## Covariate misclassification from a latent Gaussian variable with measurement error

Code and details for 'Bayesian models for missing and misclassified variables using integrated nested Laplace approximations'

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```
library(INLA)
library(dplyr)
library(ggplot2)
```

## Models adjusting for misclassification arising from a dichotomized variable with error

In this example, we simulate a continuous latent variable with measurement error, which is then dichotomized.

```
set.seed(1)
n <- 200

x_c <- rnorm(n, 0, 1) # Continuous, not observed

x_d <- ifelse(x_c > 0, 1, 0) # Dichotomized, not observed

w_c <- x_c + rnorm(n) # Continuous with cont. ME, may be observed

w_d <- ifelse(w_c > 0, 1, 0) # Dichotomized, observed
```

From the variables we have simulated, we can generate a sample misclassification matrix:

```
## observed

## unobserved 0 1

## 0 0.6603774 0.3396226

## 1 0.2872340 0.7127660
```

Lastly, we generate the response.

```
y \leftarrow 1 + x_c + rnorm(n)
```

Now, the observed data is y and  $w_d$  (often, researchers choose to use a dichotomized variable even though the continuous version is available, so perhaps  $w_c$  is also observed, but simply ignored, or it may have been

available at some point but was not sent to the statistician). For the purposes of this example, we assume that we are interested in the relationship between y and  $x_c$ .

As described in Section 3.2.2, we then have the model

$$egin{aligned} oldsymbol{y} &= eta_0 oldsymbol{1} + eta_{x_c} oldsymbol{x}_c + oldsymbol{Z}^ op oldsymbol{eta} + oldsymbol{arepsilon} \ oldsymbol{w}_c &= oldsymbol{x}_c + oldsymbol{u}_{w_c} \ , & oldsymbol{u}_{w_c} \sim \mathcal{N}(oldsymbol{0}, \sigma_{w_c}^2 oldsymbol{I}) \ , \ oldsymbol{w}_d &= \mathrm{I}(oldsymbol{w}_c > 0) \ , \ oldsymbol{x}_c &= lpha_0 oldsymbol{1} + oldsymbol{\widetilde{Z}}^ op oldsymbol{lpha} + oldsymbol{arepsilon}_{x_c} \ , & oldsymbol{arepsilon}_{x_c} \sim \mathcal{N}(oldsymbol{0}, \sigma_{x_c}^2 oldsymbol{I}) \ , \end{aligned}$$

where Z and  $\widetilde{Z}$  are both covariate matrices with whichever covariates are relevant to include, these are assumed to be without error. Now, we have that

$$\Pr(w_{d,i} = 1 \mid w_{c,i}) = \Pr(w_{c,i} > 0)$$

$$= \Pr(x_{c,i} + u_{w_c,i} > 0)$$

$$= \Pr\left(\frac{u_{w_c,i}}{\sigma_{w_c}} < \frac{x_{c,i}}{\sigma_{w_c}}\right)$$

$$= \Phi\left(\frac{x_{c,i}}{\sigma_{w_c}}\right),$$

since  $u_{w_c,i}/\sigma_{w_c} \sim \mathcal{N}(0,1)$ . This relation means that we can model  $\boldsymbol{w}_d$  as a Bernoulli variable with a probit link function. We can re-write the hierarchical model from above:

$$egin{aligned} oldsymbol{y} &= eta_0 oldsymbol{1} + eta_{x_c} oldsymbol{x}_c + oldsymbol{Z}^ op oldsymbol{eta} + oldsymbol{arepsilon} \ oldsymbol{w}_d \sim ext{Bernoulli}(\Phi(oldsymbol{x}_c/\sigma_{w_c})) \ , \ oldsymbol{x}_c &= lpha_0 oldsymbol{1} + \widetilde{oldsymbol{Z}}^ op lpha + oldsymbol{arepsilon}_{x_c} \ , \ oldsymbol{arepsilon}_{x_c} \sim \mathcal{N}(oldsymbol{0}, \sigma_{x_c}^2 oldsymbol{I}) \ , \end{aligned}$$

We use stacks in R-INLA to structure the hierarchical model:

```
stk_y <- inla.stack(data = list(y = y),</pre>
                       A = list(1),
                       effects = list(
                         list(beta.0 = rep(1, n),
                              beta.x_c = 1:n)),
                       tag = "y")
stk_w_d <- inla.stack(data = list(w_d = w_d),</pre>
                       A = list(1),
                       effects = list(
                         list(id.x_c = 1:n,
                              weight.x_c = 1),
                       tag = "w_d")
stk_x_c <- inla.stack(data = list(x_c = rep(0, n)),
                       A = list(1),
                       effects = list(
                         list(id.x_c = 1:n,
                              weight.x_c = -1,
                              alpha.0 = rep(1, n))),
                       tag = "x_c")
stk_full <- inla.stack(stk_y, stk_w_d, stk_x_c)</pre>
```

