = df.new %>% select(diagnosis, cue),   
 re\_formula = NA))  
colnames(df.ms) = df.new$condition  
  
# calculate our difference columns  
df.ms = df.ms %>%  
 mutate(  
 e = rowMeans(select(., matches(".\*\_object")), na.rm = T) -   
 rowMeans(select(., matches(".\*\_face")), na.rm = T)  
 )

Our hypothesis that saccades towards the faces have a longer latency in ASD than COMP adults was not confirmed by the data (*estimate* = 0.01 [-0.17, 0.18], *posterior probability* = 47.24%). However, our exploration revealed that latencies of saccades during a trial where the target appeared at the side of the face were faster regardless of the diagnostic group (*estimate* = 0.04 [0, 0.09], *posterior probability* = 97.52%). Specifically, the model predicted that if a saccade was produced towards a target appearing on the side of the face the latency was 5.556ms [0.008, 11.179] shorter than if a saccade was produced towards a target appearing at the side of the object cue.

### Plots

# rain cloud plot for the   
df.lat.agg %>%  
 mutate(  
 diagnosis = recode(diagnosis, "BOTH" = "ADHD+ASD")  
 ) %>%   
 ggplot(aes(diagnosis, lat, fill = cue, colour = cue)) + #  
 geom\_rain(rain.side = 'r',  
boxplot.args = list(color = "black", outlier.shape = NA, show\_guide = FALSE, alpha = .8),  
violin.args = list(color = "black", outlier.shape = NA, alpha = .8),  
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.col2) +  
 scale\_color\_manual(values = custom.col2) +  
 labs(title = "Latencies of saccades", x = "", y = "ms") +  
 theme\_bw() +   
 theme(legend.position = "bottom",   
 plot.title = element\_text(hjust = 0.5),   
 legend.direction = "horizontal",   
 text = element\_text(size = 15))

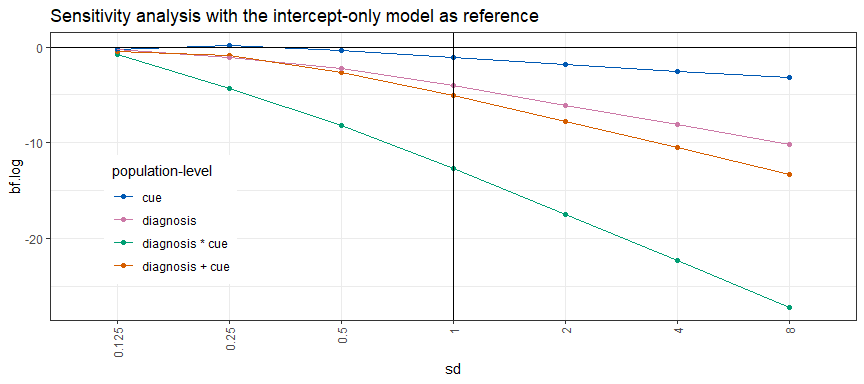


ggsave("Fig6\_latSac.pdf",   
 units = "mm",   
 width = 170,  
 height = 100,  
 dpi = 300)

## Bayes factor analysis

To complement our hypothesis testing using brms::hypothesis(), we perform a Bayes Factor analysis with models excluding some of our population-level predictors.

# set the directory in which to save results  
sense\_dir = file.path(getwd(), "\_brms\_sens\_cache")  
log\_dir = sense\_dir  
main.code = "lat-agg"  
  
# describe priors  
pr.descriptions = c("chosen",  
 "sdx2", "sdx4", "sdx8",   
 "sdx0.5", "sdx0.25", "sdx0.125"  
 )  
  
# check which have been run already  
if (file.exists(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)))) {  
 pr.done = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F) %>%  
 select(priors) %>% distinct()  
 pr.descriptions = pr.descriptions[!(pr.descriptions %in% pr.done$priors)]  
}  
  
if (length(pr.descriptions) > 0) {  
 # rerun the model with more iterations for bridgesampling  
 set.seed(5599)  
 m.lat.bf = brm(f.lat,  
 df.lat.agg, prior = priors,  
 iter = 40000, warmup = 10000,  
 backend = "cmdstanr", threads = threading(8),  
 family = "shifted\_lognormal",  
 file = "m\_lat\_agg\_bf",  
 save\_pars = save\_pars(all = TRUE)  
 )  
}  
  
# loop through them  
for (pr.desc in pr.descriptions) {  
 tryCatch({  
 # use the function  
 bf\_sens\_2int(m.lat.bf, "diagnosis", "cue", pr.desc,   
 main.code, # prefix for all models and MLL  
 file.path(log\_dir, "log\_FAB\_bf.txt"), # log file  
 sense\_dir, # where to save the models and MLL  
 reps = 5  
 )  
 },  
 error = function(err) {  
 message(sprintf("Error for %s: %s", pr.desc, err))  
 }  
 )  
}  
  
# read in the results  
df.lat.bf = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F)  
  
# check the sensitivity analysis result per model  
df.lat.bf %>%  
 filter(`population-level` != "1") %>%  
 mutate(  
 sd = as.factor(case\_when(  
 priors == "chosen" ~ "1",   
 substr(priors, 1, 3) == "sdx" ~ gsub("sdx", "", priors),  
 T ~ priors)  
 ),  
 order = case\_when(  
 priors == "chosen" ~ 1,   
 substr(priors, 1, 3) == "sdx" ~ as.numeric(gsub("sdx", "", priors)),  
 T ~ 999),  
 sd = fct\_reorder(sd, order)  
 ) %>%  
 ggplot(aes(y = bf.log,   
 x = sd,   
 group = `population-level`,   
 colour = `population-level`)) +   
 geom\_point() +  
 geom\_line() +   
 geom\_vline(xintercept = "1") +  
 geom\_hline(yintercept = 0) +  
 ggtitle("Sensitivity analysis with the intercept-only model as reference") +  
 #facet\_wrap(. ~ `population-level`, scales = "free\_y") +  
 scale\_colour\_manual(values = custom.col) +  
 theme\_bw() +  
 theme(legend.position = c(0.15,0.35),   
 axis.text.x = element\_text(angle = 90, vjust = 0.5, hjust = 1))



# print the BFs based on chosen priors  
kable(df.lat.bf %>% filter(priors == "chosen") %>% select(-priors) %>%  
 filter(`population-level` != "1") %>% arrange(desc(bf.log)), digits = 3)

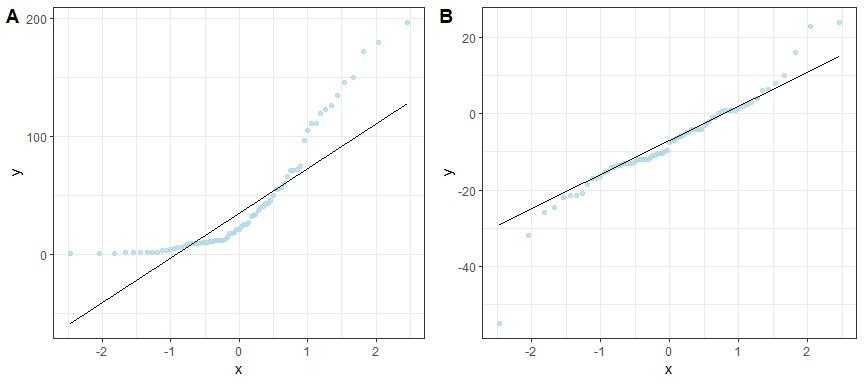
| population-level | bf.log |
| --- | --- |
| cue | -1.094 |
| diagnosis | -4.043 |
| diagnosis + cue | -5.101 |
| diagnosis \* cue | -12.680 |

Despite the effect of cue revealed by our explorative analysis, the Bayes Factor analysis revealed that the intercept-only model consistently outperformed all other models (diagnostic groups: log(*BF*) = -4.043; cue: log(*BF*) = -1.094; diagnostic group and cue: log(*BF*) = -5.101; full model: log(*BF*) = -12.68).

