# Homework 2 / 02-13-2022

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#### library(tidyverse)

# Problem 1

Develop two Monte Carlo methods for the estimation of  $\theta = \int_0^1 e^{x^2} dx$  and implement in **R**.

## Answer:

### Method 1: simple Monte Carlo integration

$$\theta = \int_0^1 e^{x^2} dx = \int_0^1 e^{x^2} \cdot 1 dx = E\{g(U)\}, U \sim U(0, 1)$$

So we can get the estimation of  $\theta$  by estimating the  $E\{g(U)\}$ 

$$E\{g(U)\} \approx \frac{1}{n} \sum_{i=1}^{n} g(U_i) = \hat{\theta}$$

```
set.seed(2022)
N = 1e5
u \leftarrow runif(N)
theta_hat_1 = mean(exp(u^2))
```

From simple Monte Carlo integration, the estimated  $\hat{\theta}_1 = 1.4632808$ 

## Method 2: Importance Sampling

Suppose X is a r.v. with density f(x), such that f(x) > 0 on the set x : g(x) > 0. Let Y be r.v. g(X)/f(X). Then we can transform the integration to the following form:

$$\theta = \int_0^1 e^{x^2} dx = \int_0^1 \frac{e^{x^2}}{f(x)} \cdot f(x) dx = E[Y]$$

Then we can estimate the integration by estimating E[Y]:

$$E[Y] = \frac{1}{m} \sum_{i=1}^{m} Y_i = \frac{1}{m} \sum_{i=1}^{m} \frac{e^{X_i^2}}{f(X_i)}$$

Where  $X_i \stackrel{i.i.d}{\sim} f(x)$ , and in this case, I chose  $f(x) = \frac{e^{-x}}{1-e^{-1}}, 0 < x < 1$ , and using inverse CDF to sample from this distribution.

```
set.seed(2022)
# invCDF of fx
fx <- -log(1-u*(1-exp(-1)))
fg = exp(fx^2)/(exp(-fx)/(1-exp(-1)));
theta_hat_2 = mean(fg)
```

From Importance Sampling, the estimated  $\hat{\theta}_2 = 1.4630146$ 

# Problem 2

Show that in estimating  $\theta = E\{\sqrt{1-U^2}\}\$  it is better to use  $U^2$  rather than U as the control variate, where  $U \sim U(0,1)$ . To do this, use simulation to approximate the necessary covariances. In addition, implement your algorithms in  $\mathbf{R}$ .

#### Answer:

Since  $E[f_1(x)] = E[U^2] = \int_0^1 x^2 dx = \frac{1}{3}, E[f_2(x)] = E[U] = 0.5$ , so we can estimate  $\theta$  by  $E[g(x)] = E\{\sqrt{1-U^2}\} = E\{\sqrt{1-U^2} + c_1 \times (U-0.5)\} = E\{\sqrt{1-U^2} + c_2 \times (U^2 - \frac{1}{3})\}$ . And we choose

$$c_i = -\frac{Cov(g(X), f_i(X))}{Var(f_i(X))}, X \sim U(0, 1)$$

To get  $c_i$ , we need  $Cov(g(U), f_i(U))$  and  $Var(f_i(U))$ . In this case, these can be estimated from a preliminary Monte Carlo experiment.

```
set.seed(2022)
gfun <- function(x) sqrt(1-x^2)
# f_1_x
ffun_1 <- function(x) x^2
# f_2 x
ffun_2 <- function(x) x
# generating X from Unif(0,1)
n_2 = 1e6
u_2 = runif(n_2)
gx = gfun(u_2)
fx_1 = ffun_1(u_2)
fx_2 = ffun_2(u_2)
# Calculate cov and var
cov_gf1 = cov(gx,fx_1)
cov_gf2 = cov(gx,fx_2)
var_f1 = var(fx_1)
var_f2 = var(fx_2)
# compute the constant c_i's
c1 = -cov_gf1/var_f1
c2 = -cov_gf2/var_f2
# get the estimates of theta
T_c1 = gx+c1*(fx_1 - 1/3)
T_c2 = gx+c2*(fx_2 - 1/2)
# compare the estimated theta and their variance
theta_c1 = mean(T_c1)
```

```
theta_c2 = mean(T_c2)
var_reduce_1 = cov_gf1^2/var_f1
var_reduce_2 = cov_gf2^2/var_f2
imp_pct = (var(T_c2)-var(T_c1))/var(T_c2)
```

