

Covariate Selection

Supplementary material

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Data pre-processing

Packages & Options

```
library(tidyverse)
library(brms)
library(projpred)
library(bayesplot)

# retrieve # of cores
ncores <- parallel::detectCores()

# for output clarity
options(scipen = 999)
```

Data

```
# load data
apes1 <- read_csv("../data/laac_data_trial.csv")
apes2 <- read_csv("../data/laac_data_task.csv")

fn0 <- function(x, ...) {
  # helper function
  # sum over correct choice variable (code)
  to_return = tibble(cogn = sum(x$code))
  return(to_return)
}

code_sum <- apes1 %>%
  # contains summed code variable [for each task, time point, session and subject]
  group_by(time_point, session, subject, task) %>%
  group_modify(fn0)
```

```

apes1_tmp <- apes1 %>%
  # helper for merging
  select(-c(date, trial_session, trial_time_point, code)) %>%
  unique(by = c("time_point", "session", "subject"))

apes1_new <-
  as_tibble(merge(apes1_tmp, code_sum, by = c("time_point", "session", "subject", "task"))) %>%
  mutate(across(c(subject, group, heat, test_day, le_present, dist_present, sex, rearing, observer), as.factor)) %>%
  mutate(observer = fct_relevel(observer, "no")) %>%
  jtools::center(., vars = c("sick_severity",
                             "le_mean",
                             "time_outdoors",
                             "age",
                             "time_in_leipzig")) %>%
  group_by(group, time_point) %>%
  mutate(rank_gmc = rank - mean(rank, na.rm = TRUE)) %>%
  ungroup() %>%
  arrange(time_point)

grp_size <- tibble(
  # number of apes for each species
  a_chimp = 20,
  b_chimp = 6,
  bonobo = 12,
  gorilla = 6,
  orangutan = 6
)

apes1_new <- apes1_new %>%
  # create rank variable depending on species
  group_by(group, time_point) %>%
  mutate(
    rel_rank = case_when(
      group == "a_chimp" ~ percent_rank(grp_size$a_chimp:1)[rank],
      group == "b_chimp" ~ percent_rank(grp_size$b_chimp:1)[rank],
      group == "bonobo" ~ percent_rank(grp_size$bonobo:1)[rank],
      group == "gorilla" ~ percent_rank(grp_size$gorilla:1)[rank],
      group == "orangutan" ~ percent_rank(grp_size$orangutan:1)[rank]
    )
  ) %>%
  ungroup()

apes1_new <- apes1_new %>%
  # create coding for heat variable
  mutate(heat_mod = case_when(
    sex == "f" & heat == "yes" ~ "_f_fheat",
    sex == "m" & heat == "yes" ~ "_m_fheat",
    sex == "f" & heat == "no" ~ "_f_noheat",
    sex == "m" & heat == "no" ~ "_m_noheat"),
    heat_mod = as_factor(heat_mod)
  ) %>%
  mutate(heat_mod = fct_relevel(heat_mod, "_f_noheat"))

```

```

apes1_new <- apes1_new %>%
  select(-heat, -heat_mod)

apes1_new <- apes1_new %>%
  # recode rearing categories: hand -> unknown
  mutate(rearing = fct_recode(rearing, "hand" = "unknown"))

t_cau <- filter(apes1_new, task == "causality")
t_inf <- filter(apes1_new, task == "inference")
t_quant <- filter(apes1_new, task == "quantity")
t_gaze <- filter(apes1_new, task == "gaze_following")

t_gaze <- t_gaze %>%
  # create dummy variable indicating if session 1 or 2
  group_by(time_point, session) %>%
  mutate(tp_mod = cur_group_id()) %>%
  ungroup() %>%
  mutate(day2 = case_when(session == 1 ~ "no",
                          session == 2 ~ "yes"),
         day2 = factor(day2)) %>%
  select(tp_mod, day2, everything())

t_gaze <- t_gaze %>%
  # remove duplicates created by day2
  group_by(subject) %>%
  filter(!duplicated(tp_mod)) %>%
  ungroup()

```

Covariate selection

```

# covariate needed for projection prediction
# placed here for easy editing of formula
all_fixed_effects <- c("sick_severity",
                      "test_day", "test_tp",
                      "rel_rank",
                      "observer",
                      "age", "time_in_leipzig",
                      "sex", "group",
                      "rearing",
                      "le_mean",
                      "dist_mean",
                      "time_outdoors",
                      "sociality")

fm <- formula(cogn ~ sick_severity +
              test_day + test_tp +
              rel_rank + # rank_gmc +
              observer +
              age + time_in_leipzig +
              sex + group +
              rearing +
              le_mean + # le_max + # le_present +

```

```

      dist_mean + # dist_max + # + dist_present +
      time_outdoors +
      sociality + # sociality_total
      # heat_mod + # heat +
      (1|subject)
    )

fm_gaze <- update(fm, . ~ . +day2)