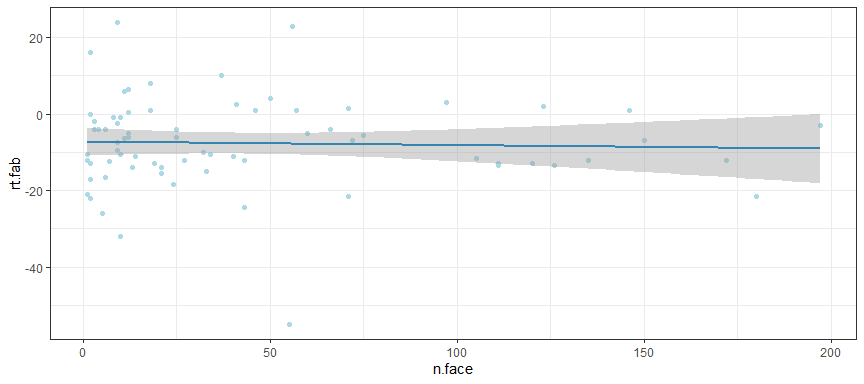
# S2.4 Correlation with reaction times: number of saccades

Last, we hypothesised that the FAB effect on reaction times may be associated with saccades produced towards the face. To investigate this, we use a Bayesian Spearman correlation as both FAB effect and number of saccades are not normally distributed.

# only keep saccades towards faces  
df.diff = df.cnt %>% filter(dir\_face == TRUE)  
  
# check the distribution plot > not normally distributed  
p1 = ggplot(df.diff, aes(sample = n.face)) +   
 stat\_qq(alpha = 0.75, colour = "lightblue") +   
 stat\_qq\_line() +   
 theme\_bw()  
p2 = ggplot(df.diff, aes(sample = rt.fab)) +   
 stat\_qq(alpha = 0.75, colour = "lightblue") +   
 stat\_qq\_line() +   
 theme\_bw()  
ggarrange(p1, p2,   
 nrow = 1, ncol = 2, labels = "AUTO")



# do a Bayesian Spearman correlation: https://osf.io/j5wud  
source("./helpers/rankBasedCommonFunctions.R")  
source("./helpers/spearmanSampler.R")  
  
# Default beta prior width is set to a = b = 1 for the sampler   
if (file.exists("CNT\_rho.rds")) {  
 rhoSamples.cnt = readRDS("CNT\_rho.rds")  
} else {  
 rhoSamples.cnt =   
 spearmanGibbsSampler(xVals = df.diff$n.face,  
 yVals = df.diff$rt.fab,   
 nSamples = 5e3)  
 saveRDS(rhoSamples.cnt, file = "CNT\_rho.rds")  
}  
  
# give the posterior samples for rho to the function below to compute BF01  
cor.bf = computeBayesFactorOneZero(rhoSamples.cnt$rhoSamples,   
 whichTest = "Spearman",  
 priorParameter = 1)  
  
# visualise it  
ggplot(data = df.diff, aes(x = n.face, y = rt.fab)) +  
 geom\_point(colour = "lightblue") +  
 geom\_smooth(method = "lm",   
 formula = y ~ x,   
 geom = "smooth", colour = c\_mid\_highlight) +  
 theme\_bw()



Furthermore, we assessed the relationship between FAB and number of saccades produced towards the face on the participant level (see supplementary materials S2.4). We used a Bayesian Spearman correlation due to both values not being normally distributed. This model revealed no association between number of saccades and face attention bias, in fact there was moderate evidence against an association between the number of saccades and face attention bias (log(*BF*) = -1.898).

# S2.5 Exploration: number of saccades towards cue

Additionally to our hypotheses, we also explored any effects of diagnostic status, cue type and their interaction on the number of saccades during the presentation of the cues.

## Specify the model

code = "CNT-cue"  
  
# set the formula  
f.cnt = brms::bf(n.cue ~ diagnosis \* dir\_face + (1 | subID))  
  
# set priors based on study design  
priors = c(  
 prior(normal(3, 1.5), class = Intercept),   
 prior(normal(0, 1.0), class = sd),  
 prior(normal(0, 1.0), class = b)  
)  
  
# set number of iterations and warmup for models  
iter = 4500  
warm = 1500

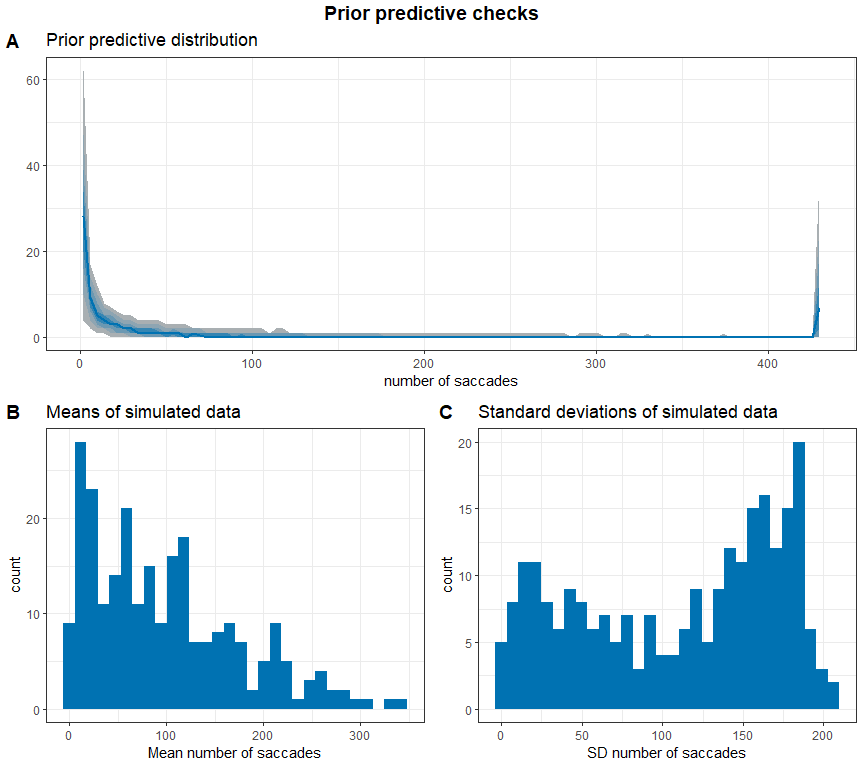
# check if the SBC already exists  
if (file.exists(file.path(cache\_dir, sprintf("df\_res\_%s.rds", code)))) {  
 # load in the results of the SBC  
 df.results = readRDS(file.path(cache\_dir, sprintf("df\_res\_%s.rds", code)))  
 df.backend = readRDS(file.path(cache\_dir, sprintf("df\_div\_%s.rds", code)))  
 dat = readRDS(file.path(cache\_dir, sprintf("dat\_%s.rds", code)))  
} else {  
 # perform the SBC  
 set.seed(2468)  
 gen = SBC\_generator\_brms(f.cnt, data = df.cnt.cue, prior = priors,   
 thin = 50, warmup = 10000, refresh = 2000,  
 generate\_lp = TRUE, family = poisson(), init = 0.1)  
 bck = SBC\_backend\_brms\_from\_generator(gen, chains = 4, thin = 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)))  
 saveRDS(res$stats,   
 file = file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 saveRDS(res$backend\_diagnostics,   
 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 5 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.

## Prior predictive checks

Next, we can plot the simulated values to perform prior predictive checks.

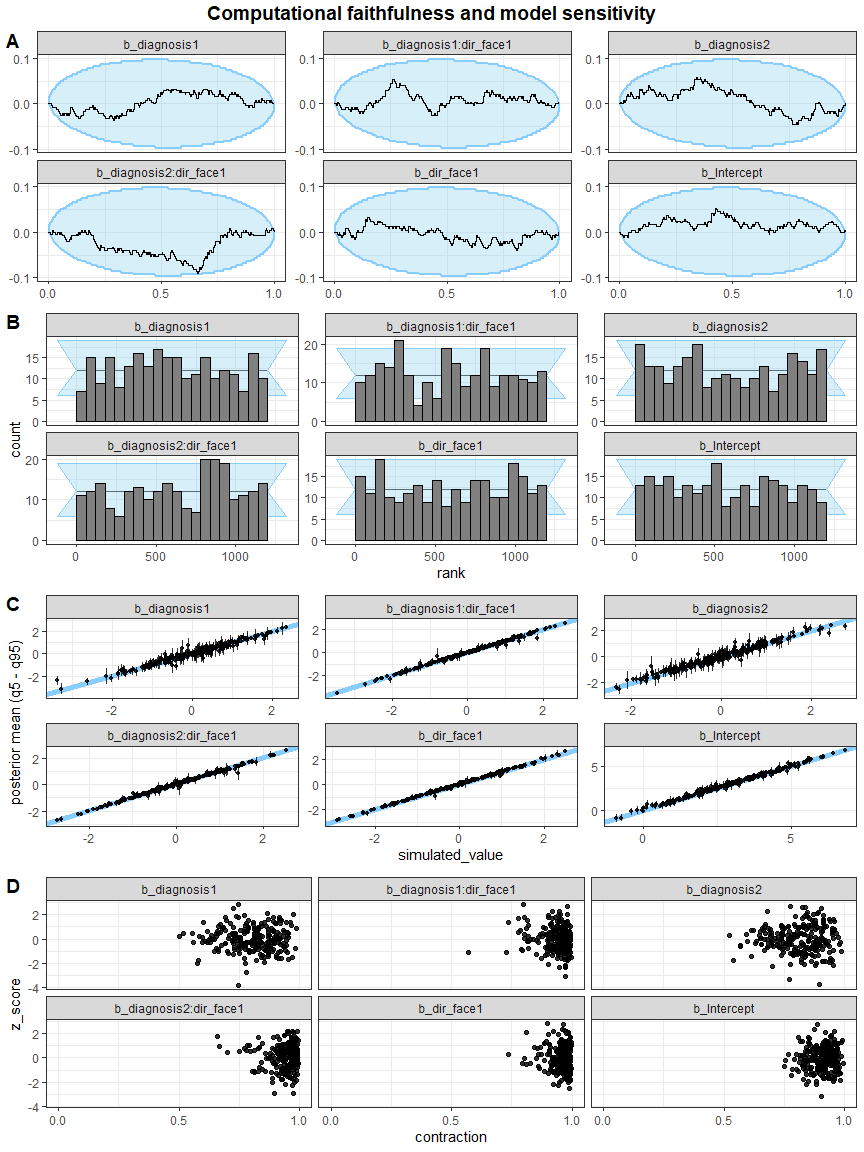
# get the true values  
truePars = dat[["variables"]]  
  