Since we set  $f_1(U) = U^2$ ,  $f_2(U) = U$ , and from Monte Carlo simulation, we got  $Cov(g(U), f_1(U)) = Cov(\sqrt{1-U^2}, U^2) = -0.065$ ,  $Cov(g(U), f_2(U)) = Cov(\sqrt{1-U^2}, U) = -0.059$ . The Var(g(U)) is reduced by  $\frac{[Cov(g(U), f_1(U))]^2}{Var(f_1(U))} = 0.048$  and  $\frac{[Cov(g(U), f_2(U))]^2}{Var(f_2(U))} = 0.042$ , so we can see that  $f_1(U) = U^2$  is a better control variate than  $f_2(U) = U$ , and the percent of improvement in variance reduction is 78.41 %. The estimated  $\theta_{ci}$  is  $\theta_{c1} = 0.785$  and  $\theta_{c2} = 0.786$ .

# Problem 3

Obtain a Monte Carlo estimate of

$$\int_{1}^{\infty} \frac{x^2}{\sqrt{2\pi}} e^{-\frac{x^2}{2}} dx$$

by importance sampling and evaluate its variance. Write a  ${f R}$  function to implement your procedure.

#### Answer:

Let  $g(x) = \frac{x^2}{\sqrt{2\pi}}e^{-\frac{x^2}{2}}dx$ , x > 1 I use the pdf of Exp(1), that is  $f(x) = e^{-x}$ , x > 1 \$ as my importance function. The integration can be written as  $\theta = \int_1^\infty \frac{x^2}{\sqrt{2\pi}}e^{-\frac{x^2}{2}}dx = \int_1^\infty \frac{g(x)}{f(x)}f(x)dx = E[\frac{g(x)}{f(x)}]$ , and we can estimate this integration by estimating  $E[\frac{g(x)}{f(x)}]$ .

```
set.seed(2022)

g <- function(x) {
    x ^ 2 / sqrt(2*pi) * exp(-x^2/2) * (x > 1)
}

f_imp <- function(x) {
    exp(-x)
}

r_imp <- function(N) {
    rexp(N)
}

# function of importance sampling, given g(x), f(x), and sampling function from f(x)
imp_sampling <- function(gfun, ffun, rffun, N = 1e7) {
    xs <- rffun(N)
    T_x <- gfun(xs)/ffun(xs)
    return(c(mean(T_x),var(T_x)))
}

imp_result <- imp_sampling(g,f_imp,r_imp,1e7)</pre>
```

The result shows that the estimated  $\hat{\theta} = 0.4006$ , and the  $Var(\hat{\theta}) = 0.3432$ . The true value of  $\theta$  can be derived by the following code:

```
integrate(g, 1, Inf)
```

## 0.400626 with absolute error < 5.7e-07

We can see that our Monte Carlo estimator is very close to the true parameter.

#### Problem 4:

Design an optimization algorithm to find the minimum of the continuously differential function

$$f(x) = -e^{-x}\sin(x)$$

on the closed interval [0, 1.5]. Write out your algorithm and implement it into R.

