

# LRMoE DemoData Generation

Spark Tseung

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## Introduction

This document contains the data generation process for the dataset **DemoData** included in the **LRMoE** package. This also serves as an example of using the **dataset.simulator** function included in the package.

## Data Simulation

### Complete Dataset

Suppose there is an auto-insurance company with two lines of business, with a total of 10,000 policies. The policyholder information includes sex (1 for Male and 0 for Female), driver's age (with range 20~80), car age (with range 0~10), and region (1 for urban and 0 for rural). We assume all covariates are uniformly and independently drawn at random.

```
set.seed(7777)
sample.size = 10000
intercept = rep(1, sample.size)
sex = rbinom(sample.size, 1, 0.5)
aged = runif(sample.size, 20, 80)
agec = runif(sample.size, 0, 10)
region = rbinom(sample.size, 1, 0.5)

X = matrix(data = c(intercept, sex, aged, agec, region),
           nrow = sample.size, byrow = FALSE,
           dimnames = list(NULL,
                          c("intercept", "sex", "agedriver", "agecar", "region")))
```

For simplicity, we assume there are two latent risk classes: low (L) and high (H). The characteristics for the high-risk class are male, young age, old car age and urban region. This is specified by the following matrix of logit regression coefficients, where the second row represents the reference class.

```
n.comp = 2
alpha = matrix( c(-0.5, 1, -0.05, 0.10, 1.25,
                  0, 0, 0, 0, 0),
               nrow = n.comp, byrow = TRUE)
```

We consider a two-dimensional response: claim frequency from the first business line, and claim severity from the second business line. For demonstration purposes and for simplicity, we don't consider the same business line to avoid the complication where zero frequency necessarily implies zero severity. The component distributions and their parameters are specified as follows.

```
dim.m = 2
comp.dist = matrix(c("poisson", "ZI-gammacount",
                    "lnorm", "invgauss"),
                  nrow = dim.m, byrow = TRUE)
```

```

zero.prob = matrix(c(0, 0.2,
                    0, 0),
                  nrow = dim.m, byrow = TRUE)

params.list = list( list(c(6), c(30, 0.5)),
                   list(c(4, 0.3), c(20, 20))
                 )

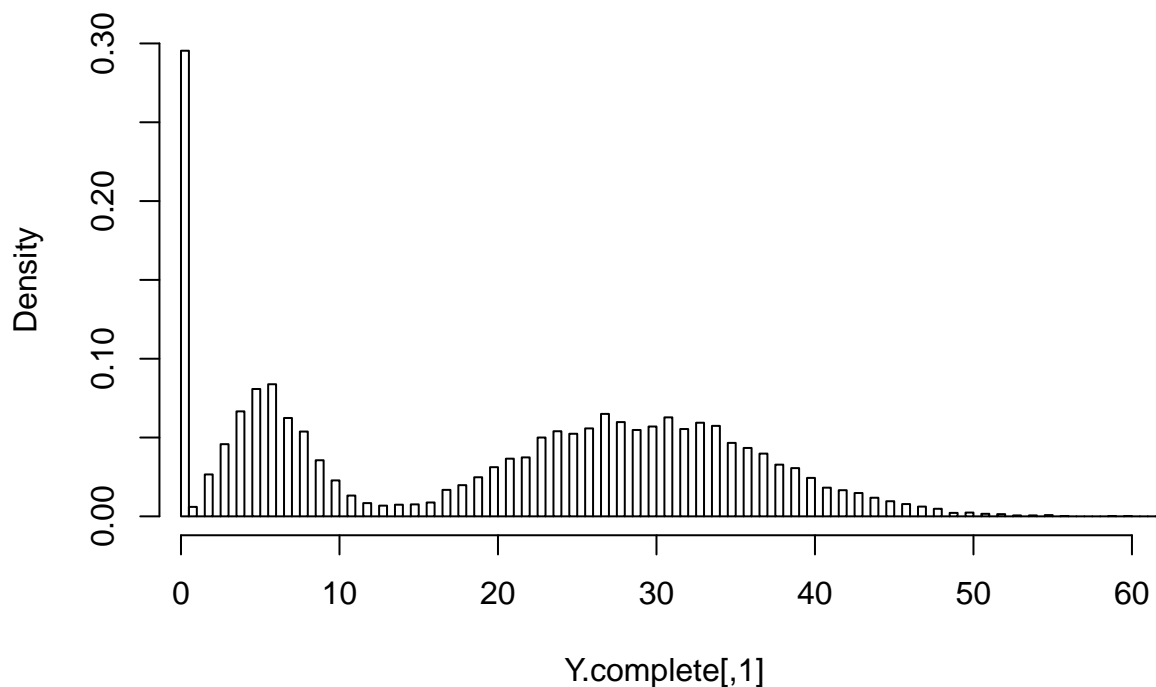
```

The LRMoe package includes a simulator. Given the covariates and parameters defined above, we can directly simulate a dataset.

