Appendix 6

R. code

```
library(simstudy)
library(cmdstanr)
library(data.table)
library(slurmR)
library(posterior)
mco <- cmdstan_model("./primary_co.stan")</pre>
#--- generating the study-specific baseline probabilities for each outcome level---#
genBaseProbs <- function(n, base, similarity, digits = 2) {</pre>
    n levels <- length(base)</pre>
    x <- gtools::rdirichlet(n, similarity * base)</pre>
    #--- ensure that each vector of probabilities sums exactly to 1
    x <- round(floor(x * 1e+08)/1e+08, digits) # round the generated probabilities
    xpart <- x[, 1:(n_levels - 1)] # delete the base prob of the final level</pre>
    partsum <- apply(xpart, 1, sum) # add the values of levels 1 to K-1
    x[, n_levels] \leftarrow 1 - partsum # the base prob of the level <math>K = 1 - sum(1:[K-1])
    return(x)
}
nsites <- 9 # assume 9 RCTs in total
basestudy \leftarrow genBaseProbs(n = nsites, base = c(0.1, 0.107,
    0.095, 0.085, 0.09, 0.09, 0.108, 0.1, 0.09, 0.075, 0.06),
    similarity = 100)
# More detailed information in Keith's blog :
# https://www.rdatagen.net/post/generating-probabilities-for-ordinal-categorical-data/
#--- data definition code ---#
defC <- defDataAdd(varname = "b", formula = 0, variance = 0.01,</pre>
    dist = "normal") # RCT specific intercept
defC <- defDataAdd(defC, varname = "size", formula = "75+75*large",</pre>
    dist = "nonrandom") # sample size; large=1: 150 patients, large=0: 75 patients
defC2 <- defDataAdd(varname = "C_rv", formula = "C * control",</pre>
    dist = "nonrandom") # 0=CP, 1= standard of care, 2=non-CP, 3=saline
defC2 <- defDataAdd(defC2, varname = "sex", formula = 0.5,</pre>
    dist = "binary")
defC2 <- defDataAdd(defC2, varname = "who_enroll", formula = "1/3;1/3;1/3",</pre>
    dist = "categorical")
defC2 <- defDataAdd(defC2, varname = "age", formula = "0.25;0.25;0.50",</pre>
    dist = "categorical")
defC2 <- defDataAdd(defC2, varname = "ss", formula = "0.2;0.2;0.2;0.2",</pre>
    dist = "categorical")
defC2 \leftarrow defDataAdd(defC2, varname = "z", formula = "0.05*(ss-1) + 0.1*sex +
                     0.075*(age-1) + 0.06*(who enroll-1) +
                     (0.3 + b) * (C rv==1) +
```

```
(0.4 + b ) * (C_rv==2) + (0.5 + b ) * (C_rv==3)",
dist = "nonrandom")
```

```
iter <- function(iternum, defC, defC2, basestudy, nsites,</pre>
    set_cmdstan_path(path = "/gpfs/share/apps/cmdstan/2.25.0")
    #--- data generation ---#
    dstudy <- genData(nsites, id = "study") # 9 RCTs</pre>
    dstudy <- trtAssign(dstudy, nTrt = 3, grpName = "C") # allocate 3 control conditions
    dstudy <- trtAssign(dstudy, nTrt = 2, strata = "C",</pre>
        grpName = "large", ratio = c(2, 1))
    dstudy <- addColumns(defC, dstudy)</pre>
    dind <- genCluster(dstudy, "study", numIndsVar = "size",</pre>
    dind <- trtAssign(dind, strata = "study", grpName = "control")</pre>
    dind <- addColumns(defC2, dind)</pre>
    setkey(dind, "id")
    dl <- lapply(1:nsites, function(i) {</pre>
        b <- basestudy[i, ]</pre>
        dx <- dind[study == i]</pre>
        dx <- genOrdCat(dx, adjVar = "z", b, catVar = "ordY")</pre>
        dx[, := (ordY, factor(ordY, levels = c(1:11)))]
        dx[]
    })
    dind <- rbindlist(dl)</pre>
    #--- model estimation ---#
    N = nrow(dind) # number of observations
    L <- dind[, length(unique(ordY))] # number of levels of outcome
    K <- dind[, length(unique(study))] # number of RCTs</pre>
    y <- as.numeric(dind$ordY) # individual outcome
    kk <- dind$study # RCT for each individual
    ctrl <- dind$control # treatment arm for individual</pre>
    cc <- dind[, .N, keyby = .(study, C)]$C # specific control arm for RCT
    x <- model.matrix(ordY ~ factor(who enroll) + factor(age) +
        factor(sex) + factor(ss), data = dind)[, -1]
    D \leftarrow ncol(x)
