# University of Edinburgh, School of Mathematics Incomplete Data Analysis, 2021/2022

#### Bivariate normal data: one variable subject to missigness

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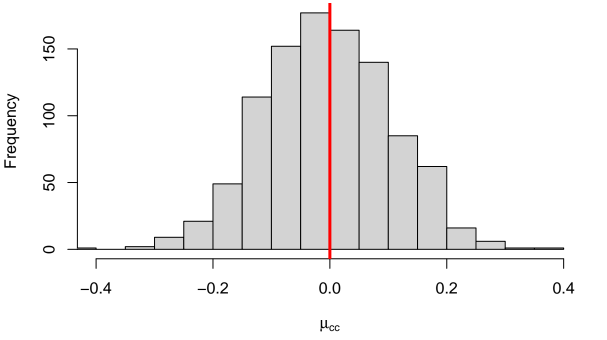
We start by simulating 1000 data sets of size n = 100 from a bivariate normal distribution with the following structure

$$\mu = (0,0)', \quad \Sigma = \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix},$$

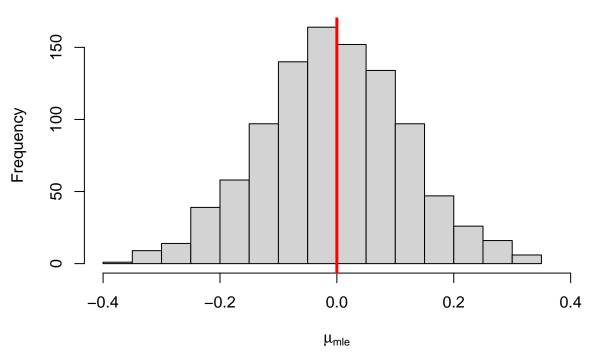
where  $\rho \in \{0, 0.5, 0.9\}$ . Note that when  $\rho = 0$ ,  $Y_1$  and  $Y_2$  are independent. I then created a function that takes as input the 1000 simulated bivariate data sets and the value of a used to induce missingness. I will start with the case of  $\rho = 0$  and I have considered a = 0.8. Code follows.

```
require(MASS)
rho <- 0
n <- 100
nsim <- 1000
mu \leftarrow c(0,0)
sigma1 <- 1
sigma2 <- 1
Sigma <- matrix(c(sigma1^2, rho*sigma1*sigma2, rho*sigma1*sigma2, sigma2^2),
                  nrow = 2, ncol = 2, byrow = T)
y \leftarrow array(0, c(n, 2, nsim))
set.seed(1)
for(l in 1:nsim){
y[, , 1] <- mvrnorm(n, mu = mu, Sigma = Sigma)
bivnorm <- function(y, a){
  n \leftarrow dim(y)[1]
  nsim \leftarrow dim(y)[3]
  y1 \leftarrow y2 \leftarrow r \leftarrow matrix(0, nrow = n, ncol = nsim)
  mu2mle <- mu2cc <- numeric(nsim)</pre>
  treshold <- qnorm(a, 0, 1)</pre>
  for(l in 1:nsim){
    y1[, 1] \leftarrow y[, 1, 1]
    y2[, 1] \leftarrow y[, 2, 1]
    r[,1] \leftarrow ifelse(y1[, 1] > treshold, 0, 1)
```