#### Answer:

I'd like to use Newton's method as the optimization algorithm. The Newton algorithm involves doing this iteratively:

$$\theta_i = \theta_{i-1} - \frac{f(\theta_{i-1})}{f'(\theta_{i-1})}$$

Until  $|f(\theta_i)|$  is sufficiently close to zero. Applying Newton's method to this problem, we got the local minimum when f'(x) = 0

$$f'(x) = e^{-x}\sin(x) - e^{-x}\cos(x) = e^{-x}(\sin(x) - \cos(x))$$
$$f''(x) = 2e^{-x}\cos(x)$$

so each time we update the  $x_i$  by the following equation:

$$x_i = x_{i-1} - \frac{e^{-x}(\sin(x) - \cos(x))}{2e^{-x}\cos(x)}$$

```
#R codes:
ffunc <- function(a){
    return(-exp(-a)*sin(a))
d_func <- function(a){</pre>
    return(exp(-a)*(sin(a)-cos(a)))
d2_func <- function(a){</pre>
    return(2*exp(-a)*cos(a))
}
i = 0
tol = 1e-10
cur <- start <- 0 # start from 0
resnewton <- c(i, cur, d_func(cur))
while(abs(d_func(cur)) > tol){
    i < -i+1
    renew <- cur - d_func(cur)/d2_func(cur)
    cur <- renew
    resnewton <- rbind(resnewton, c(i, cur, d_func(cur)))
}
min_x = resnewton[nrow(resnewton), 2]
```

From the Newton's method, we got the minimum of  $f(x) = -e^{-x}\sin(x)$  in [0, 1.5]. When x = 0.7853982, the minimum of  $f(x) = -e^{-x}\sin(x)$  in [0, 1.5] is -0.3223969.

### Problem 5:

The Poisson distribution, written as

$$P(Y = y) = \frac{\lambda^y e^{-\lambda}}{y!}$$

for  $\lambda > 0$ , is often used to model "count" data — e.g., the number of events in a given time period.

A Poisson regression model states that

$$Y_i \sim Poisson(\lambda_i),$$

where

$$\log \lambda_i = \alpha + \beta x_i$$

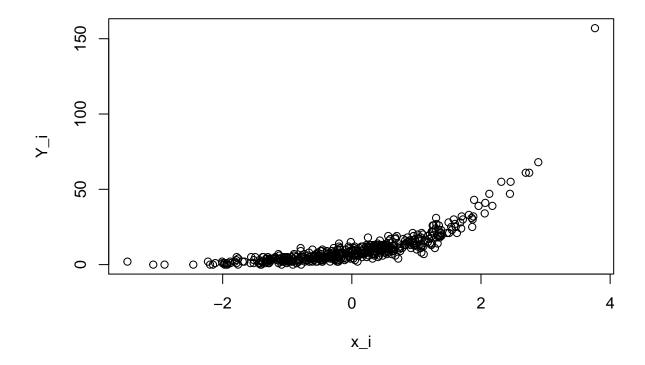
for some explanatory variable  $x_i$ . The question is how to estimate  $\alpha$  and  $\beta$  given a set of independent data  $(x_1, Y_1), (x_2, Y_2), \ldots, (x_n, Y_n)$ .

- 1. Generate a random sample  $(x_i, Y_i)$  with n = 500 from the Possion regression model above. You can choose the true parameters  $(\alpha, \beta)$  and the distribution of X.
- 2. Write out the likelihood of your simulated data, and its Gradient and Hessian functions.
- 3. Develop a modify Newton-Raphson algorithm that allows the step-halving and re-direction steps to ensure ascent directions and monotone-increasing properties.
- 4. Write down your algorithm and implement it in R to estimate  $\alpha$  and  $\beta$  from your simulated data.

#### Answer:

1. Generate a random sample  $(x_i, Y_i)$  with n = 500 from the Poisson regression model above.

Suppose the true parameters are  $\alpha = 2, \beta = 0.8$ . And  $x_i \sim N(0,1)$ . The generated data are displayed in the following plot.