# create a matrix out of generated data  
dvname = gsub(" ", "", gsub("[\\|~].\*", "", f.cnt)[1])  
dvfakemat = matrix(NA, nrow(dat[['generated']][[1]]), length(dat[['generated']]))   
for (i in 1:length(dat[['generated']])) {  
 dvfakemat[,i] = dat[['generated']][[i]][[dvname]]  
}  
  
# set very large data points to a value of 432  
dvfakematH = dvfakemat;   
dvfakematH[dvfakematH > 432] = 432  
# compute one histogram per simulated data-set   
breaks = seq(0, max(dvfakematH, na.rm=T), length.out = 100)   
binwidth = round(breaks[2] - breaks[1])  
breaks = seq(0, max(dvfakematH, na.rm=T), binwidth)   
histmat = matrix(NA, ncol = nrow(truePars) + binwidth, nrow = length(breaks)-1)   
for (i in 1:nrow(truePars)) {  
 histmat[,i] = hist(dvfakematH[,i], breaks = breaks, plot = F)$counts   
}  
# for each bin, compute quantiles across histograms   
probs = seq(0.1, 0.9, 0.1)   
quantmat= as.data.frame(matrix(NA, nrow=dim(histmat)[1], ncol = length(probs)))  
names(quantmat) = paste0("p", probs)  
for (i in 1:dim(histmat)[1]) {  
 quantmat[i,] = quantile(histmat[i,], p = probs, na.rm = T)  
}  
quantmat$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 = "Prior predictive distribution", y = "", x = "number of saccades") +  
 theme\_bw()  
  
tmpM = apply(dvfakematH, 2, mean) # mean   
tmpSD = apply(dvfakematH, 2, sd)   
p2 = ggplot() +   
 stat\_bin(aes(x = tmpM), fill = c\_dark) +   
 labs(x = "Mean number of saccades", title = "Means of simulated data") +  
 theme\_bw()  
p3 = ggplot() +   
 stat\_bin(aes(x = tmpSD), fill = c\_dark) +   
 labs(x = "SD number of saccades", title = "Standard deviations of simulated data") +  
 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", face = "bold", size = 14))



Again, we set our priors deliberately wide.

## Computational faithfulness and model sensitivity

# get simulation numbers with issues  
des\_rank = max(df.results$max\_rank)  
check = merge(df.results %>%   
 group\_by(sim\_id) %>%   
 summarise(  
 rhat = max(rhat, na.rm = T),   
 mean\_rank = max(max\_rank)  
 ) %>%   
 filter(rhat >= 1.05 | mean\_rank < des\_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$sim\_id)) %>%  
 ungroup() %>%  
 mutate(  
 max\_rank = max(rank)  
 )  
p1 = plot\_ecdf\_diff(df.results.b) + theme\_bw() + theme(legend.position = "none") +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p2 = plot\_rank\_hist(df.results.b, bins = 20) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p3 = plot\_sim\_estimated(df.results.b, alpha = .8) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p4 = plot\_contraction(df.results.b,   
 prior\_sd = setNames(  
 c(as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "Intercept",]$prior)),   
 rep(  
 as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "b",]$prior)),  
 length(unique(df.results.b$variable))-1)),   
 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))



All looks acceptable.

## Posterior predictive checks

As the next step, we fit the model and check whether the chains have converged, which they seem to have. We then perform posterior predictive checks on the model using the bayesplot package.

# fit the model  
set.seed(6644)  
m.cnt = brm(f.cnt,  
 df.cnt.cue, prior = priors,  
 iter = iter, warmup = warm,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt-cue",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
rstan::check\_hmc\_diagnostics(m.cnt$fit)

##   
## Divergences:

## 0 of 12000 iterations ended with a divergence.

##   
## Tree depth:

## 0 of 12000 iterations saturated the maximum tree depth of 10.

##   
## Energy:

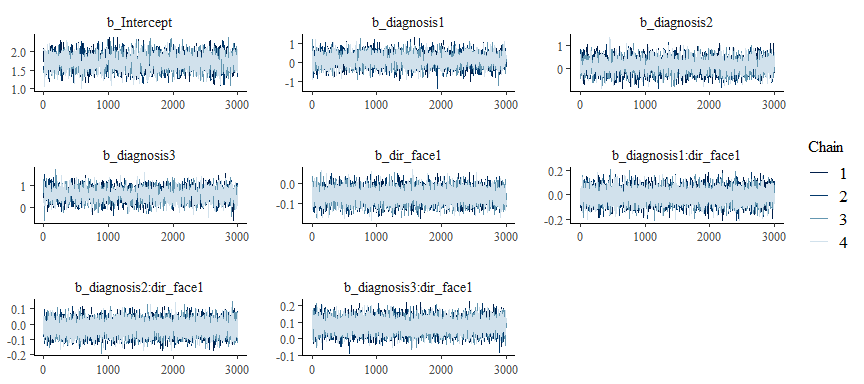
## E-BFMI indicated no pathological behavior.

# check that rhats are below 1.01  
sum(brms::rhat(m.cnt) >= 1.01, na.rm = T)

## [1] 0

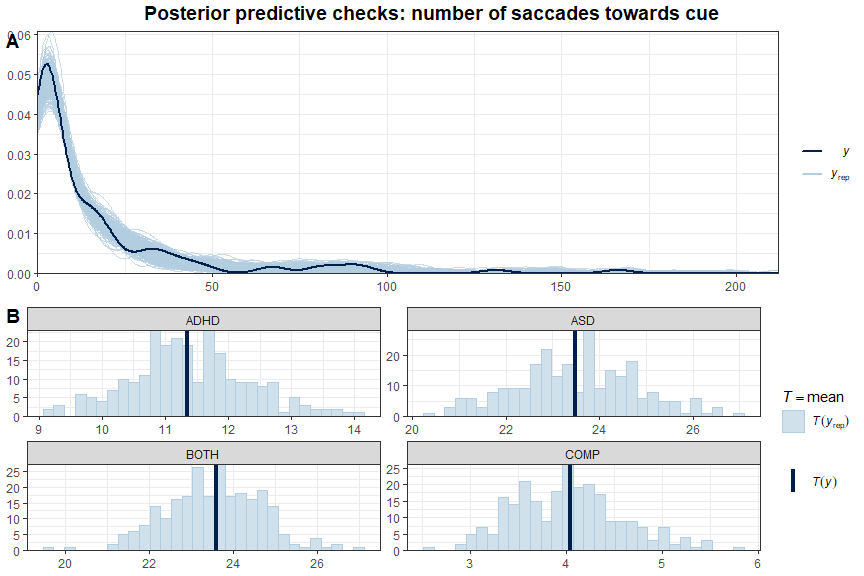
# check the trace plots  
post.draws = as\_draws\_df(m.cnt)  
mcmc\_trace(post.draws, regex\_pars = "^b\_",  
 facet\_args = list(ncol = 3)) +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))

## Scale for x is already present.  
## Adding another scale for x, which will replace the existing scale.



This model has no divergent samples and no rhats that are higher or equal to 1.01. Therefore, we go ahead and perform our posterior predictive checks.

