MLE vs OLS

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```
rm(list=ls())
#install.packages(c("MASS", "ggplot2", "tidyverse"))
library(MASS)
library(ggplot2)
## Warning: package 'ggplot2' was built under R version 4.0.5
library(tidyverse)
## Warning: package 'tidyverse' was built under R version 4.0.5
## -- Attaching packages ------ tidyverse 1.3.1 --
## v tibble 3.1.2
                    v dplyr 1.0.7
## v tidyr 1.1.3 v stringr 1.4.0
## v readr 1.4.0 v forcats 0.5.1
## v purrr 0.3.4
## Warning: package 'tibble' was built under R version 4.0.5
## Warning: package 'tidyr' was built under R version 4.0.5
## Warning: package 'readr' was built under R version 4.0.5
## Warning: package 'dplyr' was built under R version 4.0.5
## Warning: package 'forcats' was built under R version 4.0.5
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
## x dplyr::select() masks MASS::select()
#URL_tile <- c("https://faculty.washington.edu/cadolph/software/tile_0.4.14.tar.qz")
#URL_simcf <- c("https://faculty.washington.edu/cadolph/software/simcf_0.2.17.tar.gz")
#install.packages(URL_tile, repos = NULL, type="source")
#install.packages(URL_simcf, repos = NULL, type="source")
library(simcf)
library(tile)
```

Loading required package: grid

Normal MLE

Let's say our dependent variable, y_i , follows this Normal distribution:

$$y_i \sim f_{\mathcal{N}}(\mu_i, \sigma^2)$$

with a mean μ_i defined by the linear equation of covariate matrix, \mathbf{x}_i , multiplied by the coefficient matrix, β :

$$\mu_i = \mathbf{x}_i \boldsymbol{\beta}$$

This is called homoskedastic Normal distribution. However, if we allow the variance to vary across observations, say:

$$y_i \sim f_{\mathcal{N}}(\mu_i, \sigma_i^2)$$

when we define both parameters as:

$$\mu_i = \mathbf{x}_i \beta, \quad \sigma_i^2 = \langle \neg | (\mathbf{x}_i \gamma) \rangle$$

this is now heteroskedastic Normal, which we cannot model with OLS assumptions. The MLE for the heteroskedastic Normal should be:

$$P\left(\mathbf{y}|\boldsymbol{\mu},\boldsymbol{\sigma}^{2}\right) = \prod_{i=1}^{n} f_{\mathcal{N}}\left(y_{i}|\mu_{i},\sigma_{i}^{2}\right) \qquad \text{[Joint probability]}$$

$$P\left(\mathbf{y}|\boldsymbol{\mu},\boldsymbol{\sigma}^{2}\right) = \prod_{i=1}^{n} \left(2\pi\sigma_{i}^{2}\right)^{-1/2} \exp\left[\frac{-\left(y_{i}-\mu_{i}\right)^{2}}{2\sigma_{i}^{2}}\right] \qquad \text{[Expressed in Normal distribution]}$$

$$\log \mathcal{L}\left(\boldsymbol{\beta},\boldsymbol{\sigma}^{2}|\mathbf{y}\right) \propto -\frac{1}{2} \sum_{i=1}^{n} \log \sigma_{i}^{2} - \frac{1}{2} \sum_{i=1}^{n} \frac{\left(y_{i}-\mathbf{x}_{i}\boldsymbol{\beta}\right)^{2}}{\sigma_{i}^{2}} \qquad \text{[Converted to log likelihood; simplify]}$$

$$\log \mathcal{L}(\boldsymbol{\beta},\boldsymbol{\gamma}|\mathbf{y}) \propto -\frac{1}{2} \sum_{i=1}^{n} \log \mathbf{z}_{i}\boldsymbol{\gamma} - \frac{1}{2} \sum_{i=1}^{n} \frac{\left(y_{i}-\mathbf{x}_{i}\boldsymbol{\beta}\right)^{2}}{\exp\left(\mathbf{x}_{i}\boldsymbol{\gamma}\right)} \qquad \text{[Substitute in systematic components]}$$

Let's try to simulate the heteroskedastic Normal data and fit with both the OLS and MLE.

$$y_i \sim f_{\mathcal{N}}(\mu_i, \sigma_i^2)$$

 $\mu_i = 0 + 5x_{1i} + 15x_{2i}$
 $\sigma_i^2 = \exp(1 + 0x_{1i} + 3x_{2i})$

Set up the variables

```
set.seed(2020)
n <- 1000

y <- x0 <- x1 <- x2 <- vector(mode="numeric", 1000)
x0 <- rep(1, n)
x1 <- runif(n, min=0, max=1)</pre>
```