```

Reference Model: 2-level Multilevel Model (random intercepts only)

```

m_cau_2l <- brm(fm, data = t_cau,
  warmup = 1e3, iter = 4e3, cores = ncores, chains = 4,
  seed = 2021,
  save_pars = save_pars(all = TRUE)
)

```

Warning: Rows containing NAs were excluded from the model.

```

m_inf_2l <- brm(fm, data = t_inf,
  warmup = 1e3, iter = 4e3, cores = ncores, chains = 4,
  seed = 2021,
  save_pars = save_pars(all = TRUE)
)

```

Warning: Rows containing NAs were excluded from the model.

```

m_quant_2l <- brm(fm, data = t_quant,
  warmup = 1e3, iter = 4e3, cores = ncores, chains = 4,
  seed = 2021,
  save_pars = save_pars(all = TRUE)
)

```

Warning: Rows containing NAs were excluded from the model.

```

m_gaze_2l <- brm(fm_gaze, data = t_gaze,
  warmup = 1e3, iter = 4e3, cores = ncores, chains = 4,
  seed = 2021,
  save_pars = save_pars(all = TRUE)
)

```

Warning: Rows containing NAs were excluded from the model.

```
summary(m_cau_2l)
```

```

## Family: gaussian
## Links: mu = identity; sigma = identity
## Formula: cogn ~ sick_severity + test_day + test_tp + rel_rank + observer + age + time_in_leipzig + s
## Data: t_cau (Number of observations: 450)
## Samples: 4 chains, each with iter = 4000; warmup = 1000; thin = 1;
##          total post-warmup samples = 12000
##
## Group-Level Effects:
## ~subject (Number of levels: 41)
##           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)    2.19     0.32    1.65    2.89 1.00    4533    6817
##