2. Write out the likelihood of your simulated data, and its Gradient and Hessian functions.

Since  $Y_i \sim Pois(\lambda_i)$ . So the likelihood function of the data is:

$$L(\alpha, \beta | \{x_i, Y_i\}) = \prod_{i=1}^n \frac{e^{y_i(\alpha + \beta x_i)} e^{-e^{\alpha + \beta x_i}}}{y_i!}$$

Maximizing the likelihood is equivalent to maximizing the log-likelihood:

$$\ell(\alpha, \beta) = \sum_{i=1}^{n} \left( y_i(\alpha + \beta x_i) - e^{\alpha + \beta x_i} - \log y_i! \right)$$

So the gradient of this function is:

$$\nabla \ell(\alpha, \beta) = \begin{bmatrix} \frac{\partial \ell}{\partial \alpha} \\ \frac{\partial \ell}{\partial \beta} \end{bmatrix} = \begin{bmatrix} \sum_{i=1}^{n} (y_i - e^{\alpha + \beta x_i}) \\ \sum_{i=1}^{n} x_i (y_i - e^{\alpha + \beta x_i}) \end{bmatrix} = \begin{bmatrix} \sum_{i=1}^{n} (y_i - \lambda_i) \\ \sum_{i=1}^{n} x_i (y_i - \lambda_i) \end{bmatrix}$$

The Hessian is given by:

$$\nabla^2 \ell(\alpha, \beta) = - \left[ \begin{array}{c} \sum_{i=1}^n e^{\alpha + \beta x_i}, \sum_{i=1}^n x_i e^{\alpha + \beta x_i} \\ \sum_{i=1}^n x_i e^{\alpha + \beta x_i}, \sum_{i=1}^n x_i^2 e^{\alpha + \beta x_i} \end{array} \right] = - \left[ \begin{array}{c} \sum_{i=1}^n \lambda_i, \sum_{i=1}^n x_i \lambda \\ \sum_{i=1}^n x_i \lambda_i, \sum_{i=1}^n x_i^2 \lambda_i \end{array} \right]$$

3. Develop a modify Newton-Raphson algorithm that allows the step-halving and re-direction steps

To allow the step-halving to ensure monotone-increasing properties, we modified the Newton-Raphson at each step with a check-function:

$$\theta_i(\gamma) = \theta_{i-1} - \gamma [\nabla^2 \ell(\theta_{i-1})]^{-1} \nabla \ell(\theta_{i-1})$$

if  $f(\theta_i(1)) > f(\theta_{i-1})$ , then set  $\theta_i = \theta_{i-1}$ . Otherwise, search for a value  $\gamma \in (0,1)$  for which  $f(\theta_i(\gamma)) > f(\theta_{i-1})$  and set  $\theta_i = \theta_i(\gamma)$ . In step-halving, we search  $\gamma \in \{1/2, 1/4, 1/8, \ldots\}$ .

To allow re-direction steps and ensure we get ascent direction in each step, we have to make sure the Hessian matrix is negative definite. We can ensure that by changing the Newton-Raphson algorithm direction  $\mathbf{d} = -[\nabla^2 \ell(\theta_{\mathbf{i-1}})]^{-1} \nabla \ell(\theta_{\mathbf{i-1}})$  to  $\mathbf{d}' = -[\nabla^2 \ell(\theta_{\mathbf{i-1}}) - \eta \mathbf{I}]^{-1} \nabla \ell(\theta_{\mathbf{i-1}})$ , where  $\eta$  is large enough to make sure the modified Hessian is negative definite, that is, forced all eigenvalue of modified Hessian to less than zero. A simple choice of  $\eta$  is the largest positive eigenvalue of Hessian then plus one.