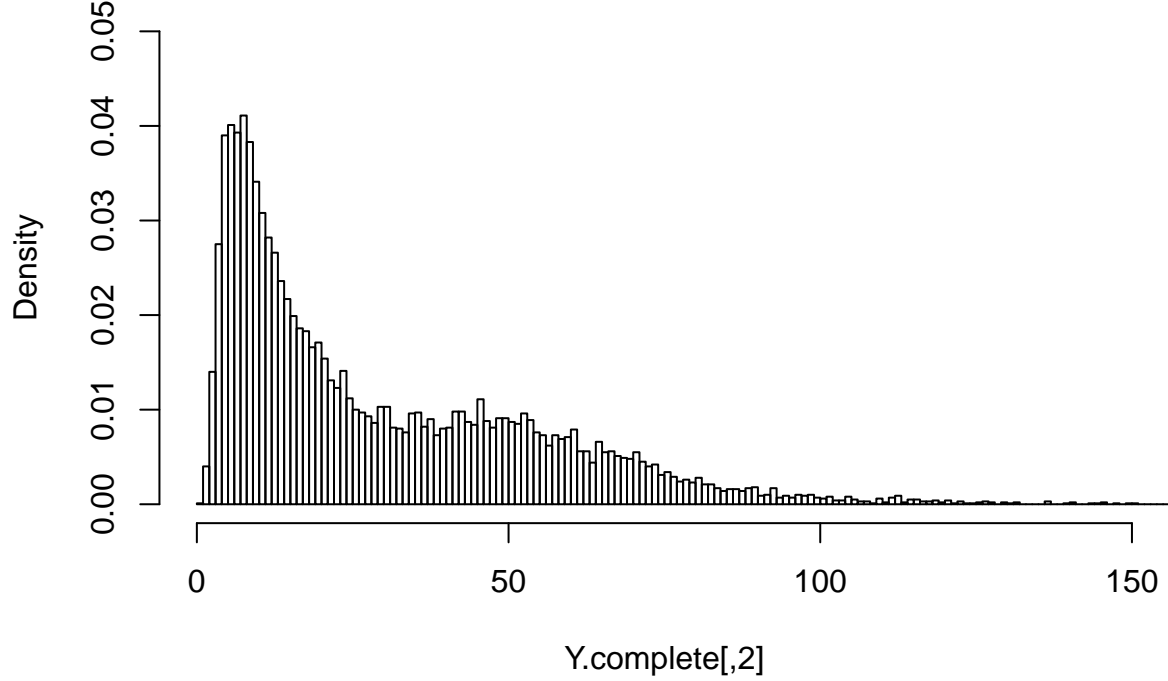
```
Y.complete = LRMoe::dataset.simulator(X, alpha, comp.dist, zero.prob, params.list)
```

The simulated values are plotted as follows. For each dimension of Y, the histogram is relatively well separated as two components. This is more or less done on purpose to demonstrate that the fitting procedure can identify the *true* model when it is known. In practice, we are usually less concerned of the underlying data generating distribution, as long as the LRMoe model provides a reasonable fit of data.

## Histogram of Dimension 1



## Histogram of Dimension 2



### Truncation and Censoring

One distinct feature of LRMoE is dealing with data truncation and censoring, which is common in insurance contexts. Consequently, instead of one single number for each dimension  $d$ , a tuple  $(t1.d, y1.d, yu.d, tu.d)$  is required, where  $t1.d/tu.d$  are the lower/upper bounds of truncation, and  $y1.d/yu.d$  are the lower/upper bounds of censoring.

For illustration purposes, we assume the dataset is subject to the following truncation and censoring.