```

```
## Population-Level Effects:
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## Intercept      7.10      1.66   3.82   10.41 1.00    7279    8270
## sick_severity   0.19      0.19  -0.19    0.56 1.00   16260    9087
## test_dayyes     0.71      1.09  -1.42    2.88 1.00   16695    9366
## test_tp         0.00      0.04  -0.08    0.08 1.00   18745    9463
## rel_rank        0.28      1.35  -2.40    2.98 1.00   10121    8424
## observeryes     0.30      0.34  -0.35    0.96 1.00   17567   10104
## age            -0.02      0.05  -0.12    0.08 1.00    7709    7819
## time_in_leipzig 0.11      0.08  -0.05    0.27 1.00    7671    8415
## sexf           0.17      0.86  -1.55    1.84 1.00    7627    8547
## groupb_chimp    3.09      1.61  -0.12    6.30 1.00    6635    7782
## groupa_chimp   -0.82      1.20  -3.18    1.53 1.00    5739    7020
## grouporangutan  1.98      1.77  -1.62    5.45 1.00    6917    7074
## groupbonobo     1.10      1.31  -1.42    3.69 1.00    6247    8045
## rearinghand     0.01      1.22  -2.41    2.36 1.00    7969    8288
## le_mean        -0.67      0.54  -1.73    0.39 1.00   18562    9531
## dist_mean      -0.36      0.21  -0.78    0.06 1.00   15668    9770
## time_outdoors  -0.07      0.06  -0.19    0.04 1.00   15460    9047
## sociality       0.07      0.24  -0.39    0.54 1.00   16087    8636
```

```
## Family Specific Parameters:
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sigma      2.22      0.08   2.07   2.38 1.00   14168    8736
```

```
## Samples were drawn using sampling(NUTS). 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).
```

```
summary(m_inf_21)
```

```
## Family: gaussian
## Links: mu = identity; sigma = identity
## Formula: cogn ~ sick_severity + test_day + test_tp + rel_rank + observer + age + time_in_leipzig + s
## Data: t_inf (Number of observations: 451)
## Samples: 4 chains, each with iter = 4000; warmup = 1000; thin = 1;
##          total post-warmup samples = 12000
##
## Group-Level Effects:
## ~subject (Number of levels: 41)
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)  2.80      0.38   2.17   3.65 1.00    4419    7020
##
## Population-Level Effects:
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## Intercept      6.31      1.89   2.63   10.07 1.00    7436    8217
## sick_severity  -0.09      0.17  -0.43    0.25 1.00   18592    9161
## test_dayyes     0.80      0.95  -1.07    2.64 1.00   21465    9046
## test_tp        -0.09      0.03  -0.16  -0.02 1.00   21423    9470
## rel_rank       -0.23      1.32  -2.82    2.36 1.00   14037    9317
## observeryes     0.51      0.29  -0.06    1.06 1.00   22336    8669
## age            -0.08      0.06  -0.19    0.05 1.00    7460    7962
## time_in_leipzig 0.37      0.10   0.17    0.56 1.00    7804    7460
## sexf           0.18      1.07  -1.98    2.27 1.00    7449    7339
## groupb_chimp    0.57      2.02  -3.43    4.58 1.00    6699    7571
```

```
## groupa_chimp      -0.01      1.50     -2.97      3.02 1.00      5908      6591
## grouporangutan    -2.17      2.12     -6.37      2.01 1.00      6876      7605
## groupbonobo       -1.58      1.67     -4.92      1.77 1.00      6433      7179
## rearinghand       -0.49      1.48     -3.43      2.43 1.00      7625      7590
## le_mean           -0.57      0.45     -1.46      0.33 1.00     22706     8269
## dist_mean         -0.11      0.18     -0.46      0.24 1.00     18597     8971
## time_outdoors     -0.01      0.05     -0.11      0.08 1.00     17195     8955
## sociality          0.03      0.20     -0.37      0.42 1.00     20660     8781
```

```
##
```

```
## Family Specific Parameters:
```

```
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
```

```
## sigma      1.86      0.07      1.74      2.00 1.00     16348     8687
```

```
##
```

```
## Samples were drawn using sampling(NUTS). 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).
```

```
summary(m_quant_2l)
```

```
## Family: gaussian
```

```
## Links: mu = identity; sigma = identity
```

```
## Formula: cogn ~ sick_severity + test_day + test_tp + rel_rank + observer + age + time_in_leipzig + s
```

```
## Data: t_quant (Number of observations: 421)
```

```
## Samples: 4 chains, each with iter = 4000; warmup = 1000; thin = 1;
```

```
## total post-warmup samples = 12000
```

```
##
```

```
## Group-Level Effects:
```

```
## ~subject (Number of levels: 41)
```

```
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
```

```
## sd(Intercept)  1.75      0.26      1.31      2.35 1.00     4406     7213
```

```
##
```

```
## Population-Level Effects:
```

```
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
```

```
## Intercept      8.74      1.34      6.13     11.38 1.00     9562     8749
```

```
## sick_severity  -0.26      0.16     -0.58      0.05 1.00    15739     9718
```

```
## test_dayyes     0.42      0.82     -1.18      2.03 1.00    16761     9939
```

```
## test_tp         0.02      0.03     -0.05      0.08 1.00    18450     9405
```

```
## rel_rank        0.33      1.08     -1.83      2.38 1.00    13256     8858
```

```
## observeryes    -0.37      0.31     -0.98      0.23 1.00    15966     9596
```

```
## age            -0.03      0.04     -0.11      0.04 1.00     9270     8017
```

```
## time_in_leipzig 0.23      0.07      0.11      0.36 1.00     9527     8539
```

```
## sexf           0.78      0.70     -0.59      2.14 1.00    10367     8996
```

```
## groupb_chimp    1.59      1.31     -1.01      4.19 1.00     8814     8486
```

```
## groupa_chimp   -1.52      0.99     -3.49      0.44 1.00     8174     8244
```

```
## grouporangutan -2.07      1.42     -4.88      0.74 1.00     8971     9208
```

```
## groupbonobo     0.11      1.07     -2.02      2.23 1.00     8239     7539
```

```
## rearinghand    -2.31      0.96     -4.25     -0.44 1.00     9596     8491
```

```
## le_mean         0.14      0.44     -0.72      1.01 1.00    18361     9509
```

```
## dist_mean      -0.04      0.19     -0.41      0.32 1.00    16275     9959
```

```
## time_outdoors  -0.01      0.05     -0.12      0.09 1.00    15895     9574
```

```
## sociality      -0.03      0.21     -0.44      0.39 1.00    18675     8957
```

```
##
```

```
## Family Specific Parameters:
```

```
##      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
```

```
## sigma      1.77      0.07      1.64      1.90 1.00    14069     8719
```

```
##
## Samples were drawn using sampling(NUTS). 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).

summary(m_gaze_2l)

## Family: gaussian
## Links: mu = identity; sigma = identity
## Formula: cogn ~ sick_severity + test_day + test_tp + rel_rank + observer + age + time_in_leipzig + s
## Data: t_gaze (Number of observations: 879)
## Samples: 4 chains, each with iter = 4000; warmup = 1000; thin = 1;
##          total post-warmup samples = 12000
##
## Group-Level Effects:
## ~subject (Number of levels: 41)
##           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)      0.75      0.10      0.58      0.98 1.00      4218      6995
##
## Population-Level Effects:
##           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## Intercept          0.56      0.50     -0.41      1.55 1.00      6860      8365
## sick_severity      -0.13      0.05     -0.23     -0.03 1.00     15844      8407
## test_dayyes         0.22      0.23     -0.22      0.66 1.00     18709      8805
## test_tp             0.01      0.01     -0.01      0.02 1.00     22027      8992
## rel_rank            -0.14      0.35     -0.82      0.54 1.00     12451      9498
## observeryes        -0.12      0.08     -0.27      0.04 1.00     18950      9392
## age                 0.02      0.02     -0.01      0.05 1.00      6351      7397
## time_in_leipzig    -0.00      0.03     -0.06      0.05 1.00      7323      7899
## sexf                0.23      0.29     -0.34      0.79 1.00      6948      7378
## groupb_chimp        0.55      0.53     -0.48      1.62 1.00      6547      7347
## groupa_chimp        0.78      0.40     -0.00      1.58 1.00      5559      6881
## grouporangutan      0.08      0.57     -1.04      1.21 1.00      6504      7480
## groupbonobo         0.59      0.44     -0.26      1.45 1.00      6055      6594
## rearinghand        -0.85      0.40     -1.63     -0.05 1.00      6825      7601
## le_mean             -0.03      0.12     -0.27      0.21 1.00     21443      8520
## dist_mean           -0.10      0.05     -0.19     -0.00 1.00     16821      9112
## time_outdoors       0.10      0.01      0.07      0.12 1.00     15600      9673
## sociality           -0.14      0.06     -0.25     -0.03 1.00     20094      8699
## day2yes             -0.12      0.05     -0.21     -0.02 1.00     20069      7768
##
## Family Specific Parameters:
##           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sigma        0.71      0.02      0.68      0.75 1.00     17879      8363
##
## Samples were drawn using sampling(NUTS). 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).
```