# get the posterior predictions  
post.pred = posterior\_predict(m.cnt, ndraws = nsim)  
  
# check the fit of the predicted data compared to the real data  
p1 = pp\_check(m.cnt, ndraws = nsim) +   
 theme\_bw()  
  
# distributions of means and sds compared to the real values per group  
p2 = ppc\_stat\_grouped(df.cnt.cue$n.cue, post.pred, df.cnt.cue$diagnosis) +   
 theme\_bw()  
  
p = ggarrange(p1, p2,  
 nrow = 2, ncol = 1, labels = "AUTO")  
annotate\_figure(p,   
 top = text\_grob("Posterior predictive checks: number of saccades towards cue",   
 face = "bold", size = 14))



The predictions based on the model capture the data well. This further increased our trust in the model.

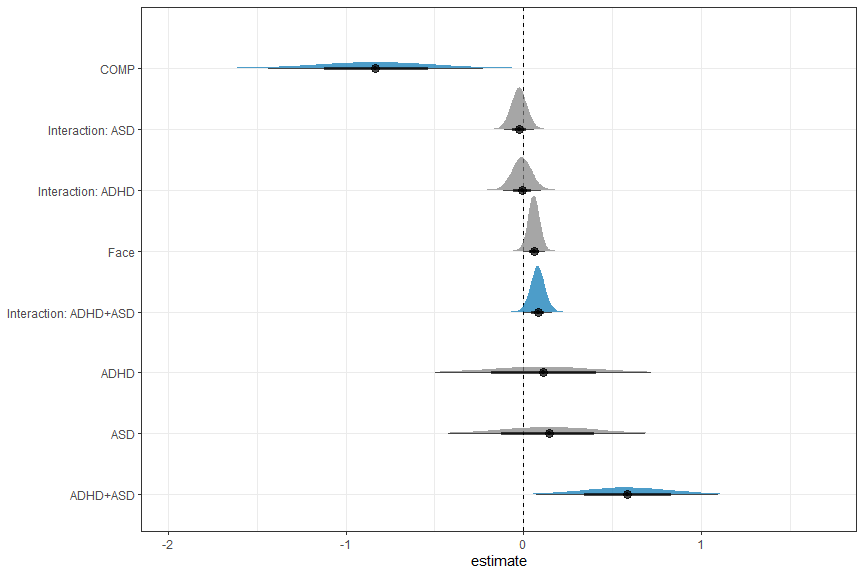
## Inferences

Now that we are convinced that we can trust our model, we have a look at the model and its estimates.

# print a summary  
summary(m.cnt)

## Family: poisson   
## Links: mu = log   
## Formula: n.cue ~ diagnosis \* dir\_face + (1 | subID)   
## Data: df.cnt.cue (Number of observations: 115)   
## Draws: 4 chains, each with iter = 4500; warmup = 1500; thin = 1;  
## total post-warmup draws = 12000  
##   
## Group-Level Effects:   
## ~subID (Number of levels: 64)   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## sd(Intercept) 1.30 0.13 1.07 1.59 1.00 2742 4947  
##   
## Population-Level Effects:   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS  
## Intercept 1.73 0.18 1.37 2.08 1.00 2546  
## diagnosis1 0.11 0.31 -0.50 0.72 1.00 2435  
## diagnosis2 0.14 0.28 -0.42 0.68 1.00 2466  
## diagnosis3 0.58 0.26 0.07 1.10 1.00 1907  
## dir\_face1 -0.06 0.03 -0.12 0.00 1.00 10629  
## diagnosis1:dir\_face1 -0.01 0.06 -0.12 0.10 1.00 21186  
## diagnosis2:dir\_face1 -0.02 0.04 -0.11 0.06 1.00 13804  
## diagnosis3:dir\_face1 0.08 0.04 0.00 0.16 1.00 13098  
## Tail\_ESS  
## Intercept 4551  
## diagnosis1 4276  
## diagnosis2 4867  
## diagnosis3 3785  
## dir\_face1 9014  
## diagnosis1:dir\_face1 8065  
## diagnosis2:dir\_face1 9655  
## diagnosis3:dir\_face1 9501  
##   
## 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.cnt = as\_draws\_df(m.cnt) %>%   
 select(starts\_with("b\_")) %>%  
 mutate(  
 b\_COMP = - b\_diagnosis1 - b\_diagnosis2 - b\_diagnosis3,  
 ASD = b\_Intercept + b\_diagnosis2,  
 ADHD = b\_Intercept + b\_diagnosis1,  
 BOTH = b\_Intercept + b\_diagnosis3,  
 b\_dir\_faceTRUE = - b\_dir\_face1,  
 COMP = b\_Intercept + b\_COMP  
 )  
  
# plot the posterior distributions  
df.m.cnt %>%   
 select(starts\_with("b\_")) %>%  
 pivot\_longer(cols = starts\_with("b\_"), names\_to = "coef", values\_to = "estimate") %>%  
 filter(coef != "b\_Intercept" & coef != "b\_dir\_face1") %>%  
 mutate(  
 coef = case\_match(coef,  
 "b\_diagnosis1" ~ "ADHD",  
 "b\_diagnosis2" ~ "ASD",  
 "b\_diagnosis3" ~ "ADHD+ASD",  
 "b\_COMP" ~ "COMP",  
 "b\_dir\_faceTRUE" ~ "Face",  
 "b\_diagnosis1:dir\_face1" ~ "Interaction: ADHD",  
 "b\_diagnosis3:dir\_face1" ~ "Interaction: ADHD+ASD",  
 "b\_diagnosis2:dir\_face1" ~ "Interaction: ASD"  
 ),  
 coef = fct\_reorder(coef, desc(estimate))  
 ) %>%   
 group\_by(coef) %>%  
 mutate(  
 cred = case\_when(  
 (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |  
 (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")



# exploration: face > object  
e1 = hypothesis(m.cnt, "0 > 2\*dir\_face1", alpha = 0.025)  
e1

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*dir\_face1) > 0 0.12 0.06 -0.01 0.24 30.33  
## Post.Prob Star  
## 1 0.97   
## ---  
## '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.

# exploration: ADHD > COMP  
e2 = hypothesis(m.cnt, "0 < 2\*diagnosis1 + diagnosis2 + diagnosis3", alpha = 0.025)  
e2

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*diagnosis1... < 0 -0.94 0.52 -1.98 0.08 27.71  
## Post.Prob Star  
## 1 0.97   
## ---  
## '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.

# exploration: ASD > COMP  
e3 = hypothesis(m.cnt, "0 < 2\*diagnosis2 + diagnosis1 + diagnosis3", alpha = 0.025)  
e3

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*diagnosis2... < 0 -0.97 0.48 -1.91 -0.02 43.94  
## Post.Prob Star  
## 1 0.98 \*  
## ---  
## '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.

# exploration: BOTH > COMP  
e4 = hypothesis(m.cnt, "0 < diagnosis2 + diagnosis1 + 2\*diagnosis3", alpha = 0.025)  
e4

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(diagnosis2+d... < 0 -1.42 0.46 -2.31 -0.5 630.58  
## Post.Prob Star  
## 1 1 \*  
## ---  
## '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.

# exploration: ASD(face) > ASD(object)  
e5 = hypothesis(m.cnt, "0 > 2\*dir\_face1 + 2\*diagnosis2:dir\_face1", alpha = 0.025)  
e5

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*dir\_face1+... > 0 0.17 0.08 0.01 0.33 53.3  
## Post.Prob Star  
## 1 0.98 \*  
## ---  
## '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.

# exploration: BOTH(face) > BOTH(object)  
e6 = hypothesis(m.cnt, "0 > 2\*dir\_face1 + 2\*diagnosis3:dir\_face1", alpha = 0.025)  
e6

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*dir\_face1+... > 0 -0.04 0.07 -0.17 0.09 0.36  
## Post.Prob Star  
## 1 0.26   
## ---  
## '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.

# exploration: ADHD(face) > ADHD(object)  
e7 = hypothesis(m.cnt, "0 > 2\*dir\_face1 + 2\*diagnosis1:dir\_face1", alpha = 0.025)  
e7

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*dir\_face1+... > 0 0.14 0.13 -0.12 0.38 6.21  
## Post.Prob Star  
## 1 0.86   
## ---  
## '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.