Index	Y.complete[,1]	Y.complete[,2]
1-6000	No truncation or censoring	No truncation or censoring
6001-8000	No truncation or censoring	Left Truncated at 5
8001-10000	No truncation or censoring	Right Censored at 100

```
# First block: 1~6000
X.obs = X[1:6000,]

t1.1 = rep(0, 6000)
y1.1 = Y.complete[1:6000, 1]
yu.1 = Y.complete[1:6000, 1]
tu.1 = rep(Inf, 6000)

t1.2 = rep(0, 6000)
y1.2 = Y.complete[1:6000, 2]
yu.2 = Y.complete[1:6000, 2]
```

```

tu.2 = rep(Inf, 6000)

# Second block: 6001-8000
keep.idx = Y.complete[6001:8000,2] >= 5
keep.length = sum(keep.idx)

X.obs = rbind(X.obs, X[6001:8000,][keep.idx,])

tl.1 = c(tl.1, rep(0, keep.length))
yl.1 = c(yl.1, Y.complete[6001:8000, 1][keep.idx])
yu.1 = c(yu.1, Y.complete[6001:8000, 1][keep.idx])
tu.1 = c(tu.1, rep(Inf, keep.length))

y.temp = Y.complete[6001:8000, 2][keep.idx]
tl.2 = c(tl.2, rep(5, keep.length))
yl.2 = c(yl.2, Y.complete[6001:8000, 2][keep.idx])
yu.2 = c(yu.2, Y.complete[6001:8000, 2][keep.idx])
tu.2 = c(tu.2, rep(Inf, keep.length))

# Third block: 8001-10000
X.obs = rbind(X.obs, X[8001:10000,])

tl.1 = c(tl.1, rep(0, 2000))
yl.1 = c(yl.1, Y.complete[8001:10000, 1])
yu.1 = c(yu.1, Y.complete[8001:10000, 1])
tu.1 = c(tu.1, rep(Inf, 2000))

y.temp = Y.complete[8001:10000, 2]
censor.idx = which(y.temp >= 100)
yl.temp = y.temp
yl.temp[censor.idx] = 100
yu.temp = y.temp
yu.temp[censor.idx] = Inf
tl.2 = c(tl.2, rep(0, 2000))
yl.2 = c(yl.2, yl.temp)
yu.2 = c(yu.2, yu.temp)
tu.2 = c(tu.2, rep(Inf, 2000))

# Put things together
Y.obs = matrix(c(tl.1, yl.1, yu.1, tu.1, tl.2, yl.2, yu.2, tu.2),
               ncol = 8, byrow = FALSE,
               dimnames = list(NULL,
                               c("tl.1", "yl.1", "yu.1", "tu.1",
                                 "tl.2", "yl.2", "yu.2", "tu.2")))

```

As a result of truncating `Y.complete[,2]`, 172 rows are discarded, leaving 9828 observations available for model fitting. Sample data points are shown below.

```

# No truncation, no censoring
Y.obs[1,]

```

##	tl.1	yl.1	yu.1	tu.1	tl.2	yl.2	yu.2	tu.2
##	0.000000	27.000000	27.000000	Inf	0.000000	8.145064	8.145064	Inf

```
# Y.2 is left-truncated
```

```
Y.obs[6002,]
```

```
##      tl.1      yl.1      yu.1      tu.1      tl.2      yl.2      yu.2      tu.2
## 0.00000 4.00000 4.00000      Inf 5.00000 62.02375 62.02375      Inf
```

```
# Y.2 is right-censored
```

```
Y.obs[7884,]
```

```
##      tl.1      yl.1      yu.1      tu.1      tl.2      yl.2      yu.2      tu.2
## 0.000000 28.000000 28.000000      Inf 0.000000 5.887506 5.887506      Inf
```

We will export both the complete and incomplete datasets to the LRMOE package.

```
Y = matrix(c(rep(0,10000), Y.complete[,1], Y.complete[,1], rep(Inf,10000),
              rep(0,10000), Y.complete[,2], Y.complete[,2], rep(Inf,10000)),
           ncol = 8, byrow = FALSE,
           dimnames = list(NULL,
                           c("tl.1", "yl.1", "yu.1", "tu.1",
                              "tl.2", "yl.2", "yu.2", "tu.2")))
```

```
save(X, Y, X.obs, Y.obs, file = "DemoData.RData")
```