Predictive Projection

```
# delay random intercept to last place so that it doesn't soak up all the variance
s_terms <- c("1", all_fixed_effects,
             paste0(paste(all_fixed_effects, collapse = " + "), " + (1 | subject)"))
```

```

s_terms_gaze <- c("1", c(all_fixed_effects, "day2"),
  paste0(paste(c(all_fixed_effects, "day2"), collapse = " + "), " + (1 | subject)"))

refM_cau <- get_refmodel(m_cau_2l)
refM_inf <- get_refmodel(m_inf_2l)
refM_quant <- get_refmodel(m_quant_2l)
refM_gaze <- get_refmodel(m_gaze_2l)

vs_cau <- varsel(refM_cau, search_terms = s_terms)
summary(vs_cau); plot(vs_cau, stats = c('elpd', 'rmse'))

randint_ind_vscau <- length(solution_terms(vs_cau))
relevant_cov_vscau <- c(1, 2, 3, 4, 5, randint_ind_vscau)

# proj_cau <- project(vs_cau, solution_terms = relevant_cov_vscau)
# mcmc_areas(as.matrix(proj_cau), pars = solution_terms(vs_cau)[relevant_cov_vscau])

vs_inf <- varsel(refM_inf, search_terms = s_terms)
summary(vs_inf); plot(vs_inf, stats = c('elpd', 'rmse'))

randint_ind_vsinf <- length(solution_terms(vs_inf))
relevant_cov_vsinf <- c(1, 2, 3, 4, 5, randint_ind_vsinf)

# proj_inf <- project(vs_inf, solution_terms = relevant_cov_vsinf)
# mcmc_areas(as.matrix(proj_inf), pars = solution_terms(vs_inf)[relevant_cov_vsinf])

vs_quant <- varsel(refM_quant, search_terms = s_terms)
summary(vs_quant); plot(vs_quant, stats = c('elpd', 'rmse'))

randint_ind_vsquant <- length(solution_terms(vs_quant))
relevant_cov_vsquant <- c(1, 2, 3, 4, 5, randint_ind_vsquant)

# proj_quant <- project(vs_quant, solution_terms = relevant_cov_vsquant)
# mcmc_areas(as.matrix(proj_quant), pars = solution_terms(vs_quant)[relevant_cov_vsquant])

vs_gaze <- varsel(refM_gaze, search_terms = s_terms_gaze)
summary(vs_gaze); plot(vs_gaze, stats = c('elpd', 'rmse'))

randint_ind_vsgaze <- length(solution_terms(vs_gaze))
relevant_cov_vsgaze <- c(1, 2, 3, 4, 5, randint_ind_vsgaze)

# proj_gaze <- project(vs_gaze, solution_terms = relevant_cov_vsgaze)
# mcmc_areas(as.matrix(proj_gaze), pars = solution_terms(vs_gaze)[relevant_cov_vsgaze])

cvs_cau <- cv_varsel(refM_cau,
  search_terms = s_terms, cv_method = "LOO", method = "forward",
  seed = 2020)

## [1] "Computing LOOs..."
## |

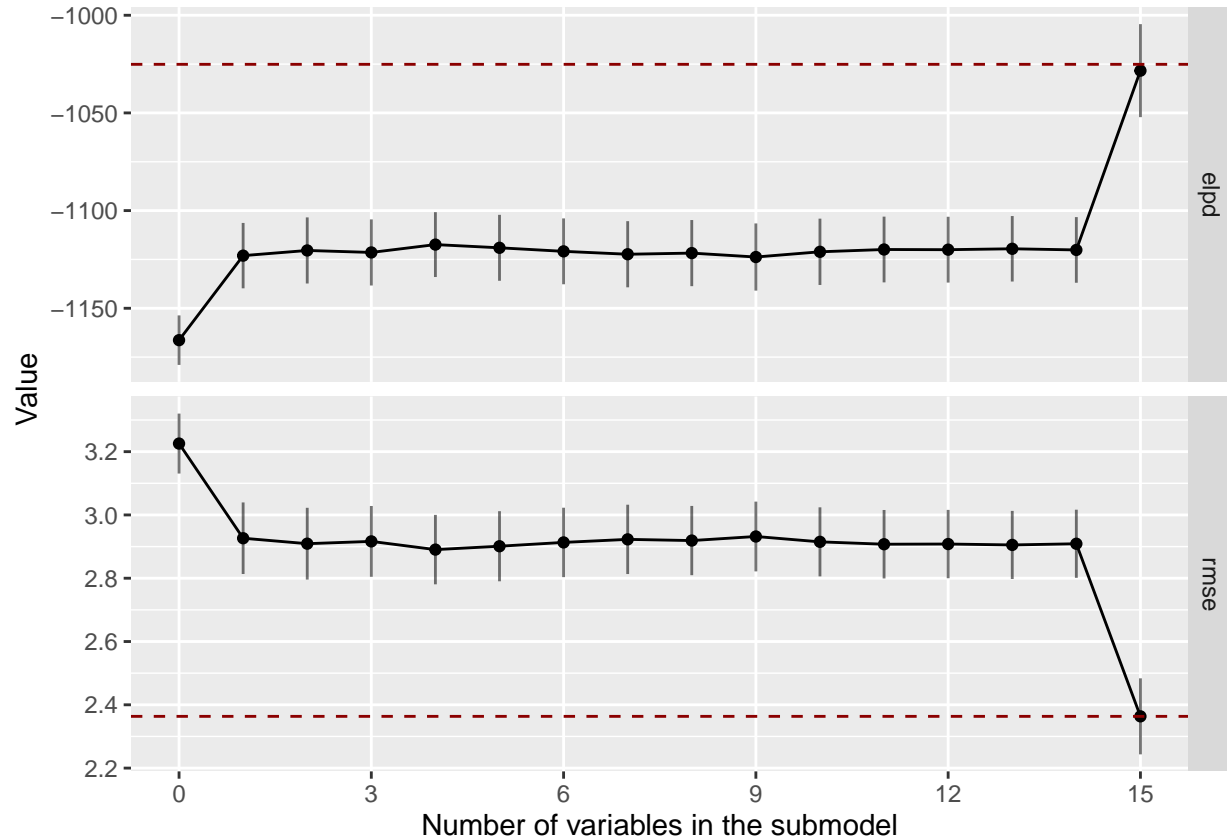
```



```
## [1] "50% of terms selected."
## [1] "60% of terms selected."
## [1] "70% of terms selected."
## [1] "80% of terms selected."
## [1] "90% of terms selected."
## [1] "100% of terms selected."
## [1] "Done."
```

```
summary(cvs_cau); plot(cvs_cau, stats = c('elpd', 'rmse'))
```

