```
## Loading required package: StanHeaders

## Loading required package: ggplot2

## rstan (Version 2.21.8, GitRev: 2e1f913d3ca3)
```

## For execution on a local, multicore CPU with excess RAM we recommend calling
## options(mc.cores = parallel::detectCores()).
## To avoid recompilation of unchanged Stan programs, we recommend calling
## rstan\_options(auto\_write = TRUE)

```
options(mc.cores = parallel::detectCores())
rstan_options(auto_write = T)

# Get list of files in 'data_2' folder with the pattern "riskytimed"
files <- dir(path = "data_2", pattern="riskytimed")

# Read all csv files in the list
data_list <- lapply(paste0("data_2/", files), read.table, header = TRUE, skip = 0, fill = TRUE, sep= ";")

# Concatenate rows of all items in the list into a data frame
dat <- do.call("rbind", data_list)</pre>
```

```
# gamble characteristics
dat$eva = dat$oa1*dat$pa1+dat$oa2*dat$pa2 + dat$oa3*dat$pa3+dat$oa4*dat$pa4
dat$evb = dat$ob1*dat$pb1+dat$ob2*dat$pb2 + dat$ob3*dat$pb3+dat$ob4*dat$pb4
dat$evd = dat$evb - dat$eva
dat$sda = sqrt((dat$oa1-dat$eva)^2*dat$pa1 + (dat$oa2-dat$eva)^2*dat$pa2 + (dat$oa
3-dat$eva)^2*dat$pa3 + (dat$oa4-dat$eva)^2*dat$pa4)
dat$sdb = sqrt((dat$ob1-dat$evb)^2*dat$pb1 + (dat$ob2-dat$evb)^2*dat$pb2 + (dat$ob
3-dat$evb)^2*dat$pb3 + (dat$ob4-dat$evb)^2*dat$pb4)
dat$sdd = dat$sdb - dat$sda
dat$evdummy = ifelse(dat$evd>0,1,0)
```

```
# transform to +/- 1; safe - 1, risky +1
dat$cho <- ifelse(dat$choice==0,-1,ifelse(dat$choice==1,1,NA))
dat$cho2 <- ifelse(dat$choice==0,1,ifelse(dat$choice==1,0,NA))
ids <- unique(dat$id)
for(j in 1:length(ids)){
   dat$tid[dat$id==ids[j]] <- j
}
tids <- unique(dat$tid)
# only control data
control_dat <- dat[dat$cond=="control",]
# remove fast RTs
rcontrol_dat <- control_dat[control_dat$rt>1,]
# only condition no time pressure
dataList = list(cho = rcontrol_dat$cho, accuracy_flipped = rcontrol_dat$cho2, rt
= rcontrol_dat$rt, participant = rcontrol_dat$tid,N=nrow(rcontrol_dat), L = lengt
h(tids), evd = rcontrol_dat$evd, sdd = rcontrol_dat$sdd)
```

```
oa = as.matrix(rcontrol_dat[, c("oa1", "oa2", "oa3", "oa4")])
ob = as.matrix(rcontrol_dat[, c("ob1", "ob2", "ob3", "ob4")])
pa = as.matrix(rcontrol_dat[, c("pa1", "pa2", "pa3", "pa4")])
pb = as.matrix(rcontrol_dat[, c("pb1", "pb2", "pb3", "pb4")])
```

```
initFunc <-function (i) {</pre>
  initList=list()
  for (ll in 1:i){
    initList[[ll]] = list(
                          mu alpha = runif(1,-1.4587,2.5413),
                          sd_alpha = runif(1,0,1),
                          mu threshold = runif(1,-0.5, 2.5),
                          sd threshold = runif(1,0,1),
                          mu_ndt = runif(1, -1.5, 0),
                          sd_ndt = runif(1, 0, 1),
                          mu_{theta} = runif(1,0,6),
                          sd theta = runif(1,0,1),
                          mu_beta = runif(1,-5, 5),
                          sd_beta = runif(1, 0, 1),
                          z = runif(length(tids), -0.1, 0.1),
                          z theta = runif(length(tids),-0.1,0.1),
                          z_threshold = runif(length(tids),-0.1,0.1),
                          z_ndt = runif(length(tids),-0.1,0.1),
                          z_beta = runif(length(tids),-0.1,0.1)
    )
  }
  return(initList)
}
```