```
x2 <- runif(n, min=0, max=1)

x <- cbind(x0,x1,x2)

beta <- c(0,5,15)
gamma <- c(1,0,3)</pre>
```

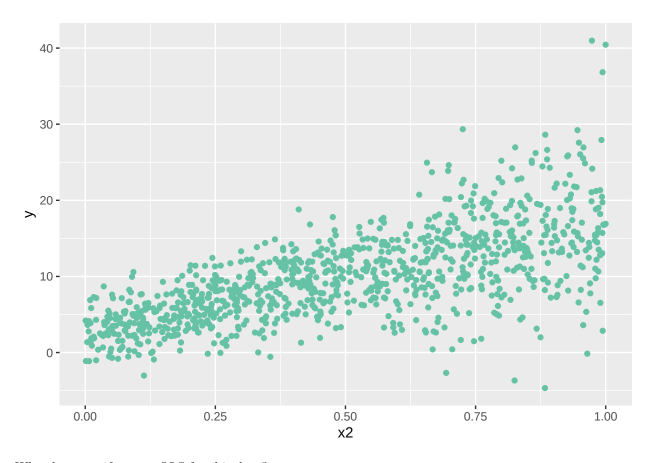
Create y variables according to our model

Let's plot the y_i over x_{2i} .

```
col <- RColorBrewer::brewer.pal(6, "Set2")

true <- ggplot(data, aes(x2, y)) +
   geom_point(color=col[1]) +
   theme(legend.position = "none")

true</pre>
```



What happens if we use OLS for this data?

```
lm <- lm(y ~ x1 + x2, data=data)
summary(lm)</pre>
```

```
##
## Call:
## lm(formula = y ~ x1 + x2, data = data)
##
## Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                           Max
## -21.3971 -2.2322 -0.1876
                               2.1940 25.9313
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
                                    0.444
## (Intercept)
                0.1633
                           0.3678
                                             0.657
## x1
                 4.8264
                            0.4626 10.433
                                             <2e-16 ***
## x2
                14.8285
                            0.4660 31.823
                                            <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4.199 on 997 degrees of freedom
## Multiple R-squared: 0.5157, Adjusted R-squared: 0.5147
## F-statistic: 530.7 on 2 and 997 DF, p-value: < 2.2e-16
```

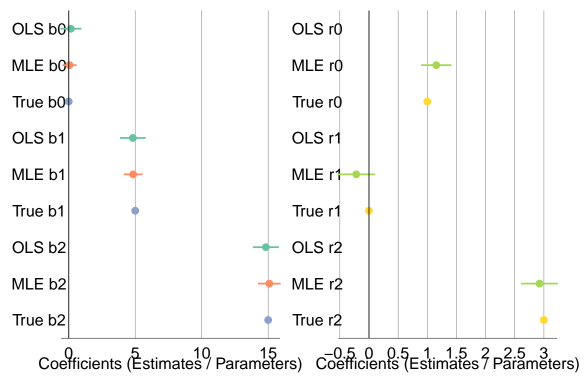
What if we use MLE?