##	size	solution_terms	elpd	elpd.se
## 2	0	<NA>	-1166.320	12.74171
## 3	1	group	-1123.058	16.82876
## 4	2	time_in_leipzig	-1120.403	17.01587
## 5	3	sociality	-1121.423	17.01735
## 6	4	dist_mean	-1117.403	16.68867
## 7	5	sick_severity	-1119.044	16.97399
## 8	6	age	-1120.857	16.95057
## 9	7	le_mean	-1122.342	17.02853
## 10	8	test_day	-1121.755	17.02313
## 11	9	rel_rank	-1123.767	17.29413
## 12	10	observer	-1121.095	17.07342
## 13	11	time_outdoors	-1119.914	16.92239
## 14	12	sex	-1120.004	16.94353
## 15	13	rearing	-1119.549	16.85887
## 16	14	test_tp	-1120.135	16.89832
## 17	15	(1 subject)	-1028.359	23.94456



```
# proj_cau_cv <- project(cvs_cau, solution_terms = c(1, 2, 3, 14))
# mcmc_areas(as.matrix(proj_cau_cv), pars = solution_terms(cvs_cau)[c(1, 2, 3, 14)])
```

relevant covariates: (1 | subject), group

```
cvs_inf <- cv_varsel(refM_inf,
  search_terms = s_terms, cv_method = "L00", method = "forward",
  seed = 2020)
```