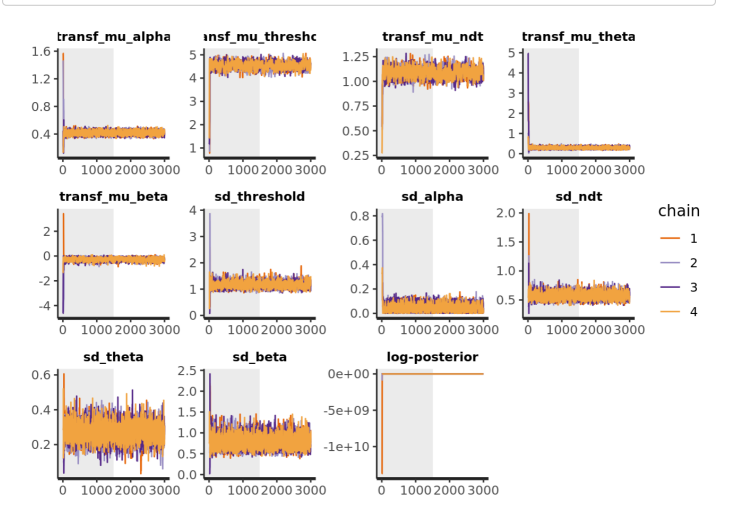
```
m <- stan_model("EU_discount.stan")</pre>
```

## recompiling to avoid crashing R session

```
## Trying to compile a simple C file
```

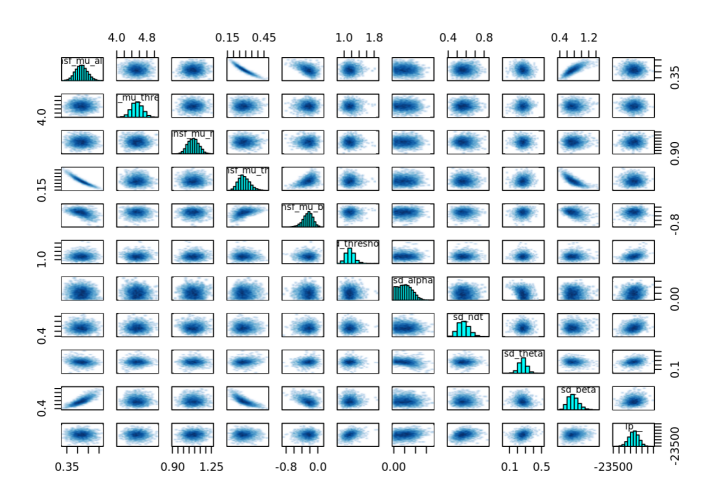
```
## Running /usr/lib/R/bin/R CMD SHLIB foo.c
## using C compiler: 'qcc (Ubuntu 11.4.0-1ubuntu1~22.04) 11.4.0'
## gcc -I"/usr/share/R/include" -DNDEBUG
                                         -I"/usr/local/lib/R/site-library/Rcpp/i
nclude/" -I"/usr/local/lib/R/site-library/RcppEigen/include/" -I"/usr/local/lib/
R/site-library/RcppEigen/include/unsupported" -I"/usr/local/lib/R/site-library/B
H/include" -I"/usr/local/lib/R/site-library/StanHeaders/include/src/" -I"/usr/loc
al/lib/R/site-library/StanHeaders/include/" -I"/usr/local/lib/R/site-library/Rcpp
Parallel/include/" -I"/usr/local/lib/R/site-library/rstan/include" -DEIGEN NO DEB
   -DBOOST_DISABLE_ASSERTS -DBOOST_PENDING_INTEGER_LOG2_HPP -DSTAN_THREADS -DB
OOST_NO_AUTO_PTR -include '/usr/local/lib/R/site-library/StanHeaders/include/sta
n/math/prim/mat/fun/Eigen.hpp' -D_REENTRANT -DRCPP_PARALLEL_USE_TBB=1
                                                                             -fpic
-g -O2 -ffile-prefix-map=/build/r-base-MHXHhT/r-base-4.3.1=. -fstack-protector-str
ong -Wformat -Werror=format-security -Wdate-time -D FORTIFY SOURCE=2 -c foo.c -o
## In file included from /usr/local/lib/R/site-library/RcppEigen/include/Eigen/Cor
e:88,
##
                    from /usr/local/lib/R/site-library/RcppEigen/include/Eigen/Den
se:1,
##
                    from /usr/local/lib/R/site-library/StanHeaders/include/stan/ma
th/prim/mat/fun/Eigen.hpp:13,
                    from <command-line>:
## /usr/local/lib/R/site-library/RcppEigen/include/Eigen/src/Core/util/Macros.h:62
8:1: error: unknown type name 'namespace'
     628 | namespace Eigen {
##
##
## /usr/local/lib/R/site-library/RcppEigen/include/Eigen/src/Core/util/Macros.h:62
8:17: error: expected '=', ',', ';', 'asm' or '__attribute__' before '{' token
     628 | namespace Eigen {
##
##
## In file included from /usr/local/lib/R/site-library/RcppEigen/include/Eigen/Den
se:1,
##
                    from /usr/local/lib/R/site-library/StanHeaders/include/stan/ma
th/prim/mat/fun/Eigen.hpp:13,
##
                    from <command-line>:
## /usr/local/lib/R/site-library/RcppEigen/include/Eigen/Core:96:10: fatal error:
complex: No such file or directory
##
      96 | #include <complex>
##
## compilation terminated.
## make: *** [/usr/lib/R/etc/Makeconf:191: foo.o] Error 1
```

