refitME: Tutorial for fitting measurement error models using Monte Carlo Expectation Maximization in R

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This tutorial documents fitting an MCEM algorithm via the refitME R-package. For more specific details see: refitME: Measurement Error Modelling using Monte Carlo Expectation Maximization in R. Also, see ?refitME for further details on the fitting function, input arguments and output.

Example 1: A simple GLM example taken from Carroll et al. (2006).

We use the Framingham heart study data set. In addition to the naive model, we also fit a simex model and compare it with MCEM. Computational times for both models are also reported.

Load data and R-packages.

```
suppressWarnings(suppressMessages(library(refitME)))
suppressWarnings(suppressMessages(library(simex)))
set.seed(2020)
data(Framinghamdata)
```

Fit the naive model.

The first stored variable w1 is the error contaminated predictor as used in the original Carroll $et\ al.$ (2006) analysis.

Setup measurement error (ME) variance and all tuning parameters.

```
sigma.sq.u \leftarrow 0.01259/2 \ \# \ \textit{ME variance, as obtained from Carroll et al. (2006)}. \\ B \leftarrow 100 \ \# \ \textit{The number of Monte Carlo replication values/SIMEX simulations}.
```

Fit the SIMEX model.

Fit the MCEM model.

```
start <- Sys.time()
glm_MCEM <- refitME(glm_naiv, sigma.sq.u, B)

## One specified error-contaminated predictor/covariate.

## convergence :-)

## [1] 5

end <- Sys.time()
t2 <- difftime(end, start, units = "secs")
comp.time <- c(comp.time, t2)</pre>
```

Report model estimates and compare computational times.

```
est.beta <- rbind(coef(glm_naiv), coef(glm_simex), coef(glm_MCEM))</pre>
est.beta.se <- rbind(sqrt(diag(vcov(glm_naiv))),</pre>
                  sqrt(diag(glm_simex$variance.jackknife)), sqrt(diag(vcov(glm_MCEM))))
row.names(est.beta) = row.names(est.beta.se) <- c("Naive GLM", "SIMEX", "MCEM")
colnames(est.beta) = colnames(est.beta.se) <- c("(Intercept)", "SBP", "chol. level",</pre>
                                               "age", "smoke")
round(est.beta, digits = 3)
             (Intercept)
                          SBP chol. level
                                            age smoke
## Naive GLM
              -14.951 1.707 0.008 0.055 0.592
                -15.919 1.947
                                    0.008 0.053 0.598
## SIMEX
## MCEM
                -16.059 1.955
                                    0.008 0.056 0.594
round(est.beta.se, digits = 3) # Standard error estimates.
            (Intercept) SBP chol. level age smoke
## Naive GLM 1.900 0.418 0.002 0.012 0.250
## SIMEX
                 2.043 0.456
                                  0.002 0.012 0.251
                 2.185 0.487 0.002 0.012 0.250
## MCEM
```

```
names(comp.time) <- c("SIMEX", "MCEM")
comp.time # SIMEX and MCEM.

## Time differences in secs
## SIMEX MCEM
## 7.894655 5.059130
```

Example 2: A GAM example taken from Ganguli et al. (2005).

The Milan mortality air pollution data set is part of the SemiPar package. Here, we fit GAM models via the mgcv package where one predictor (daily total suspended particles measurements) is error-contaminated.

Load data and R-packages.

```
suppressWarnings(suppressMessages(library(refitME)))
suppressWarnings(suppressMessages(library(SemiPar)))
suppressWarnings(suppressMessages(library(mgcv)))
set.seed(2020)
data(milan.mort)
dat.air <- milan.mort</pre>
```

Setup all variables.

```
Y <- dat.air[, 6] # Mortality counts.

n <- length(Y)

z1 <- (dat.air[, 1])

z2 <- (dat.air[, 4])

z3 <- (dat.air[, 5])

w1 <- log(dat.air[, 9]) # The error-contaminated predictor (total suspended particles).
dat <- data.frame(cbind(Y, w1, z1, z2, z3))
```

Fit the naive model.

```
gam_naiv \leftarrow gam(Y \sim s(w1) + s(z1, k = 25) + s(z2) + s(z3), family = "poisson", data = dat)
```

Fit the MCEM model.