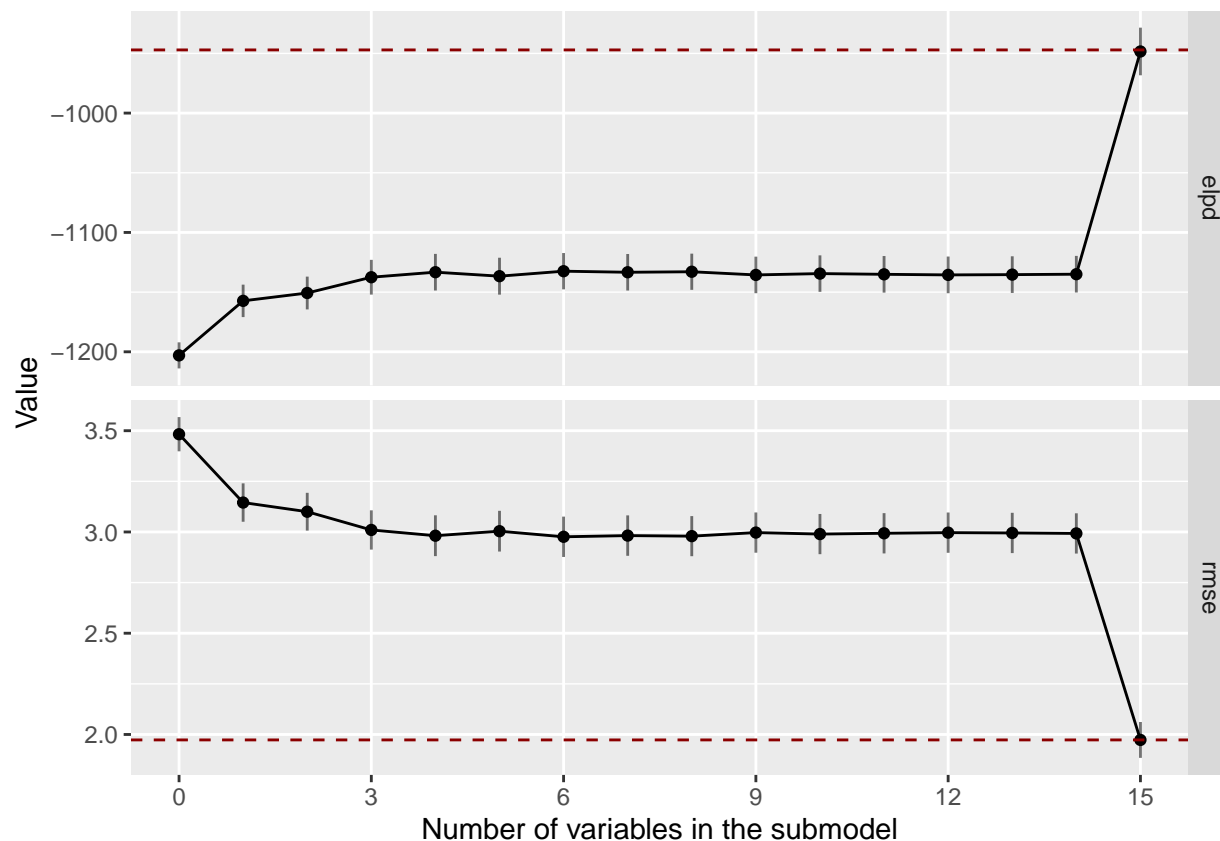
```
## [1] "Computing L00s..."
```

```
## |
```

```
|
```

```
summary(cvs_inf); plot(cvs_inf, stats = c('elpd', 'rmse'))
```

```
##   size  solution_terms      elpd  elpd.se
## 2     0              <NA> -1202.9857 10.97886
## 3     1 time_in_leipzig -1157.3144 13.67677
## 4     2              group -1150.7333 13.81341
## 5     3              age -1137.5141 14.65069
## 6     4 sick_severity -1133.2952 15.42105
## 7     5      test_day -1136.6343 15.54014
## 8     6 time_outdoors -1132.4628 15.29881
## 9     7      observer -1133.3510 15.38904
## 10    8      rearing -1132.9248 15.24541
## 11    9    sociality -1135.5923 15.40453
## 12   10     rel_rank -1134.4788 15.36408
## 13   11    dist_mean -1135.0805 15.42556
## 14   12     test_tp -1135.5634 15.46405
## 15   13     le_mean -1135.3409 15.44985
## 16   14       sex -1134.9778 15.42959
## 17   15 (1 | subject) -948.3915 20.07552
```



```
# proj_inf_cv <- project(cvs_inf, solution_terms = c(1, 2, 14))
# mcmc_areas(as.matrix(proj_inf_cv), pars = solution_terms(cvs_inf)[c(1, 2, 14)])
```

relevant covariates: (1 | subject), time_in_leipzig, group, age

```
cvs_quant <- cv_varselect(refM_quant,
  search_terms = s_terms, cv_method = "L00", method = "forward",
  seed = 2020)
```