# exploration: COMP(face) > COMP(object)  
e8 = hypothesis(m.cnt, "0 > 2\*(dir\_face1 - diagnosis1:dir\_face1 -   
 diagnosis2:dir\_face1 - diagnosis3:dir\_face1)",  
 alpha = 0.025)  
e8

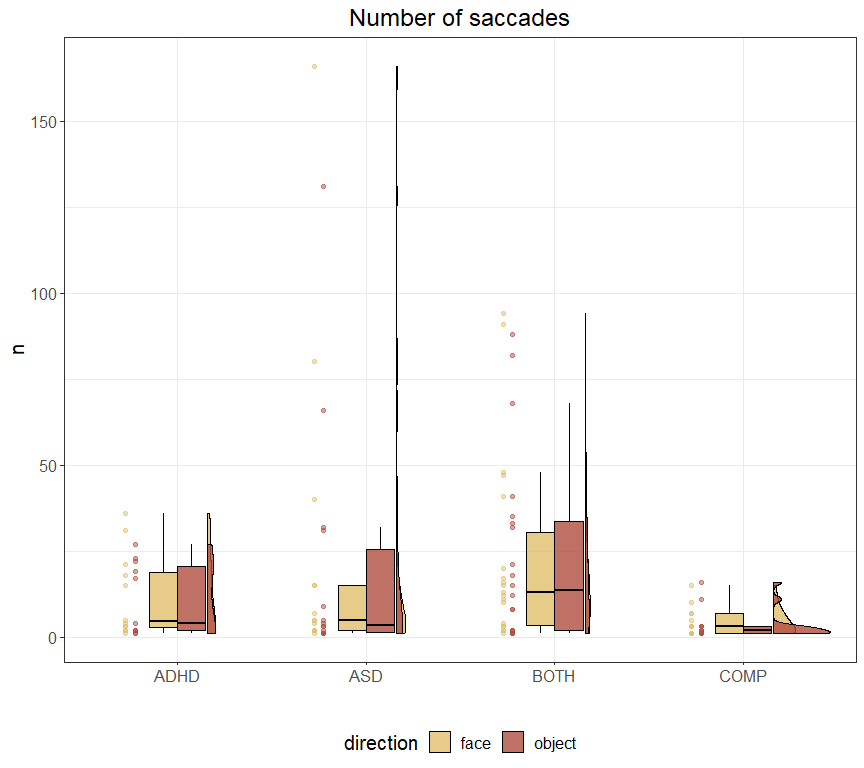
## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(2\*(dir\_face1... > 0 0.22 0.2 -0.17 0.61 6.1  
## Post.Prob Star  
## 1 0.86   
## ---  
## 'CI': 95%-CI for one-sided and 97.5%-CI for two-sided hypotheses.  
## '\*': For one-sided hypotheses, 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   
df.new = df.cnt.cue %>%   
 select(diagnosis, dir\_face) %>%   
 distinct() %>%  
 mutate(  
 condition = paste(diagnosis, dir\_face, sep = "\_")  
 )  
df.est = as.data.frame(  
 fitted(m.cnt, summary = F,   
 newdata = df.new %>% select(diagnosis, dir\_face),   
 re\_formula = NA))  
colnames(df.est) = df.new$condition

Exploring on the saccades elicited by the presentation of the cue, we find that our ADHD+ASD and our ASD group produced more saccades than our COMP group (ADHD+ASD vs. COMP: *estimate* = -1.42 [-2.31, -0.5], *posterior probability* = 99.84%; ASD vs. COMP: *estimate* = -0.97 [-1.91, -0.02], *posterior probability* = 97.78%). There was no credible effect between the ADHD and the COMP group (*estimate* = -0.94 [-1.98, 0.08], *posterior probability* = 96.52%). Additionally, there was a credible increase of saccades produced towards the face compared to the object cues in the ADHD+ASD (*estimate* = -0.04 [-0.17, 0.09], *posterior probability* = 26.28%) and the ASD (*estimate* = 0.17 [0.01, 0.33], *posterior probability* = 98.16%) group but not in the COMP (*estimate* = 0.22 [-0.17, 0.61], *posterior probability* = 85.91%) or the ADHD group (*estimate* = 0.14 [-0.12, 0.38], *posterior probability* = 86.13%).

## Plots

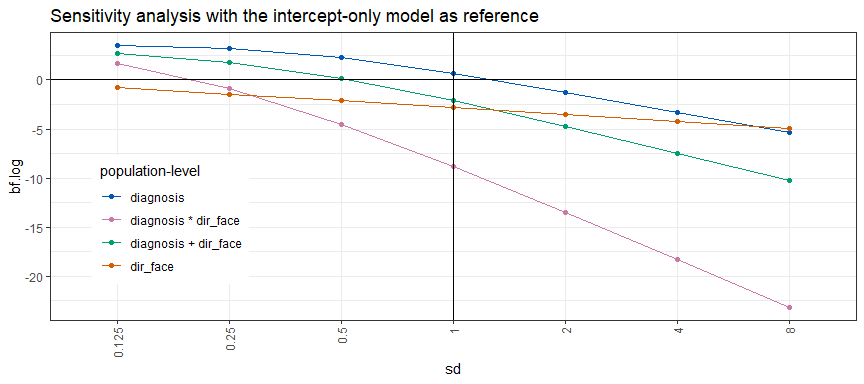
# rain cloud plot  
  
df.cnt.cue %>%  
 mutate(direction = if\_else(dir\_face == T, "face", "object")) %>%  
 ggplot(aes(diagnosis, n.cue, fill = direction, colour = direction)) + #  
 geom\_rain(rain.side = 'r',  
boxplot.args = list(color = "black", outlier.shape = NA, show\_guide = FALSE, alpha = .8),  
violin.args = list(color = "black", outlier.shape = NA, alpha = .8),  
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.col2) +  
 scale\_color\_manual(values = custom.col2) +  
 labs(title = "Number of saccades", x = "", y = "n") +  
 theme\_bw() +   
 theme(legend.position = "bottom",   
 plot.title = element\_text(hjust = 0.5),   
 legend.direction = "horizontal",   
 text = element\_text(size = 15))



## Bayes factor analysis

To complement our hypothesis testing using brms::hypothesis(), we perform a Bayes Factor analysis with models excluding some of our population-level predictors.

# set the directory in which to save results  
sense\_dir = file.path(getwd(), "\_brms\_sens\_cache")  
log\_dir = sense\_dir  
main.code = "cnt-cue"  
  
# describe priors  
pr.descriptions = c("chosen",  
 "sdx2", "sdx4", "sdx8",   
 "sdx0.5", "sdx0.25", "sdx0.125"  
 )  
  
# check which have been run already  
if (file.exists(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)))) {  
 pr.done = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F) %>%  
 select(priors) %>% distinct()  
 pr.descriptions = pr.descriptions[!(pr.descriptions %in% pr.done$priors)]  
}  
  
if (length(pr.descriptions) > 0) {  
 # rerun the model with more iterations for bridgesampling  
 set.seed(3311)  
 m.cnt.bf = brm(f.cnt,  
 df.cnt.cue, prior = priors,  
 iter = 40000, warmup = 10000,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt-cue\_bf",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
}  
  
# loop through them  
for (pr.desc in pr.descriptions) {  
 tryCatch({  
 # use the function  
 bf\_sens\_2int(m.cnt.bf, "diagnosis", "dir\_face", pr.desc,   
 main.code, # prefix for all models and MLL  
 file.path(log\_dir, "log\_FAB\_bf.txt"), # log file  
 sense\_dir, # where to save the models and MLL  
 reps = 5  
 )  
 },  
 error = function(err) {  
 message(sprintf("Error for %s: %s", pr.desc, err))  
 }  
 )  
}  
  
# read in the results  
df.cnt.bf = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F)  
  
# check the sensitivity analysis result per model  
df.cnt.bf %>%  
 filter(`population-level` != "1") %>%  
 mutate(  
 sd = as.factor(case\_when(  
 priors == "chosen" ~ "1",   
 substr(priors, 1, 3) == "sdx" ~ gsub("sdx", "", priors),  
 T ~ priors)  
 ),  
 order = case\_when(  
 priors == "chosen" ~ 1,   
 substr(priors, 1, 3) == "sdx" ~ as.numeric(gsub("sdx", "", priors)),  
 T ~ 999),  
 sd = fct\_reorder(sd, order)  
 ) %>%  
 ggplot(aes(y = bf.log,   
 x = sd,   
 group = `population-level`,   
 colour = `population-level`)) +   
 geom\_point() +  
 geom\_line() +   
 geom\_vline(xintercept = "1") +  
 geom\_hline(yintercept = 0) +  
 ggtitle("Sensitivity analysis with the intercept-only model as reference") +  
 #facet\_wrap(. ~ `population-level`, scales = "free\_y") +  
 scale\_colour\_manual(values = custom.col) +  
 theme\_bw() +  
 theme(legend.position = c(0.15,0.35),   
 axis.text.x = element\_text(angle = 90, vjust = 0.5, hjust = 1))