#### Complete cases, rho = 0

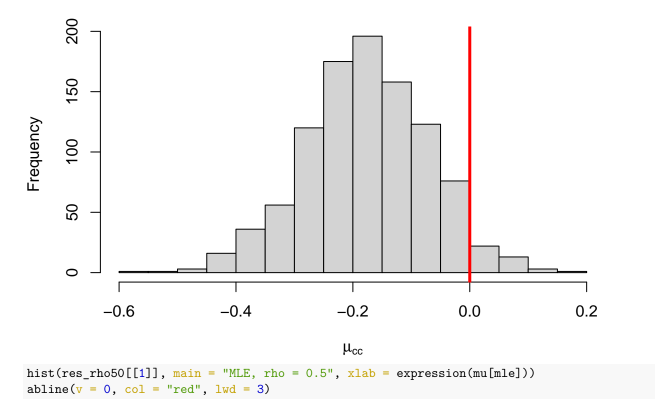


#### MLE, rho = 0

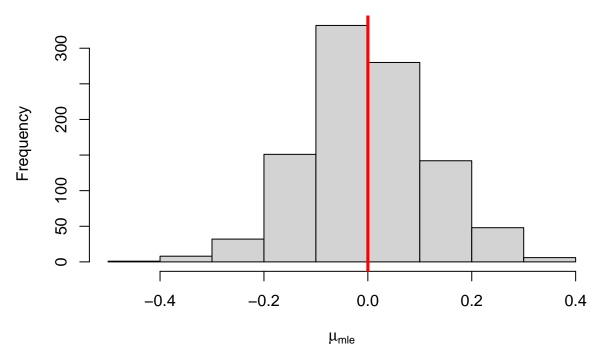


The results depicted in the histograms are not surprising, as in the case of independence between  $Y_1$  and  $Y_2$ ,  $s_{12}$  should be close to 0, and therefore  $\hat{\beta}_1 \approx 0$ ,  $\hat{\beta}_1 \approx \bar{y}_2$ , and thus both the complete cases and the mle estimators coincide. We will now simulate data with  $\rho = 0.5$  and we expect the performance of the complete cases estimator to deteriorate, whereas the mle should provide correct estimates.

### Complete cases, rho = 0.5

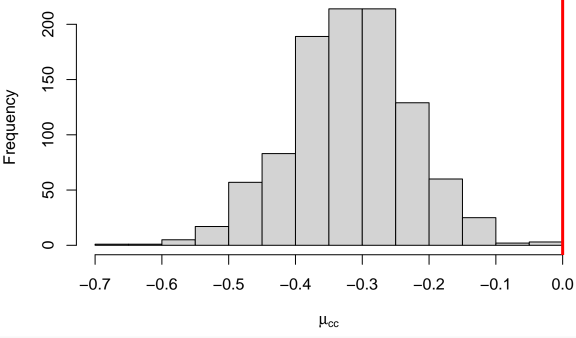


## MLE, rho = 0.5



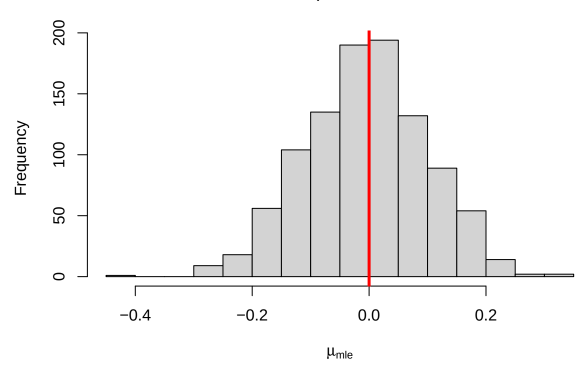
The plots confirm what we just stated we should expect. Now we will try  $\rho = 0.9$  and we should expect the complete cases estimator to perform even worse.

### Complete cases, rho = 0.9



hist(res\_rho90[[1]], main = "MLE, rho = 0.9", xlab = expression(mu[mle]))
abline(v = 0, col = "red", lwd = 3)