rstan::traceplot(dsamples, pars=c("transf\_mu\_alpha","transf\_mu\_threshold","transf\_
mu\_ndt", "transf\_mu\_theta",'transf\_mu\_beta', 'sd\_threshold',"sd\_alpha","sd\_ndt", '
sd\_theta', 'sd\_beta', "lp\_\_"), inc\_warmup = TRUE, nrow = 3)



pairs(dsamples, pars = c( "transf\_mu\_alpha","transf\_mu\_threshold","transf\_mu\_ndt",
 "transf\_mu\_theta",'transf\_mu\_beta', 'sd\_threshold',"sd\_alpha","sd\_ndt", 'sd\_thet
 a', 'sd\_beta', "lp\_\_"))

## Warning in par(usr): argument 1 does not name a graphical parameter
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print(dsamples, pars = c("transf\_mu\_alpha","transf\_mu\_threshold","transf\_mu\_ndt",
"transf\_mu\_theta",'transf\_mu\_beta', 'sd\_threshold',"sd\_alpha","sd\_ndt", 'sd\_thet
a', 'sd\_beta', "lp\_\_"))

```
## Inference for Stan model: EU discount.
## 4 chains, each with iter=3000; warmup=1500; thin=1;
## post-warmup draws per chain=1500, total post-warmup draws=6000.
##
##
                                                    2.5%
                                                               25%
                                                                          50%
                            mean se mean
                                            sd
## transf mu alpha
                            0.42
                                    0.00
                                          0.03
                                                    0.36
                                                              0.40
                                                                         0.42
## transf_mu_threshold
                            4.53
                                    0.01
                                          0.15
                                                    4.25
                                                              4.44
                                                                         4.53
## transf mu ndt
                            1.09
                                    0.00 0.05
                                                    0.99
                                                              1.05
                                                                        1.09
## transf mu theta
                            0.29
                                    0.00 0.05
                                                    0.21
                                                              0.25
                                                                        0.28
## transf_mu_beta
                                    0.00 0.12
                                                  -0.53
                                                             -0.33
                                                                       -0.24
                           -0.25
## sd threshold
                                    0.00 0.11
                                                    0.96
                                                              1.08
                           1.16
                                                                        1.15
                                    0.00 0.03
                                                    0.00
## sd alpha
                            0.05
                                                              0.03
                                                                        0.05
## sd ndt
                            0.57
                                    0.00 0.06
                                                    0.46
                                                              0.53
                                                                         0.57
## sd theta
                            0.27
                                    0.00 0.05
                                                    0.17
                                                              0.24
                                                                         0.27
## sd beta
                            0.77
                                    0.00 0.15
                                                    0.53
                                                              0.67
                                                                         0.76
## lp__
                      -23428.87
                                    0.52 17.05 -23463.48 -23439.99 -23428.57
##
                             75%
                                    97.5% n eff Rhat
                                      0.47 2659 1.00
## transf mu alpha
                            0.44
                                             442 1.01
## transf mu threshold
                            4.63
                                      4.82
## transf mu ndt
                           1.12
                                      1.18
                                           398 1.01
## transf mu theta
                            0.32
                                      0.39 2654 1.00
## transf mu beta
                                     -0.04
                                           763 1.00
                          -0.17
## sd_threshold
                                             795 1.00
                           1.23
                                      1.42
## sd alpha
                            0.07
                                      0.11 523 1.01
                                      0.70 1141 1.00
## sd_ndt
                            0.61
                                      0.36 1100 1.00
## sd theta
                            0.30
## sd beta
                            0.86
                                      1.10 1916 1.00
## lp
                       -23417.34 -23396.06 1086 1.00
##
## Samples were drawn using NUTS(diag e) at Wed Oct 25 02:09:12 2023.
## For each parameter, n_eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor on split chains (at
## convergence, Rhat=1).
```

## library(bayesplot)

```
## This is bayesplot version 1.10.0
```

```
## - Online documentation and vignettes at mc-stan.org/bayesplot
```

```
## - bayesplot theme set to bayesplot::theme_default()
```

