# Nested Pair-Design Hierarchical Model

### A short simulation study

Version 0.1

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#### **Contents**

1	Organise R Session	2
2	Simulation of Measurements	3
3	Modeling	5
4	Results	7

## 1 Organise R Session

## Organise R Session: ####
rm(list = ls())
library("countreg")
library("rcartocolor")
library("brms")

#### 2 Simulation of Measurements

Design:

- ten sites
- · one to five pairs per site
- · one measurement for each element of a pair

$$y_i \sim \text{Normal}\left(\mu_i, \sigma^2\right)$$

index i for the basic observation units, population variance  $\sigma^2=1$ , and:

$$\mu_i = 0 + 0.5 \cdot I_{\{x_i = \text{Treatment}\}} + \gamma_i,$$

with indicator function I:

$$I_{\{\text{Condition}\}} = \begin{cases} 0, & \text{if condition is not met} \\ 1, & \text{if condition is met} \end{cases}$$

and

$$\gamma_i = \sum_{k=1}^{10} \left( \mathrm{I}_{\{\mathsf{site}_i = k\}} \left( \gamma_k + \sum_{j=1}^{n_k} \mathrm{I}_{\{\mathsf{site}_i = k \text{ and } \mathsf{pair}_i = j\}} \gamma_{k_j} \right) \right)$$

with

$$\gamma_k \sim ext{Normal}\left(0, \sigma_{ ext{site}}^2\right), \qquad \gamma_{k_j} \sim ext{Normal}\left(0, \sigma_{ ext{pair}}^2\right).$$

```
set.seed(123456789)
sites <- LETTERS[1:10]
n_pairs_per_site <- sample(1:5, length(sites), replace = T)</pre>
df <- data.frame(site = NULL, pair = NULL, mu_k = NULL, mu_j = NULL,</pre>
                 stringsAsFactors = FALSE)
mu_k <- rnorm(length(sites))</pre>
for (k in 1:length(sites)) {
 mu_j <- rnorm(n_pairs_per_site[k])</pre>
 for (j in 1:n_pairs_per_site[k]) {
    df_k <- data.frame(site = LETTERS[k],</pre>
                        pair = letters[j],
                        x = c("Control", "Treatment"),
                        mu_k = mu_k[k],
                        mu_j = mu_j[j],
                        mu_x = c(0, 0.5),
                        stringsAsFactors = FALSE)
    df <- rbind(df, df_k)
df$y <- df$mu_k + df$mu_j + df$mu_x + rnorm(nrow(df))</pre>
```

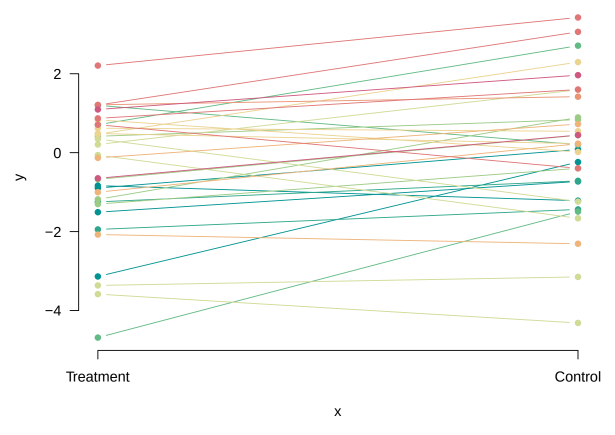


Figure 1: Simulated data of pairs (one element treatment with one element control) one ten different sited (colors).