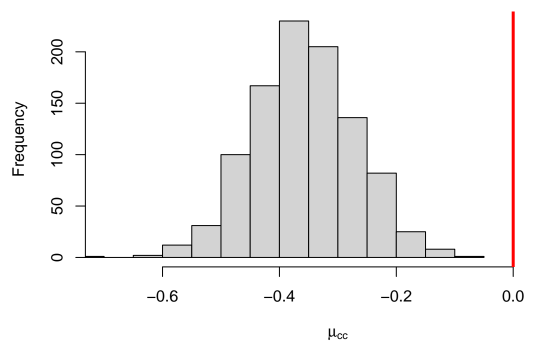
#### MLE, rho = 0.9



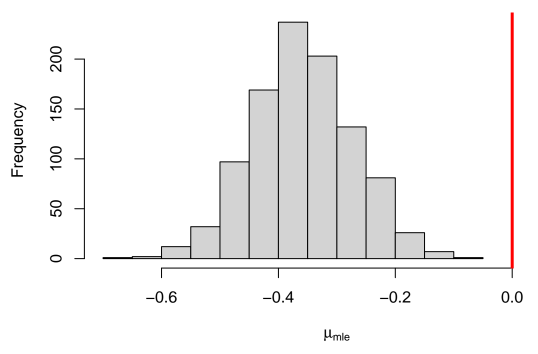
We will now break the assumption of MAR data and, instead, simulate MNAR data. We follow exactly the same procedure, with the only difference being that  $r_i = 0$  (i.e.,  $y_{2i}$  is missing) if  $y_{2i} > z_p$ , where  $z_p = \Phi^{-1}(a)$ , and a = 0.8. As we know from one of the last week examples, we should expect the mle to be severely biased in this situation.

```
rho <- 0
Sigma <- matrix(c(sigma1^2, rho*sigma1*sigma2, rho*sigma1*sigma2, sigma2^2),
                  nrow = 2, ncol = 2, byrow = T)
y \leftarrow array(0, c(n, 2, nsim))
set.seed(1)
for(l in 1:nsim){
y[, , 1] <- mvrnorm(n, mu = mu, Sigma = Sigma)
bivnorm_mnar <- function(y, a){</pre>
  n \leftarrow dim(y)[1]
  nsim \leftarrow dim(y)[3]
  y1 \leftarrow y2 \leftarrow r \leftarrow matrix(0, nrow = n, ncol = nsim)
  mu2mle <- mu2cc <- numeric(nsim)</pre>
  treshold <- qnorm(a, 0, 1)</pre>
  for(l in 1:nsim){
    y1[, 1] <- y[, 1, 1]
    y2[, 1] \leftarrow y[, 2, 1]
    r[,1] \leftarrow ifelse(y2[, 1] > treshold, 0, 1)
    s1squared \leftarrow mean((y1[r[, 1] == 1, 1] - mean(y1[r[, 1] == 1, 1]))^2)
    s12 \leftarrow mean((y1[r[, 1] == 1, 1] -
                      mean(y1[r[, 1] == 1, 1]))*(y2[r[, 1] == 1, 1] - mean(y2[r[, 1] == 1, 1])))
```

### Complete cases, rho = 0, MNAR

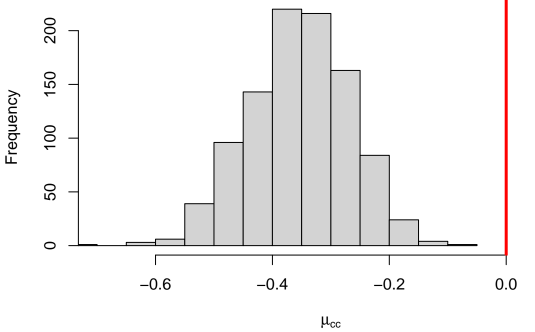


#### MLE, rho = 0, MNAR

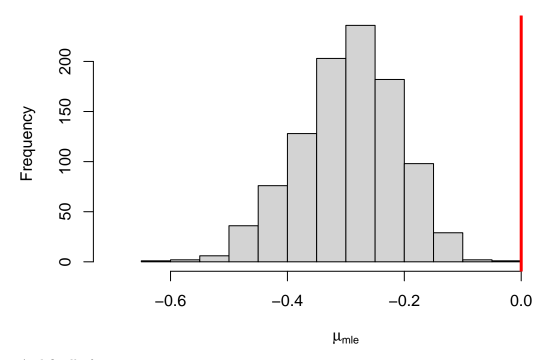


Now trying  $\rho = 0.5$ .

### Complete cases, rho = 0.5, MNAR

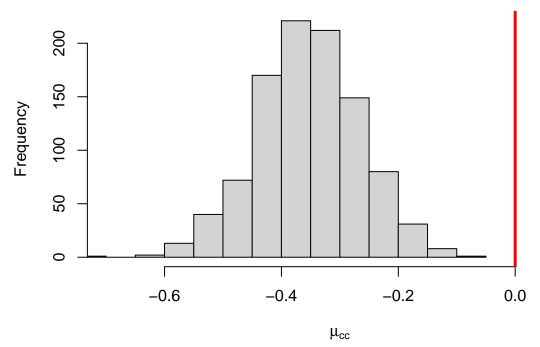


MLE, rho = 0.5, MNAR



And finally for  $\rho = 0.9$ .

#### Complete cases, rho = 0.9, MNAR



## MLE, rho = 0.9, MNAR