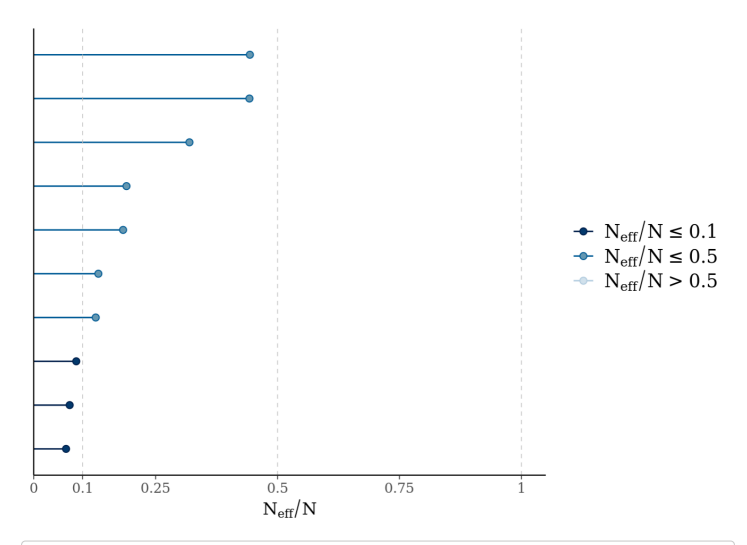
```
## * Does _not_ affect other ggplot2 plots
```

```
## * See ?bayesplot_theme_set for details on theme setting
```

```
ratios_cp <- neff_ratio(dsamples, pars = c("transf_mu_alpha","transf_mu_threshol
d","transf_mu_ndt", "transf_mu_theta",'transf_mu_beta', 'sd_threshold',"sd_alph
a","sd_ndt", 'sd_theta', 'sd_beta'))
df_ratios_cp <- as.data.frame(ratios_cp)
print(df_ratios_cp)</pre>
```

```
##
                        ratios_cp
## transf_mu_alpha
                       0.44324826
## transf mu threshold 0.07369538
## transf mu ndt
                       0.06632715
## transf mu theta
                       0.44234585
## transf_mu_beta
                       0.12712217
## sd_threshold
                       0.13256569
## sd alpha
                       0.08721803
## sd ndt
                       0.19024066
## sd_theta
                       0.18325915
## sd beta
                       0.31933757
```

```
mcmc_neff(ratios_cp, size = 2)
```



```
library(ggplot2)
library(tidyverse) # for the gather function
```

```
## — Attaching core tidyverse packages —
                                                          ----- tidyverse 2.0.0 -
## ✓ dplyr
             1.1.1
                         ✓ readr
                                     2.1.4
## ✓ forcats
               1.0.0

✓ stringr

                                     1.5.0
## ✓ lubridate 1.9.2

✓ tibble

                                     3.2.1
## ✓ purrr
               1.0.1

✓ tidyr

                                     1.3.0
## — Conflicts —
                                                          - tidyverse_conflicts() -
## * tidyr::extract() masks rstan::extract()
## * dplyr::filter() masks stats::filter()
## * dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all conf
licts to become errors
```

```
## Warning: Using an external vector in selections was deprecated in tidyselect 1.
1.0.
## i Please use `all_of()` or `any_of()` instead.
      # Was:
##
##
     data %>% select(parameters)
##
##
      # Now:
##
     data %>% select(all_of(parameters))
##
## See <a href="https://tidyselect.r-lib.org/reference/faq-external-vector.html">https://tidyselect.r-lib.org/reference/faq-external-vector.html</a>.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
```