```
llk.hetnormlin <- function(param, y, x, z) {</pre>
  x <- as.matrix(x)</pre>
                                        # x (some covariates) as a matrix
  z <- as.matrix(z)</pre>
                                       # z (some covariates) as a matrix
  os \leftarrow rep(1, nrow(x))
                                       # Set the intercept as 1 (constant)
  x <- cbind(os, x)
                                       # Add intercept to covariates x
 z <- cbind(os, z)</pre>
                                       # Add intercept to covariates z
 b <- param[1 : ncol(x)]
                                                  # Parameters for x
  g \leftarrow param[(ncol(x) + 1) : (ncol(x) + ncol(z))] # Parameters for z
 xb <- x %*% b
                                         # Systematic components for mean
 s2 \leftarrow exp(z \% *\% g)
                                         # Systematic components for variance
  sum(0.5 * (log(s2) + (y - xb)^2 / s2)) # Likelihood we want to maximize
                                         # optim is a minimizer by default
                                         \# To maximize lnL is to minimize -lnL
                                         # so the +/- signs are reversed
}
stval \leftarrow c(0, 0, 0, 0, 0, 0)
xcovariate <- cbind(x1,x2)</pre>
# Run ML, and get the output we need
hetnorm.result <- optim(stval,
                                       # Initial quesses
                        llk.hetnormlin, # Likelihood function
                        method = "BFGS", # Gradient method
                       hessian = TRUE, # Return Hessian matrix
                                       # Outcome variable
                       x = xcovariate, # Covariates x (w/o constant)
                        z = xcovariate # Covariates z (w/o constant)
                                    # Point estimates
pe <- hetnorm.result$par</pre>
round(pe, 3)
## [1] 0.059 4.846 15.086 1.154 -0.218 2.925
vc <- solve(hetnorm.result$hessian) # Var-cov matrix (for computing s.e.)
round(vc, 5)
                   [,2]
                              [,3]
            [,1]
                                       [, 4]
                                                [,5]
                                                         [,6]
## [1,] 0.05610 -0.06059 -0.05223 0.00087 -0.00068 -0.00107
## [2,] -0.06059 0.10771 0.01117 -0.00064 0.00098 0.00032
## [4,] 0.00087 -0.00064 -0.00184 0.01570 -0.01387 -0.01376
## [5,] -0.00068  0.00098  0.00055 -0.01387  0.02406  0.00406
## [6,] -0.00107  0.00032  0.00315 -0.01376  0.00406  0.02355
se <- sqrt(diag(vc))</pre>
                                    # To compute standard errors (s.e.)
                                    # take the diagonal of the Hessian;
                                    # then take square root
round(se, 3)
```

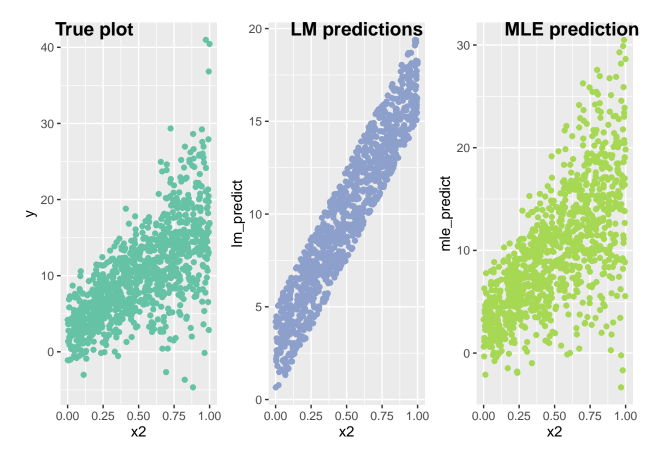
Comparison between MLE and OLS? Let's show with a ladderplot:

```
lm_coef <- c(summary(lm)$coefficients[,1], rep(NA,3))</pre>
lm_se <- c(summary(lm)$coefficients[,2], rep(NA,3))</pre>
mle coef <- round(pe,3)</pre>
mle se <- round(se,3)</pre>
true_coef <- c(0,5,15,1,0,3)
lm_lower <- lm_coef+lm_se*-1.96</pre>
mle_lower <- mle_coef+mle_se*-1.96</pre>
lm_upper <- lm_coef+lm_se*1.96</pre>
mle_upper <- mle_coef+mle_se*1.96</pre>
pe_all <- c(lm_coef,mle_coef,true_coef)</pre>
pe_all \leftarrow pe_all[c(1,7,13,2,8,14,3,9,15,4,10,16,5,11,17,6,12,18)]
lower all <- c(lm lower,mle lower,true coef)</pre>
lower_all <- lower_all[c(1,7,13,2,8,14,3,9,15,4,10,16,5,11,17,6,12,18)]
upper_all <- c(lm_upper,mle_upper,true_coef)</pre>
upper_all <- upper_all[c(1,7,13,2,8,14,3,9,15,4,10,16,5,11,17,6,12,18)]
result <- data.frame(label = c("OLS b0", "MLE b0", "True b0",
                                 "OLS b1", "MLE b1", "True b1",
                                 "OLS b2", "MLE b2", "True b2",
                                 "OLS r0", "MLE r0", "True r0",
                                 "OLS r1", "MLE r1", "True r1",
                                 "OLS r2", "MLE r2", "True r2"),
                        pe = pe_all, lower = lower_all, upper = upper_all)
B012 <- ropeladder(x=result[1:9,]$pe, lower=result[1:9,]$lower, upper=result[1:9,]$upper,
                      labels = c("OLS b0", "MLE b0", "True b0",
                                 "OLS b1", "MLE b1", "True b1",
                                 "OLS b2", "MLE b2", "True b2"),
                      size=0.65.
                      lex=1.75,
                      lineend="square", plot=1, col=col[c(1:3)])
R012 <- ropeladder(x=result[10:18,]$pe, lower=result[10:18,]$lower, upper=result[10:18,]$upper,
                      labels = c("OLS r0", "MLE r0", "True r0",
                                "OLS r1", "MLE r1", "True r1",
                                 "OLS r2", "MLE r2", "True r2"),
                      size=0.65.
                      lex=1.75,
                      lineend="square", plot=2, col=col[c(4:6)])
# Make reference line trace for first diffs (at 0)
line1 <- linesTile(x=c(0,0), y=c(0,1), plot=1)
line2 <- linesTile(x=c(0,0), y=c(0,1), plot=2)
# Set tick marks for x axis
```

Differences in coefs and 95% intervals



Finally, let's compare the prediction result between OLS and MLE:



GLM vs MLE logit

Technically, GLM fits a logit model with MLE methods. The basic difference between the linear and logit model is the existence of an error term:

$$y_i \sim f_{Bern}(\pi_i)$$

$$\pi_i = \text{AdJ} \approx^{-1} (X_i \beta) = \frac{e^{X_i \beta}}{1 + e^{X_i \beta}} = \frac{1}{1 + e^{-X_i \beta}}$$

The MLE for the logit model is as follows:

$$\begin{split} \mathcal{L}(\pi|\curvearrowright) &\propto \prod_{i=1}^n \pi_i^{y_i} (1-\pi_i)^{1-y_i} \\ \mathcal{L}(\beta|\curvearrowright) &\propto \prod_{i=1}^n (\frac{1}{1+\curvearrowleft(-\curvearrowright_i\beta)})^{y_i} (1-\frac{1}{1+\curvearrowleft(-\curvearrowright_i\beta)})^{1-y_i} \\ \mathcal{L}(\beta|\curvearrowright) &\propto \prod_{i=1}^n (1+\curvearrowleft(-\curvearrowright_i\beta))^{-y_i} (1+\curvearrowleft(\curvearrowright_i\beta))^{-(1-y_i)} \\ &\lessdot \forall \eth \mathcal{L}(\beta|\curvearrowright) &\propto \sum_{i=1}^n y_i \lessdot \forall \eth (1+\curvearrowleft(-\curvearrowright_i\beta)) - (1-y_i) \lessdot \forall \eth (1+\curvearrowleft(\curvearrowright\beta)) \end{split}$$

Let's just compare the results between GLM and MLE. Let's say we have a true model of a Bernoulli random variable which is:

$$\pi_i = < \times \eth \beth \approx^{-1} (0.5 + x_{1i} + 1.5 x_{2i}) = \frac{e^{X_i \beta}}{1 + e^{X_i \beta}} = \frac{1}{1 + e^{-X_i \beta}}$$