## 3 Modeling

For comparison, a reduced model is constructed without including the structural information by the pair design:

$$\gamma_i = \sum_{k=1}^{10} \mathbf{I}_{\{\text{site}_i = k\}} \gamma_k$$

In such a model, the present variation by  $\gamma_{k_j}$  is not correctly incorporated. As per site, the variation by pairs is symmetrical and central at 0, this non-inclusion should lead to an inflation of the population variation. This model compared to the correctly constructed model as defined by the simulation build-up. As the strength of the true *signal* of interest – the influence of x on the expected value of y – is not changed between the models, such an inflation of the estimated population variation should lead to a lower precision in the estimation of the signal.

```
# frmla_0 <- bf(y ~ x + (1 | site))
# frmla <- bf(y ~ x + (1 | site/pair))</pre>
# pr_0 <- get_prior(frmla_0, data = df, family = "normal")</pre>
# pr <- get_prior(frmla, data = df, family = "normal")</pre>
# pr_0[pr_0$class == "b" & pr_0$coef == "xTreatment", "prior"] <- "normal(0, 2)"</pre>
# pr[pr$class == "b" & pr$coef == "xTreatment", "prior"] <- "normal(0, 2)"</pre>
# # dummy_model_0 <- brm(frmla_0, data = df, family = "normal",
                           prior = pr_0, chains = 0)
# # dummy_model <- brm(frmla, data = df, family = "normal",</pre>
# # prior = pr, chains = 0)
# # save(dummy_model_0, file = "dummy_model_0_paired_design_20200622.RData")
# # save(dummy_model, file = "dummy_model_paired_design_20200622.RData")
# load(file = "dummy_model_0_paired_design_20200622.RData")
# load(file = "dummy_model_paired_design_20200622.RData")
# results <- vector("list", R)</pre>
# set.seed(123456789)
# for (r in 1:R) {
   n_pairs_per_site <- sample(1:5, length(sites), replace = T)</pre>
    df <- data.frame(site = NULL, pair = NULL, mu_k = NULL, mu_j = NULL,</pre>
                       stringsAsFactors = FALSE)
    mu k <- rnorm(length(sites))
#
    for (k in 1:length(sites)) {
      mu_j <- rnorm(n_pairs_per_site[k])</pre>
      for (j in 1:n_pairs_per_site[k]) {
         df_k <- data.frame(site = LETTERS[k],</pre>
                              pair = letters[j],
                              x = c("Control", "Treatment"),
                              mu_k = mu_k[k],
                              mu_j = mu_j[j],
                              mu_x = c(0, 0.5),
#
                              stringsAsFactors = FALSE)
#
         df <- rbind(df, df_k)
#
    \label{eq:dfsmu_k + dfsmu_j + dfsmu_x + rnorm(nrow(df))} df\$y <- df\$mu_k + df\$mu_j + df\$mu_x + rnorm(nrow(df))
#
    fit_0 <- update(dummy_model_0, newdata = df, recompile = FALSE,</pre>
                      control = list(adapt_delta = 0.8, max_treedepth = 10),
                      chains = 4.
                      cores = 4,
                      iter = 2000
                      warmup = 1000,
                      sample_prior = "no",
                      seed = 123456789
                      refresh = 1000)
    fit <- update(dummy_model, newdata = df, recompile = FALSE,</pre>
                    control = list(adapt_delta = 0.8, max_treedepth = 10),
                    chains = 4,
                    cores = 4.
                    iter = 2000
                    warmup = 1000,
                    sample_prior = "no".
                    seed = 123456789,
```

```
refresh = 1000)
#
#
   tmp_0 <- fitted(fit_0, re_formula = NA, summary = FALSE,</pre>
                  newdata = data.frame(x = c("Control", "Treatment")))
#
   #
   res_0 <- apply(tmp_0, MAR = 1, FUN = diff)
   res <- apply(tmp, MAR = 1, FUN = diff)
results[[r]] <- cbind(res_0, res,
                        as.matrix(fit_0)[, "sigma"], as.matrix(fit)[, "sigma"],
                        as.matrix(fit_0)[, "sd_site__Intercept"], as.matrix(fit)[, "sd_site__Intercept"])
#
# }
# save(results, file = "results_paired_design_20200622.RData")
load(file = "results_paired_design_20200622.RData")
```