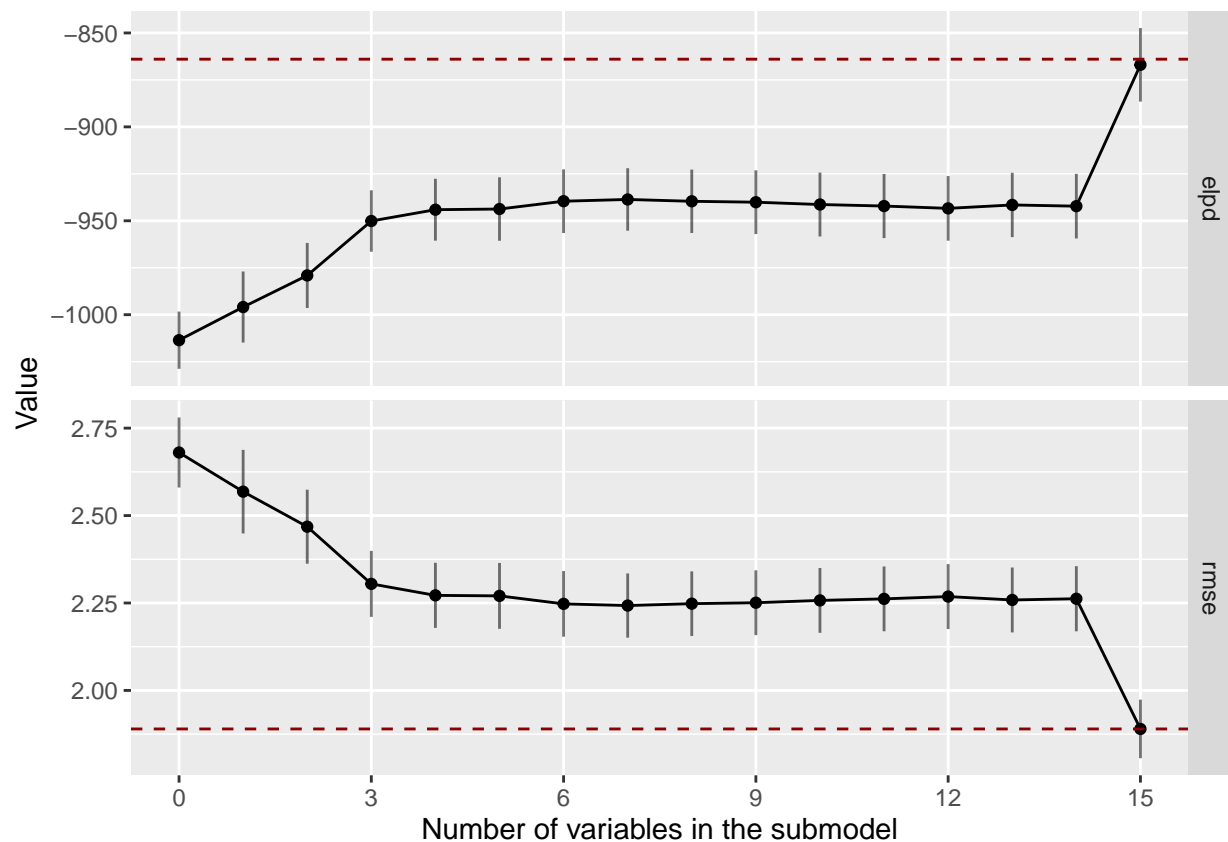
```
## [1] "Computing LOOs..."
```

```
## |
```

```
summary(cvs_quant); plot(cvs_quant, stats = c('elpd', 'rmse'))
```

```
## size solution_terms elpd elpd.se
## 2 0 <NA> -1013.5630 15.32066
```

```
## 3      1 time_in_leipzig -995.9227 19.01982
## 4      2      rearing   -979.0997 17.41291
## 5      3        group   -950.1172 16.43463
## 6      4      observer  -944.0947 16.56630
## 7      5         sex    -943.7346 16.98652
## 8      6      rel_rank  -939.5734 17.04140
## 9      7         age    -938.6473 16.73218
## 10     8 sick_severity -939.6164 16.96110
## 11     9      test_tp   -940.1058 17.06342
## 12    10     test_day   -941.3380 17.09880
## 13    11 time_outdoors -942.1494 17.15309
## 14    12      dist_mean -943.3841 17.29724
## 15    13        le_mean -941.5656 17.21523
## 16    14     sociality  -942.2124 17.29028
## 17    15 (1 | subject) -866.9580 19.68460
```



```
# proj_quant_cv <- project(cvs_quant, solution_terms = c(1, 2, 3, 15))
# mcmc_areas(as.matrix(proj_quant_cv), pars = solution_terms(cvs_quant)[c(1, 2, 3, 15)])
```

relevant covariates: (1 | subject), time_in_leipzig, rearing, group

```
cvs_gaze <- cv_varsel(refM_gaze,
  search_terms = s_terms_gaze, cv_method = "LOO", method = "forward",
  seed = 2020)
```