# print the BFs based on chosen priors  
kable(df.cnt.bf %>% filter(priors == "chosen") %>% select(-priors) %>%  
 filter(`population-level` != "1") %>% arrange(desc(bf.log)), digits = 3)

| population-level | bf.log |
| --- | --- |
| diagnosis | 0.666 |
| diagnosis + dir\_face | -2.138 |
| dir\_face | -2.817 |
| diagnosis \* dir\_face | -8.854 |

However, in the Bayes Factor analysis, the model including the diagnosis performs only slightly better than the intercept-only model for our chosen priors, with anecdotal evidence in favour of the model including diagnostic group (comparisons with intercept-only model, diagnostic groups: log(*BF*) = 0.666; saccade direction: log(*BF*) = -2.817; diagnostic group and saccade direction: log(*BF*) = -2.138; full model: log(*BF*) = -8.854). The sensitivity analysis shows that this effect is stable for priors which were narrower than our deliberately wide chosen priors.

# S2.6 Exploration: number of saccades towards target

Similarly, we explore the number of saccades during the target presentation and if the number of saccades differs depending on whether the target appears on the side of the face or the object cue.

## Specify the model

code = "CNT-tar"  
  
# set the formula  
f.cnt = brms::bf(n.tar ~ diagnosis \* cue + (1 | subID))  
  
# set priors based on study design  
priors = c(  
 prior(normal(3, 1.5), class = Intercept),   
 prior(normal(0, 1.0), class = sd),  
 prior(normal(0, 1.0), class = b)  
)  
  
# set number of iterations and warmup for models  
iter = 4500  
warm = 1500

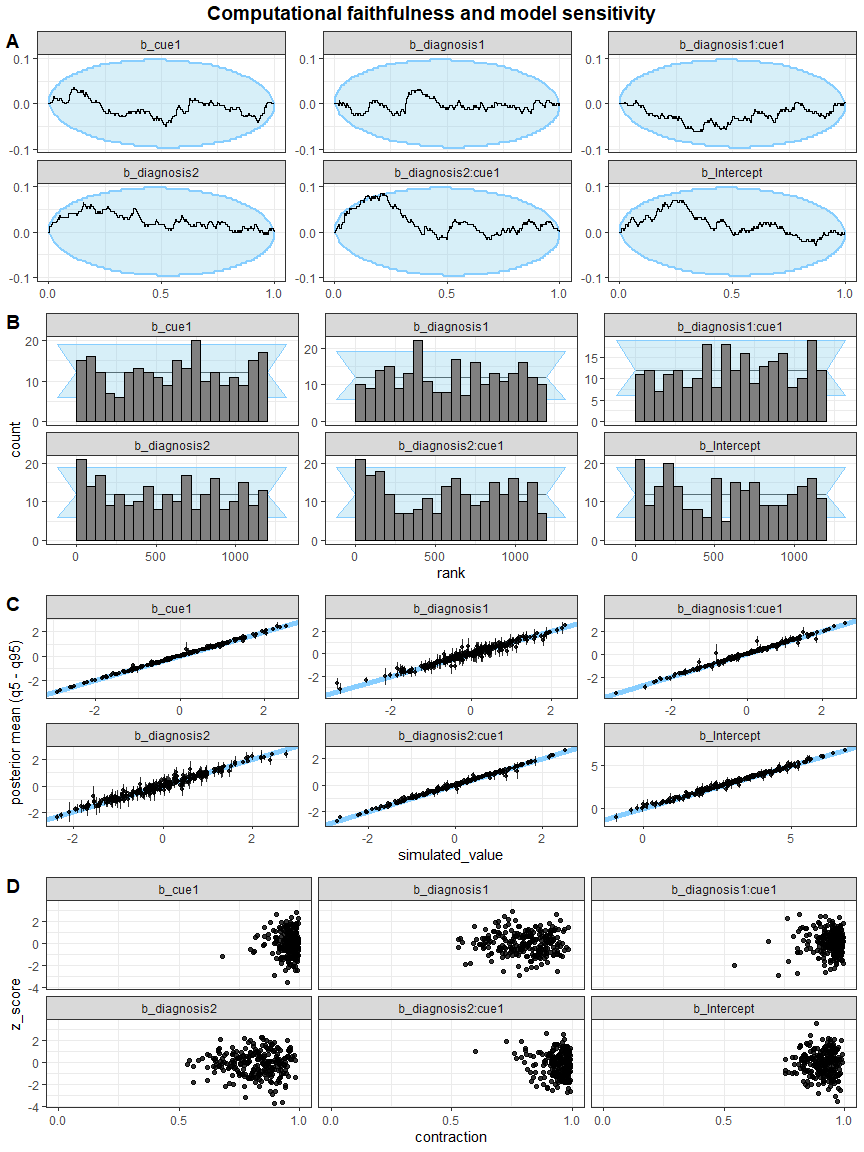
## SBC

# 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 {  
 # perform the SBC  
 set.seed(2468)  
 gen = SBC\_generator\_brms(f.cnt, data = df.cnt.tar, prior = priors,   
 thin = 50, warmup = 10000, refresh = 2000,  
 generate\_lp = TRUE, family = poisson(), init = 0.1)  
 bck = SBC\_backend\_brms\_from\_generator(gen, chains = 4, thin = 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)))  
 saveRDS(res$stats,   
 file = file.path(cache\_dir, paste0("df\_res\_", code, ".rds")))  
 saveRDS(res$backend\_diagnostics,   
 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 5 of 250 simulations had at least one parameter that had an rhat of at least 1.05, and only 1 models had divergent samples. This suggests that this model performs well.

## Computational faithfulness and model sensitivity

# get simulation numbers with issues  
des\_rank = max(df.results$max\_rank)  
check = merge(df.results %>%   
 group\_by(sim\_id) %>%   
 summarise(  
 rhat = max(rhat, na.rm = T),   
 mean\_rank = max(max\_rank)  
 ) %>%   
 filter(rhat >= 1.05 | mean\_rank < des\_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$sim\_id)) %>%  
 ungroup() %>%  
 mutate(  
 max\_rank = max(rank)  
 )  
p1 = plot\_ecdf\_diff(df.results.b) + theme\_bw() + theme(legend.position = "none") +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p2 = plot\_rank\_hist(df.results.b, bins = 20) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p3 = plot\_sim\_estimated(df.results.b, alpha = .8) + theme\_bw() +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))  
p4 = plot\_contraction(df.results.b,   
 prior\_sd = setNames(  
 c(as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "Intercept",]$prior)),   
 rep(  
 as.numeric(  
 gsub(".\*, (.+)\\).\*", "\\1",   
 priors[priors$class == "b",]$prior)),  
 length(unique(df.results.b$variable))-1)),   
 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))



This looks good.

## Posterior predictive checks

As the next step, we fit the model and check whether the chains have converged, which they seem to have. We then perform posterior predictive checks on the model using the bayesplot package.

# fit the maximal model  
set.seed(7744)  
m.cnt.tar = brm(f.cnt,  
 df.cnt.tar, prior = priors,  
 iter = iter, warmup = warm,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt-tar",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
rstan::check\_hmc\_diagnostics(m.cnt$fit)

##   
## Divergences:

## 0 of 12000 iterations ended with a divergence.

##   
## Tree depth:

## 0 of 12000 iterations saturated the maximum tree depth of 10.

##   
## Energy:

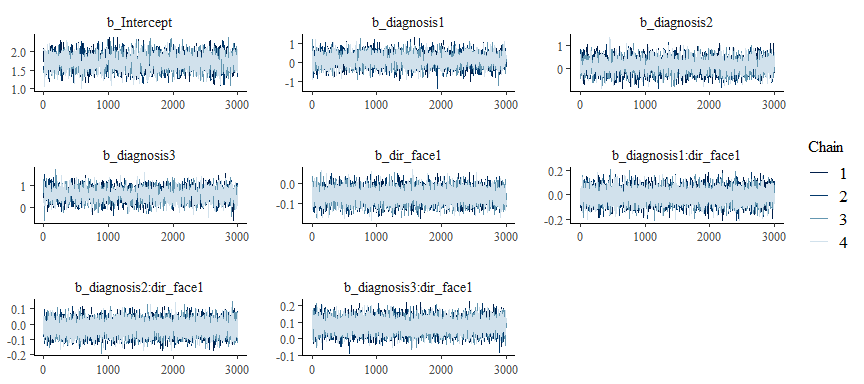
## E-BFMI indicated no pathological behavior.