### 4 Results

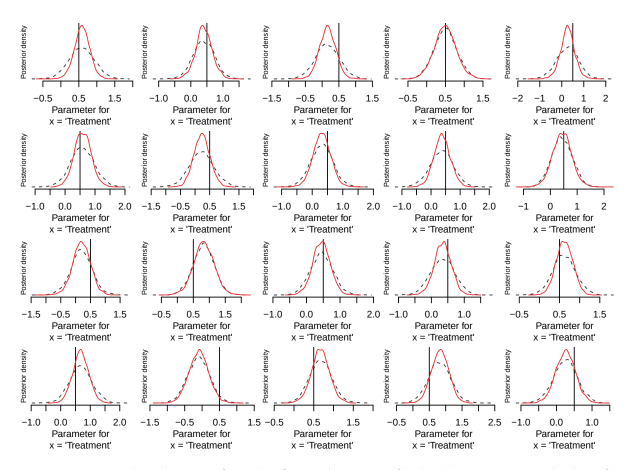


Figure 2: Posterior distribution – for each of 20 replications – for the change in expected value of y when changing x from control to treatment – underlying truth is 0.5. Each dashed black line is a kernel density estimator of the distribution estimated by model without including the structural information by the pair design, solid red line is the respective density estimator from the model including the structural pair information.

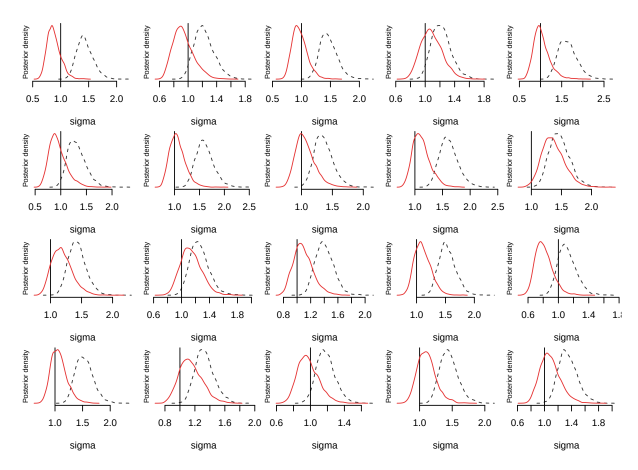


Figure 3: Posterior distribution – for each of 20 replications – for the population variation. Each dashed black line is a kernel density estimator of the distribution estimated by model without including the structural information by the pair design, solid red line is the respective density estimator from the model including this structural pair information.

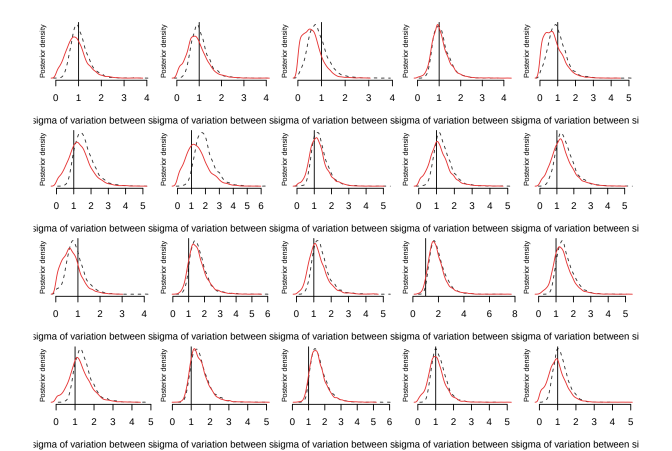


Figure 4: Posterior distribution – for each of 20 replications – for the variation between sites. Each dashed black line is a kernel density estimator of the distribution estimated by model without including the structural information by the pair design, solid red line is the respective density estimator from the model including this structural pair information.