```
n <- 1000
x1 <- runif(n, 0, 1)
x2 <- runif(n, 0, 1)
y <- pi <- vector(mode="numeric", length=n)

# rbernoulli(1, 0.5)

for (i in 1:n) {
    xb <- 0.5 + x1[i] + 1.5*x2[i]
    pi[i] <- 1 / (1+exp(-xb))
    y[i] <- rbernoulli(1, pi[i])
}

sum(y)/length(y)</pre>
```

[1] 0.857

Let's fit the model with glm()

```
glm <- glm(y ~ x1 + x2, family="binomial")
summary(glm)</pre>
```

```
## x1
                 1.1873
                             0.3215
                                      3.693 0.000222 ***
## x2
                 2.1057
                             0.3450 6.103 1.04e-09 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 820.74 on 999 degrees of freedom
## Residual deviance: 766.10 on 997 degrees of freedom
## AIC: 772.1
## Number of Fisher Scoring iterations: 5
What about MLE?
llk.logit <- function(param,y,x) {</pre>
  os <- rep(1,length(x[,1]))
  x <- cbind(os,x)
  b <- param[ 1 : ncol(x) ]
  xb <- x\*\b
  sum(y*log(1+exp(-xb)) + (1-y)*log(1+exp(xb)));
               # optim is a minimizer, so min -ln L(param/y)
}
stval <- glm$coefficients
xcovariates <- cbind(x1, x2)
mle logit <- optim(stval,</pre>
                          llk.logit,
                          method="BFGS",
                          hessian=TRUE,
                          y=y,
                          x=xcovariates)
pe_2 <- mle_logit$par</pre>
se_2 <- sqrt(diag(solve(mle_logit$hessian)))</pre>
results_logit <- rbind(pe_2, se_2,
                       summary(glm)$coefficients[,1],
                       summary(glm)$coefficients[,2])
rownames(results_logit) <- c("Optim coefficients", "Optim SE", "GLM coefficients", "GLM SE")
results_logit %>% knitr::kable()
```

	(Intercept)	x1	x2
Optim coefficients	0.3179567	1.1873193	2.1057311
Optim SE	0.2161397	0.3215426	0.3450204
GLM coefficients	0.3179567	1.1873193	2.1057311
GLM SE	0.2161396	0.3215425	0.3450202

They are just same! What a surprise!