# check that rhats are below 1.01  
sum(brms::rhat(m.cnt) >= 1.01, na.rm = T)

## [1] 0

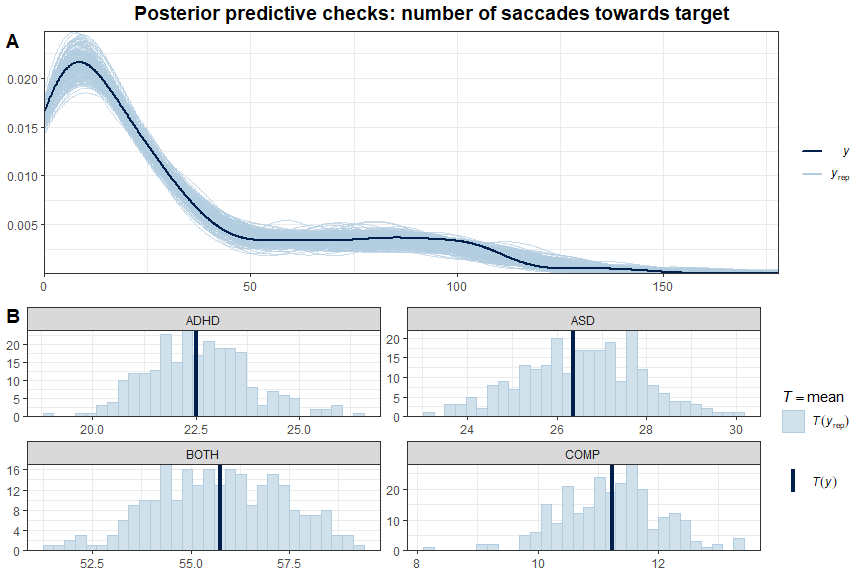
# check the trace plots  
post.draws = as\_draws\_df(m.cnt)  
mcmc\_trace(post.draws, regex\_pars = "^b\_",  
 facet\_args = list(ncol = 3)) +  
 scale\_x\_continuous(breaks=scales::pretty\_breaks(n = 3)) +  
 scale\_y\_continuous(breaks=scales::pretty\_breaks(n = 3))

## Scale for x is already present.  
## Adding another scale for x, which will replace the existing scale.



This model has no divergent samples and no rhats that are higher or equal to 1.01. Therefore, we go ahead and perform our posterior predictive checks.

# get the posterior predictions  
post.pred = posterior\_predict(m.cnt.tar, ndraws = nsim)  
  
# check the fit of the predicted data compared to the real data  
p1 = pp\_check(m.cnt.tar, ndraws = nsim) +   
 theme\_bw()  
  
# distributions of means and sds compared to the real values per group  
p2 = ppc\_stat\_grouped(df.cnt.tar$n.tar, post.pred, df.cnt.tar$diagnosis) +   
 theme\_bw()  
  
p = ggarrange(p1, p2,  
 nrow = 2, ncol = 1, labels = "AUTO")  
annotate\_figure(p,   
 top = text\_grob("Posterior predictive checks: number of saccades towards target",   
 face = "bold", size = 14))



The predictions based on the model capture the data well. This further increased our trust in the model.

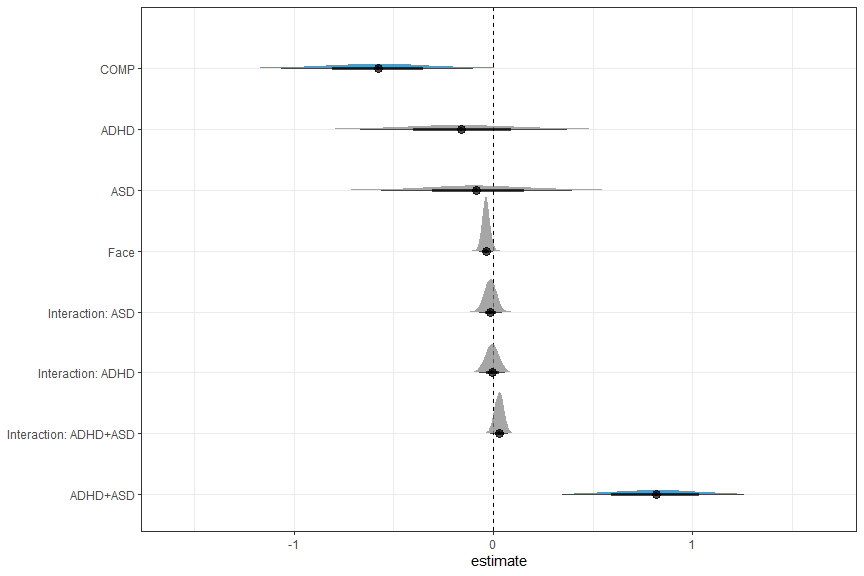
## Model summary

Now that we are convinced that we can trust our model, we have a look at the model and its estimates.

# print a summary  
summary(m.cnt.tar)

## Family: poisson   
## Links: mu = log   
## Formula: n.tar ~ diagnosis \* cue + (1 | subID)   
## Data: df.cnt.tar (Number of observations: 144)   
## Draws: 4 chains, each with iter = 4500; warmup = 1500; thin = 1;  
## total post-warmup draws = 12000  
##   
## Group-Level Effects:   
## ~subID (Number of levels: 76)   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## sd(Intercept) 1.22 0.11 1.03 1.47 1.00 2356 4302  
##   
## Population-Level Effects:   
## Estimate Est.Error l-95% CI u-95% CI Rhat Bulk\_ESS Tail\_ESS  
## Intercept 2.62 0.15 2.33 2.91 1.00 2146 3050  
## diagnosis1 -0.16 0.26 -0.67 0.37 1.00 1879 3287  
## diagnosis2 -0.08 0.24 -0.56 0.40 1.00 1953 2842  
## diagnosis3 0.82 0.23 0.35 1.26 1.00 1645 2689  
## cue1 -0.04 0.02 -0.07 0.00 1.00 12115 9440  
## diagnosis1:cue1 -0.00 0.03 -0.07 0.06 1.00 13961 9242  
## diagnosis2:cue1 -0.01 0.03 -0.07 0.04 1.00 13194 8965  
## diagnosis3:cue1 0.03 0.02 -0.02 0.08 1.00 11665 9265  
##   
## 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).

# plot the posterior distributions  
as\_draws\_df(m.cnt.tar) %>%   
 select(starts\_with("b\_")) %>%  
 mutate(  
 b\_COMP = - b\_diagnosis1 - b\_diagnosis2 - b\_diagnosis3  
 ) %>%  
 pivot\_longer(cols = starts\_with("b\_"), names\_to = "coef", values\_to = "estimate") %>%  
 filter(coef != "b\_Intercept") %>%  
 mutate(  
 coef = case\_match(coef,  
 "b\_diagnosis1" ~ "ADHD",  
 "b\_diagnosis2" ~ "ASD",  
 "b\_diagnosis3" ~ "ADHD+ASD",  
 "b\_COMP" ~ "COMP",  
 "b\_cue1" ~ "Face",  
 "b\_diagnosis1:cue1" ~ "Interaction: ADHD",  
 "b\_diagnosis2:cue1" ~ "Interaction: ASD",  
 "b\_diagnosis3:cue1" ~ "Interaction: ADHD+ASD"  
 ),  
 coef = fct\_reorder(coef, desc(estimate))  
 ) %>%   
 group\_by(coef) %>%  
 mutate(  
 cred = case\_when(  
 (mean(estimate) < 0 & quantile(estimate, probs = 0.975) < 0) |  
 (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")



# explore: face > object  
e1 = hypothesis(m.cnt.tar, "0 > 2\*cue1", alpha = 0.025)  
e1

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio Post.Prob  
## 1 (0)-(2\*cue1) > 0 0.07 0.04 0 0.14 31.09 0.97  
## Star  
## 1   
## ---  
## '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.

# explore: BOTH > COMP  
e2 = hypothesis(m.cnt.tar, "0 < diagnosis1 + diagnosis2 + 2\*diagnosis3", alpha = 0.025)  
e2

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(diagnosis1+d... < 0 -1.39 0.38 -2.15 -0.63 11999  
## Post.Prob Star  
## 1 1 \*  
## ---  
## '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.