```
# Convert hpd interval to a data frame and name the columns
hpd_interval_sub <- hpd_interval[parameters, ]</pre>
hpd df <- as.data.frame(hpd interval sub)</pre>
colnames(hpd_df) <- c("lower", "upper")</pre>
rownames(hpd df) <- parameters</pre>
hpd_df$parameter <- rownames(hpd_df)</pre>
# Aesthetic enhancements
theme_set(theme_minimal(base_size = 14)) # Set the default theme
custom_palette <- c("density_fill" = "lightgray",</pre>
                     "mean_line" = "blue",
                     "hpd line" = "darkgreen")
# Add text labels for mean, lower, and upper HPD values
df_long <- df_long %>%
  group_by(parameter) %>%
 mutate(mean = means[parameter])
hpd df <- hpd df %>%
  mutate(mid = (lower + upper) / 2)
p \leftarrow ggplot(df long, aes(x = value)) +
  geom_density(aes(fill = "density_fill")) +
  scale fill manual(values = custom palette, guide = FALSE) +
  geom_vline(aes(xintercept = mean, color = "mean_line"), linetype = "dashed", siz
e = 1, alpha = 0.7) +
  geom_text(data = df_long, aes(x = mean, y = 0, label = round(mean, 2)), vjust =
-0.5, hjust = 0.5, size = 4, color = custom palette["mean line"]) +
  geom_vline(data = hpd_df, aes(xintercept = lower, color = "hpd_line"), linetype
= "solid", size = 1, alpha = 0.5) +
  geom_text(data = hpd_df, aes(x = lower, y = 0, label = round(lower, 2)), vjust =
-0.5, hjust = -0.5, size = 4, color = custom_palette["hpd_line"]) +
  geom_vline(data = hpd_df, aes(xintercept = upper, color = "hpd_line"), linetype
= "solid", size = 1, alpha = 0.5) +
  geom_text(data = hpd_df, aes(x = upper, y = 0, label = round(upper, 2)), vjust =
-0.5, hjust = 1.5, size = 4, color = custom_palette["hpd_line"]) +
  facet_wrap(~ parameter, scales = "free", ncol = 2) +
  scale_color_manual(values = custom_palette, guide = 'none') +
  labs(title = "Posterior distributions")
```

```
## Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use `linewidth` instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
```

print(p)

```
## Warning: The `guide` argument in `scale_*()` cannot be `FALSE`. This was deprec
ated in
## ggplot2 3.3.4.
## i Please use "none" instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
```

## Posterior distributions