    prior_div <- 8</pre>
    prior_Delta_sd <- 0.354</pre>
    eta <- 0.1
    prior eta 0 <- 0.25
    prior_beta_sd <- 2.5</pre>
    studydata \leftarrow list(N = N, L = L, K = K, y = y, ctrl = ctrl,
        cc = cc, kk = kk, prior_div = prior_div, prior_Delta_sd = prior_Delta_sd,
        eta = eta, prior_eta_0 = prior_eta_0, x = x, D = D,
        prior_beta_sd = prior_beta_sd)
    fit_co <- mco$sample(step_size = 0.1, data = studydata,</pre>
```

```
#--- Replication ---#
job <- Slurm_lapply(1:2520, iter, defC = defC, defC2 = defC2,
    basestudy = basestudy, nsites = nsites, mco = mco, njobs = 90,
    mc.cores = 4, tmp_path = "/gpfs/scratch/...", job_name = "sim_1",
    sbatch_opt = list(time = "24:00:00"), plan = "wait",
    overwrite = TRUE)

site_plasma_all <- Slurm_collect(job)
site_plasma <- rbindlist(site_plasma)

date_stamp <- gsub("-", "", Sys.Date())
dir.create(file.path("/gpfs/home/.../r/", date_stamp), showWarnings = FALSE)
save(site_plasma, file = paste0("/gpfs/home/.../r/", date_stamp,
    "/model_co.rda"))</pre>
```

Stan code

```
data {
 int<lower=0> N;
                                // number of observations
                                // number of WHO categories
  int<lower=2> L;
                                // number of RCTs
  int<lower=1> K;
  int<lower=1,upper=L> y[N];
                                // vector of categorical outcomes
  int<lower=1,upper=K> kk[N];
                                // RCT for individual
  int<lower=0,upper=1> ctrl[N]; // treatment or control
                                // specific control for RCT
  int<lower=1,upper=3> cc[K];
  int<lower=1> D;
                                // number of covariates
 row_vector[D] x[N];
                                // strata indicators N x D matrix
 real<lower=0> prior_div;
                                // prior sd of tau
  real<lower=0> prior_Delta_sd;
                                // prior sd of overall control effect
  real<lower=0> eta;
                                // prior sd of delta
  real<lower=0> prior_beta_sd;
                                // prior sd of beta
 real<lower=0> prior_eta_0;
                                 //prior sd of eta_0 (the sd of delta_k)
parameters {
 real alpha;
                                // overall intercept for treatment
 ordered[L-1] tau[K];
                                // cut-points for cumulative odds model (K X [L-1] matrix)
 real<lower=0> eta_0;
                                // sd of delta_k (around delta)
```

```
// non-central parameterization
  vector[K] z_ran_rx;
  vector[3] z_delta;
  vector[D] z_beta;
  real z_Delta;
transformed parameters{
  vector[3] delta;
                                // control-specific effect
                                 // RCT-specific treatment effect
  vector[K] delta_k;
  vector[D] beta;
                                // covariate estimates
                                // overall control effect
  real Delta;
  vector[N] yhat;
  Delta = prior_Delta_sd * z_Delta;
  delta = eta * z_delta + Delta;
  beta = prior_beta_sd * z_beta;
  for (k in 1:K)
    delta_k[k] = eta_0 * z_ran_rx[k] + delta[cc[k]];
    yhat[i] = alpha + ctrl[i] * delta_k[kk[i]] + x[i] * beta;
model {
  // priors
  alpha \sim normal(0,0.1);
  z_ran_rx ~ std_normal();
  z_delta ~ std_normal();
  z_beta ~ std_normal();
  z_Delta ~ std_normal();
  eta_0 ~ student_t(3,0,prior_eta_0);
  for (k in 1:K)
    for (1 in 1:(L-1))
      tau[k, 1] ~ student_t(3, 0, prior_div);
  // outcome model
  for (i in 1:N)
    y[i] ~ ordered_logistic(yhat[i], tau[kk[i]]);
generated quantities {
 real OR;
                           // overall CCP effect (odds ratio)
  real negDelta;
                           // overall CCP effect on log scale
```

```
OR = exp(-Delta);
negDelta=-1*Delta;
}
```