# explore: BOTH > ADHD  
e3 = hypothesis(m.cnt.tar, "0 < -2\*diagnosis1 + 2\*diagnosis3", alpha = 0.025)  
e3

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(-2\*diagnosis... < 0 -1.94 0.81 -3.51 -0.31 105.19  
## Post.Prob Star  
## 1 0.99 \*  
## ---  
## '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.

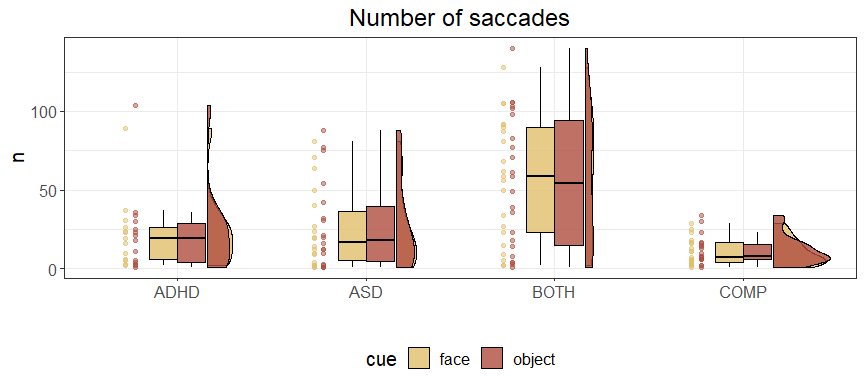
# explore: BOTH > ASD  
e4 = hypothesis(m.cnt.tar, "0 < -2\*diagnosis2 + 2\*diagnosis3", alpha = 0.025)  
e4

## Hypothesis Tests for class b:  
## Hypothesis Estimate Est.Error CI.Lower CI.Upper Evid.Ratio  
## 1 (0)-(-2\*diagnosis... < 0 -1.79 0.76 -3.24 -0.24 92.02  
## Post.Prob Star  
## 1 0.99 \*  
## ---  
## '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.

This model again suggests a larger number of saccades produced by the ADHD+ASD group than the other groups (ADHD+ASD vs. COMP: *estimate* = -1.39 [-2.15, -0.63], *posterior probability* = 99.99%; ADHD+ASD vs. ADHD: *estimate* = -1.94 [-3.51, -0.31], *posterior probability* = 99.06%; ADHD+ASD vs. ASD: *estimate* = -1.79 [-3.24, -0.24], *posterior probability* = 98.92%). There seems to be no credible effect of cue, i.e., whether the saccade was produced towards a target appearing on the side of the face or the object (*estimate* = 0.07 [0, 0.14], *posterior probability* = 96.88%).

## Plots

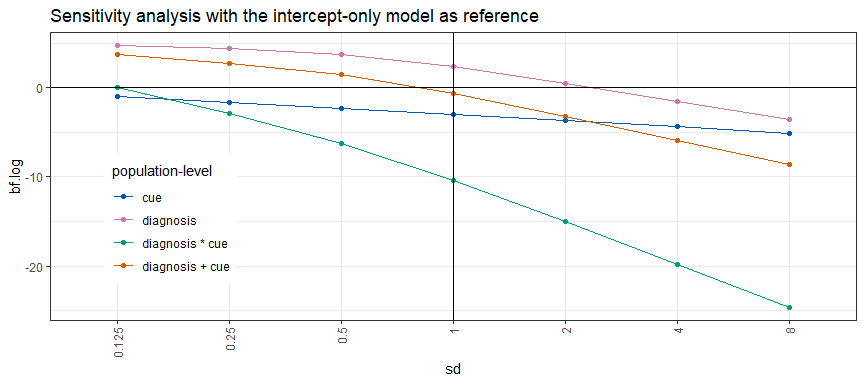
# rain cloud plot  
  
df.cnt.tar %>%  
 ggplot(aes(diagnosis, n.tar, fill = cue, colour = cue)) + #  
 geom\_rain(rain.side = 'r',  
boxplot.args = list(color = "black", outlier.shape = NA, show\_guide = FALSE, alpha = .8),  
violin.args = list(color = "black", outlier.shape = NA, alpha = .8),  
boxplot.args.pos = 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.col2) +  
 scale\_color\_manual(values = custom.col2) +  
 labs(title = "Number of saccades", x = "", y = "n") +  
 theme\_bw() +   
 theme(legend.position = "bottom", plot.title = element\_text(hjust = 0.5),   
 legend.direction = "horizontal",   
 text = element\_text(size = 15))



## Bayes factor analysis

To complement our hypothesis testing using brms::hypothesis(), we perform a Bayes Factor analysis with models excluding some of our population-level predictors.

# set the directory in which to save results  
sense\_dir = file.path(getwd(), "\_brms\_sens\_cache")  
log\_dir = sense\_dir  
main.code = "cnt-tar"  
  
# describe priors  
pr.descriptions = c("chosen",  
 "sdx2", "sdx4", "sdx8",   
 "sdx0.5", "sdx0.25", "sdx0.125"  
 )  
  
# check which have been run already  
if (file.exists(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)))) {  
 pr.done = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F) %>%  
 select(priors) %>% distinct()  
 pr.descriptions = pr.descriptions[!(pr.descriptions %in% pr.done$priors)]  
}  
  
if (length(pr.descriptions) > 0) {  
 # rerun the model with more iterations for bridgesampling  
 set.seed(2244)  
 m.cnt.bf = brm(f.cnt,  
 df.cnt.tar, prior = priors,  
 iter = 40000, warmup = 10000,  
 backend = "cmdstanr", threads = threading(8),  
 file = "m\_cnt-tar\_bf",  
 family = "poisson",   
 save\_pars = save\_pars(all = TRUE)  
 )  
}  
  
# loop through them  
for (pr.desc in pr.descriptions) {  
 tryCatch({  
 # use the function  
 bf\_sens\_2int(m.cnt.bf, "diagnosis", "cue", pr.desc,   
 main.code, # prefix for all models and MLL  
 file.path(log\_dir, "log\_FAB\_bf.txt"), # log file  
 sense\_dir, # where to save the models and MLL  
 reps = 5  
 )  
 },  
 error = function(err) {  
 message(sprintf("Error for %s: %s", pr.desc, err))  
 }  
 )  
}  
  
# read in the results  
df.cnt.bf = read\_csv(file.path(sense\_dir, sprintf("df\_%s\_bf.csv", main.code)),   
 show\_col\_types = F)  
  
# check the sensitivity analysis result per model  
df.cnt.bf %>%  
 filter(`population-level` != "1") %>%  
 mutate(  
 sd = as.factor(case\_when(  
 priors == "chosen" ~ "1",   
 substr(priors, 1, 3) == "sdx" ~ gsub("sdx", "", priors),  
 T ~ priors)  
 ),  
 order = case\_when(  
 priors == "chosen" ~ 1,   
 substr(priors, 1, 3) == "sdx" ~ as.numeric(gsub("sdx", "", priors)),  
 T ~ 999),  
 sd = fct\_reorder(sd, order)  
 ) %>%  
 ggplot(aes(y = bf.log,   
 x = sd,   
 group = `population-level`,   
 colour = `population-level`)) +   
 geom\_point() +  
 geom\_line() +   
 geom\_vline(xintercept = "1") +  
 geom\_hline(yintercept = 0) +  
 ggtitle("Sensitivity analysis with the intercept-only model as reference") +  
 #facet\_wrap(. ~ `population-level`, scales = "free\_y") +  
 scale\_colour\_manual(values = custom.col) +  
 theme\_bw() +  
 theme(legend.position = c(0.15,0.35),   
 axis.text.x = element\_text(angle = 90, vjust = 0.5, hjust = 1))



# print the BFs based on chosen priors  
kable(df.cnt.bf %>% filter(priors == "chosen") %>% select(-priors) %>%  
 filter(`population-level` != "1") %>% arrange(desc(bf.log)), digits = 3)

| population-level | bf.log |
| --- | --- |
| diagnosis | 2.387 |
| diagnosis + cue | -0.643 |
| cue | -3.004 |
| diagnosis \* cue | -10.405 |

These effects were mirrored by the Bayes Factor analysis where the model only including the main effect diagnostic group outperforms the other models with the exception of very wide priors (all comparisons with intercept-only model; diagnostic groups: log(*BF*) = 2.387; cue: log(*BF*) = -3.004; diagnostic group and cue: log(*BF*) = -0.643; full model: log(*BF*) = -10.405).