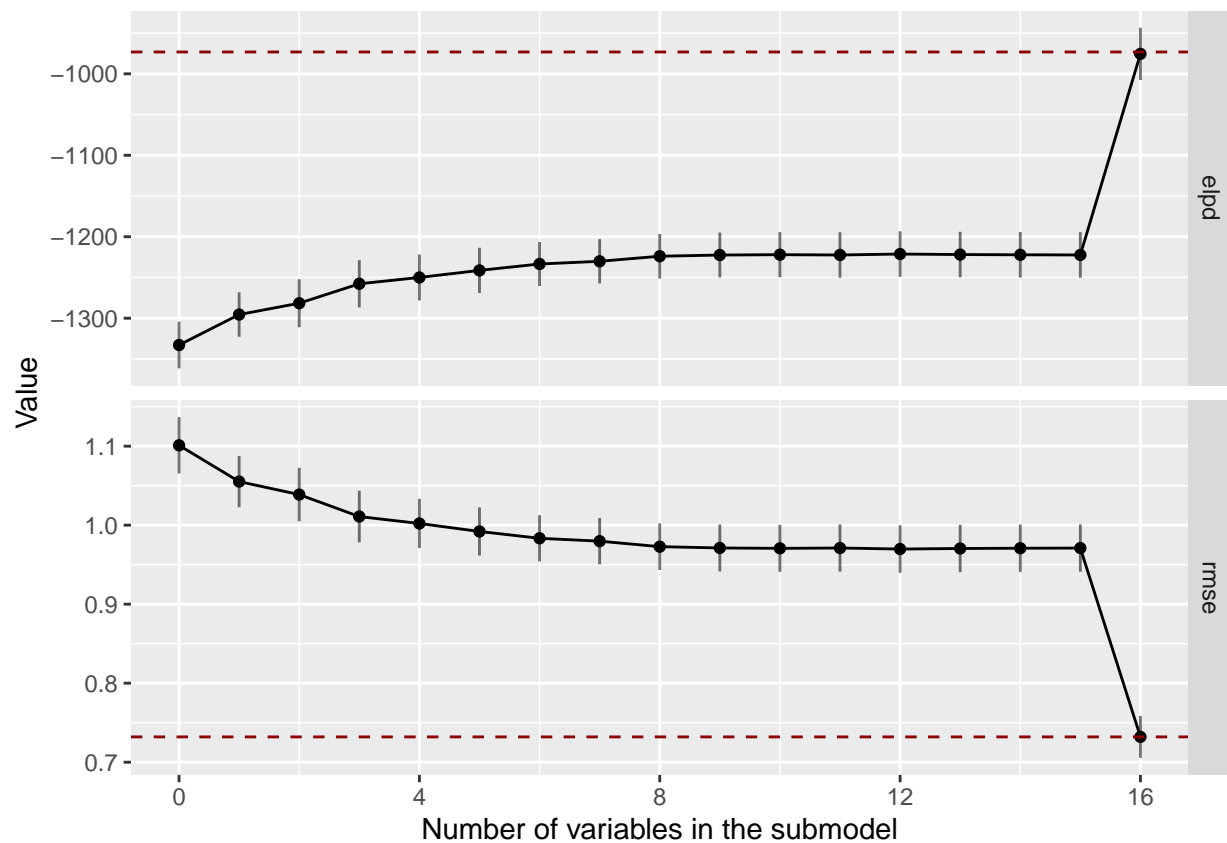
```
## [1] "Computing LOOs..."
## |
```

|

```
## [1] "10% of terms selected."
## [1] "20% of terms selected."
## [1] "30% of terms selected."
## [1] "40% of terms selected."
## [1] "50% of terms selected."
## [1] "60% of terms selected."
## [1] "70% of terms selected."
## [1] "80% of terms selected."
## [1] "90% of terms selected."
## [1] "100% of terms selected."
## [1] "Done."
```

```
summary(cvs_gaze); plot(cvs_gaze, stats = c('elpd', 'rmse'))
```

```
##      size solution_terms      elpd elpd.se
## 2      0      <NA> -1332.926 28.85612
## 3      1      group -1295.514 27.45796
## 4      2      rearing -1281.612 29.61006
## 5      3 time_outdoors -1257.730 29.16308
## 6      4      age -1249.990 28.41721
## 7      5      sociality -1241.305 27.96193
## 8      6      sex -1233.446 27.09530
## 9      7 sick_severity -1230.140 27.31724
## 10     8      observer -1223.934 27.57003
## 11     9 time_in_leipzig -1222.454 27.68608
## 12    10      day2 -1222.011 27.80184
## 13    11      dist_mean -1222.380 28.01366
## 14    12      test_day -1221.232 28.10263
## 15    13      rel_rank -1221.860 28.08296
## 16    14      test_tp -1222.162 28.09163
## 17    15      le_mean -1222.340 28.09300
## 18    16 (1 | subject) -975.499 32.36394
```



```
# proj_gaze_cv <- project(cvs_gaze, solution_terms = c(1, 2, 3, 4, 5, 6, 7, 15))
# mcmc_areas(as.matrix(proj_gaze_cv), pars = solution_terms(cvs_gaze)[c(1, 2, 3, 4, 5, 6, 7, 15)])
```

relevant covariates: (1 | subject), group, rearing, time_outdoors, age, sociality, sex