```
formula \leftarrow list(y, w_d, x_c) \sim -1 + beta.0 + alpha.0 +
  f(beta.x_c, copy = "id.x_c",
    hyper = list(beta = list(param = c(0, 1/1000), fixed = FALSE))) +
  f(id.x_c, weight.x_c, model = "iid", values = 1:n,
    hyper = list(prec = list(initial = -15, fixed = TRUE)))
res <- inla(formula, data = inla.stack.data(stk_full), Ntrials = rep(1, n),
            family = c("gaussian", "binomial", "gaussian"),
            control.family = list(
              list(hyper = list(prec = list(initial = log(1),
                                             param = c(10, 9),
                                             fixed = FALSE))),
              list(link = "probit"),
              list(hyper = list(prec = list(initial = log(1),
                                             param = c(10, 9),
                                             fixed = FALSE)))
            control.predictor = list(compute = TRUE))
```

## Looking at the results

##

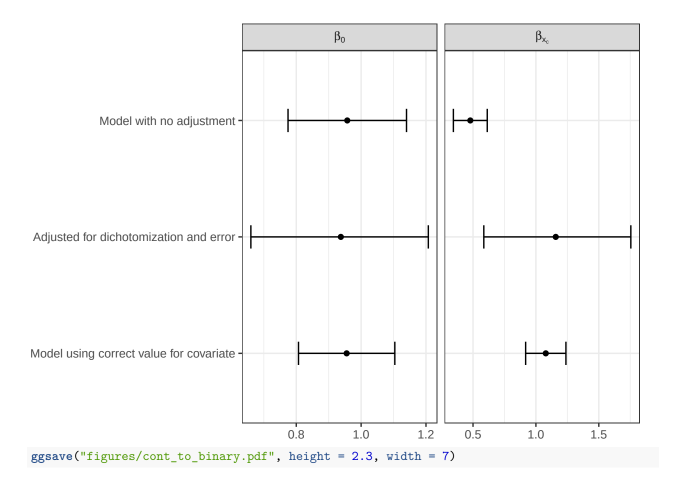
```
data \leftarrow data.frame(y = y, x_c = x_c, w_c = w_c, w_d = w_d, x_d = x_d)
Results from using correct continuous variable:
correct_model <- inla(y ~ x_c, data = data)</pre>
correct_model$summary.fixed
##
                                  sd 0.025quant 0.5quant 0.975quant
                    mean
## (Intercept) 0.9553833 0.07565846 0.8068701 0.9553833 1.103896 0.9553833
               1.0785024 0.08157634 0.9183726 1.0785024 1.238632 1.0785024
## x_c
##
## (Intercept) 1.943436e-09
               1.943381e-09
Results from using dichotomized version but continuous latent variable (our proposed model):
res$summary.fixed["beta.0",]
##
                           sd 0.025quant 0.5quant 0.975quant
                                                                                  kld
## beta.0 0.9374329 0.138637 0.6600015 0.938767 1.207244 0.938742 7.090626e-08
res$summary.hyperpar["Beta for beta.x_c",]
                                      sd 0.025quant 0.5quant 0.975quant
                          mean
## Beta for beta.x_c 1.157789 0.2969229 0.5857856 1.153582
                                                                 1.754753 1.135542
Results from using continuous variable with measurement error with no correction:
naive_model <- inla(y ~ w_c, data = data)</pre>
naive_model$summary.fixed
                                  sd 0.025quant 0.5quant 0.975quant
                    mean
## (Intercept) 0.9572666 0.09306920 0.7745771 0.9572666 1.1399560 0.9572666
```

 $0.4784398 \ 0.06858374 \quad 0.3438138 \ 0.4784398 \quad 0.6130657 \ 0.4784398$ 

kld

## (Intercept) 1.942828e-09

```
## w c
               1.942792e-09
adjusted_results <- rbind("(Intercept)" = res$summary.fixed["beta.0",1:5],</pre>
                           "x_c" = res$summary.hyperpar["Beta for beta.x_c",1:5])
adjusted_results$variable <- rownames(adjusted_results)</pre>
naive_results <- naive_model$summary.fixed</pre>
rownames(naive_results) <- c("(Intercept)", "x_c")</pre>
naive results$variable <- rownames(naive results)</pre>
correct_results <- correct_model$summary.fixed</pre>
correct_results$variable <- rownames(correct_results)</pre>
all res <- dplyr::bind rows(adjusted = adjusted results,
                             naive = naive_results,
                             correct = correct_results, .id = "Model")
all_res$labels <- paste0("beta", "[", c(0, "x[c]"), "]")</pre>
all_res$Model <- factor(all_res$Model, levels = c("naive", "adjusted", "correct"))
all_res$Model <- plyr::revalue(all_res$Model,
                                c("naive" = "Model with no adjustment",
                                  "adjusted" = "Adjusted for dichotomization and error",
                                  "correct" = "Model using correct value for covariate"))
ggplot(all_res, aes(y = Model)) +
  geom point(aes(x = mean)) +
  geom_errorbarh(aes(xmin = .data$"0.025quant", xmax = .data$"0.975quant"), height = .2) +
  scale_y_discrete(limits = rev) +
  facet_wrap(vars(labels), scales = "free_x", labeller = label_parsed) +
  theme bw() +
  theme(axis.title = element_blank())
```



## How do different measurement error variances correspond to different misclassification probabilities?

Using simulated data as well as the exact sensitivity and specificity calculated given our simulation setup, we can take a look at how the sensitivity and specificity of the classification of the continuous variable change for increasing measurement error variance.

First, we simulate data for different levels of measurement error:

```
simulate_dichotomized <- function(me_variance = 1, n = 200){
    x_c <- rnorm(n, 0, 1) # Continuous, not observed
    x_d <- ifelse(x_c > 0, 1, 0) # Dichotomized, not observed
    w_c <- x_c + rnorm(n, 0, sqrt(me_variance)) # Continuous with cont. ME, may be observed
    w_d <- ifelse(w_c > 0, 1, 0) # Dichotomized, observed

return(list(x_d = x_d, w_d = w_d))
}

calculate_mc <- function(x_d, w_d){
    pix0 <- sum(x_d==0)
    pix1 <- sum(x_d==1)
    mc_tab <- table(unobserved = x_d, observed = w_d)/
    matrix(c(pix0, pix0, pix1, pix1), byrow = TRUE, nrow = 2)
    return(list(spec = mc_tab[1,1], sens = mc_tab[2,2]))
}</pre>
```

```
me_variances <- seq(0.01, 2, by = 0.05)
result_df <- data.frame(me_var = NA, sens = NA, spec = NA)
for(i in 1:length(me_variances)){
   var_list <- simulate_dichotomized(me_variance = me_variances[i])
   sens_spec <- calculate_mc(var_list$x_d, var_list$w_d)
   result_df[i,] <- c(me_variances[i], sens_spec$sens, sens_spec$spec)
}</pre>
```

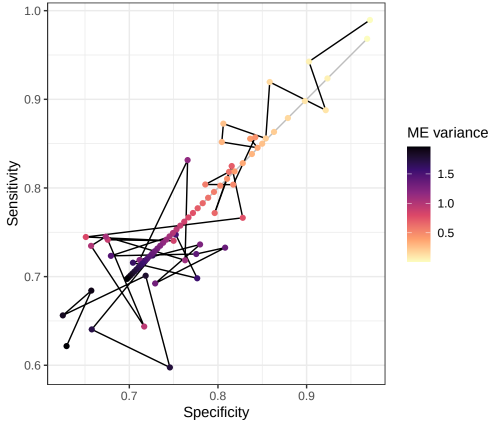
Next, we also calculate the expected sensitivity and specificity for different values of measurement error variance given the simulation setup:

```
sens_spec_exact <- function(sigma_u2 = 1){</pre>
  mu x <- 0
  mu_v <- mu_x
  sigma x2 <- 1
  sigma_v2 <- sigma_x2 + sigma_u2
  cov_v_x \leftarrow sigma_x^2 + 2*mu_x^2
  sigma_mat <- matrix(c(sigma_v2, cov_v_x, sigma_x2, cov_v_x), byrow = TRUE, nrow = 2)</pre>
  # P(V>0, X>0)
  p_v x \leftarrow mvtnorm::pmvnorm(c(0, 0), mean = c(mu_v, mu_x), sigma = sigma_mat)
  # P(X>0)
  p_x <- pnorm(0, mean = mu_x, sd = sqrt(sigma_x2), lower.tail = FALSE)</pre>
  \# P(V>0)
  p_v <- pnorm(0, mean = mu_v, sd = sqrt(sigma_v2), lower.tail = FALSE)</pre>
  # Sensitivity
  sens \leftarrow p_v_x/p_x
  # Specificity
  spec <- (1-(p_v + p_x - p_v_x))/(1-p_x)
  return(list(sens_calc = sens, spec_calc = spec, me_var = sigma_u2))
sens_spec_list <- lapply(me_variances, sens_spec_exact)</pre>
exact_sens_spec <- as.data.frame(do.call(rbind, sens_spec_list)) %>%
  mutate(across(everything(), as.numeric))
```

We combine the results and plot the sensitivity and specificity for different levels of measurement error variance:

```
sim_and_calc <- inner_join(exact_sens_spec, result_df, by = "me_var")

ggplot(sim_and_calc) +
   geom_path(aes(x = spec_calc, y = sens_calc), color = "grey") +
   geom_path(aes(x = spec, y = sens)) +
   geom_point(aes(x = spec, y = sens, color = me_var)) +
   geom_point(aes(x = spec_calc, y = sens_calc, color = me_var)) +
   scale_color_viridis_c(option = "magma", direction = -1) +
   coord_equal() +
   labs(x = "Specificity", y = "Sensitivity", color = "ME variance") +
   theme_bw()</pre>
```



Specificity This plot shows the resulting sensitivity and specificity when the measurement error variance goes from 0.01 to 2.