Note this will take a fair while to run if B is large (consider decreasing B if you want a rough answer).

```
sigma.sq.u <- 0.0915 # This gives a reliability ratio of 0.7.
rel.rat <- round((1 - sigma.sq.u/var(dat$w1))*100, digits = 0)
gam_MCEM1 <- refitME(gam_naiv, sigma.sq.u)
## One specified error-contaminated predictor/covariate.
## convergence :-)</pre>
## [1] 14
```

MCEM (Poisson GAM) fitted to the air pollution data.

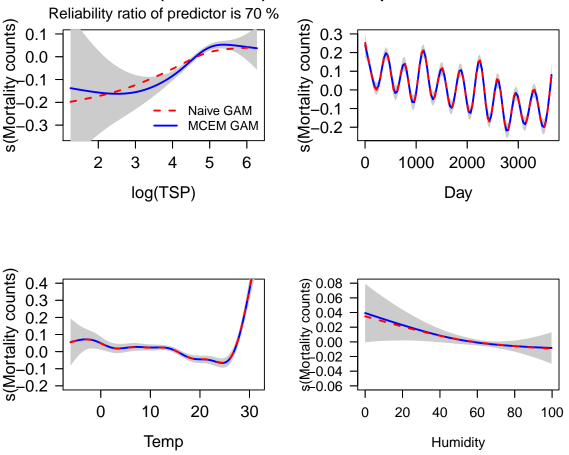


Figure 1: Plots of smooths against each predictor. TSP (top left is the error contaminated variable).

Example 3: A point-process model using presence-only data

We use the *Corymbia eximia* presence-only data set from Renner and Warton (2013). Here, we fit a naive models point-process model (PPM) and an MCEM PPM.

Load data and R-packages.

```
suppressWarnings(suppressMessages(library(refitME)))
suppressWarnings(suppressMessages(library(caret)))
data(Corymbiaeximiadata)
dat <- Corymbiaeximiadata</pre>
```

Setup all variables.

Fit the naive (PPM) model.

```
PPM_naiv1 <- glm(Y ~ X1 + X2 + Z1 + Z2 + Z3 + Z4, family = "poisson",
    weights = p.wt, data = dat1)</pre>
```

Warning: glm.fit: fitted rates numerically 0 occurred

Fit the MCEM PPM.

```
sigma.sq.u <- 0.25

PPM_MCEM1 <- refitME(PPM_naiv1, sigma.sq.u)

## One specified error-contaminated predictor/covariate.

## convergence :-)

## [1] 4

## Warning in limits.and.aspect(default.prepanel, prepanel = prepanel, have.xlim = ## have.xlim, : 'aspect="iso"' approximate since 'relation="free"'</pre>
```

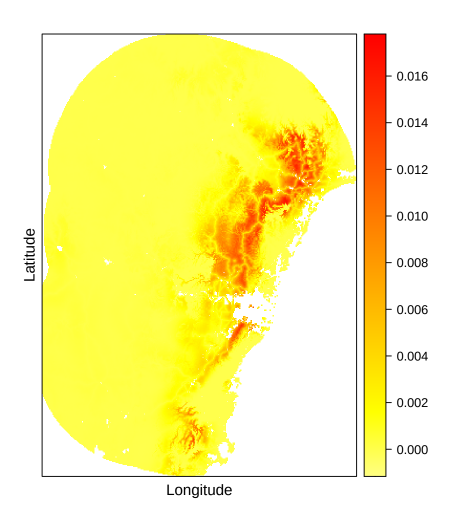


Figure 2: Plot of predicted presences of Corymbia eximia using presence-only data when fitting the MCEM model. Here the max temperature predictor is assumed to be error-contaminated.

Example 4: A VGAM example using the Prinia flaviventris capture-recapture data.

We use the *Prinia flaviventris* capture-recapture data set from Hwang, Huang and Wang (2007). Here, we fit naive vglm and vgam capture-recapture models, and the MCEM capture-recapture model. For all models we used the posbinomial() family provided in VGAM.

Load data and R-packages.

```
suppressWarnings(suppressMessages(library(refitME)))
suppressMessages(library(VGAM))
data(Priniadata)
```

Setup all variables.

```
tau <- 17 # No. of capture occasions.
w1 <- Priniadata$w1
```

Fit the naive vglm and vgam capture-recapture models.

Fit the MCEM capture-recapture model.

```
sigma.sq.u <- 0.37/var(w1) # Measurement error variance.
B <- 100
CR_MCEM <- refitME(CR_naiv2, sigma.sq.u, B)

## [1] "convergence :-)"
## [1] 9</pre>
```