4. Write down your algorithm and implement it in R to estimate  $\alpha$  and  $\beta$  from your simulated data.

The code of the modified Newton-Raphson algorithm is in the following trunk.

```
# function giving gradient and Hessian
poissonstuff <- function(dat, betavec){</pre>
    u \leftarrow betavec[1] + betavec[2]*dat$x
    lambda \leftarrow exp(u)
    # loglik at betavec
    loglik \leftarrow sum(dat\$y * u - exp(u) - log(factorial(dat\$y)))
    # get the gradient at betavec
    grad \leftarrow c(sum(dat\$y - lambda), sum(dat\$x*(dat\$y - lambda)))
    # hessian matrix at betavec
    Hess <- -matrix(</pre>
        c(sum(lambda), sum(dat$x*lambda), sum(dat$x*lambda), sum((dat$x)^2*lambda)), ncol = 2
    )
    return(list(loglik = loglik, grad = grad, Hess = Hess))
}
ModifiedNewtonRaphson <- function(dat, func, start, tol = 1e-10, maxiter = 1000){
    i <- 0
    cur <- start
    stuff <- func(dat, cur)
    res \leftarrow (c(0, stuff$loglik, cur))
    prevloglik <- -Inf
    while (i < maxiter & abs(stuff\$loglik - prevloglik) > tol) {
        i < -i + 1
        prevloglik <- stuff$loglik</pre>
        prev <- cur
        Hess <- stuff$Hess
        e_hess <- eigen(Hess)$values
        # modified algorithm 1: re-direction
        # check if the hess is negative definite
        if (sum(e_hess) > 0){
             eta \leftarrow max(e hess) + 1
             Hess \leftarrow Hess - eta*diag(x = 1, dim(Hess))
        # modified algorithm 2: step-halving
        cur <- prev - solve(Hess) %*% stuff$qrad
        j <- 0
        while (func(dat, cur)$loglik < func(dat, prev)$loglik) {
```

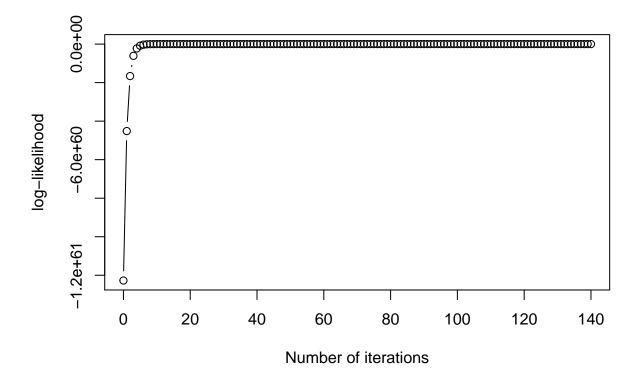
```
    j <- j + 1
        Gamma <- 0.5^(j)
        cur <- prev - Gamma * solve(Hess) %*% stuff$grad
}

# modification procedure ends
    stuff <- func(dat, cur)
    res <- rbind(res, c(i, stuff$loglik, cur))
}

return(res)
}</pre>
```

I choose a extreme combination of  $\{\alpha, \beta\}$  at the start value to test the robustness of my algorithm.

```
ans <- ModifiedNewtonRaphson(samples, poissonstuff, c(-47, -54))
ans <- data.frame(ans)
colnames(ans) <- c("iterations", "loglik", "alpha", "beta")
plot(ans$iterations, ans$loglik, type = "b", xlab = "Number of iterations", ylab = "log-likelihood")
```



```
alpha.hat <- ans[nrow(ans),]$alpha
beta.hat <- ans[nrow(ans),]$beta
```

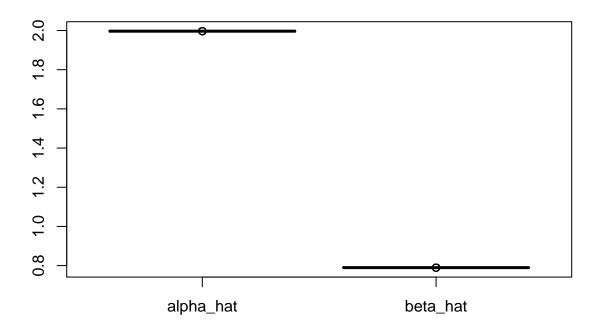
We can show the trace of the log-likelihood of the optimization procedure.

There are total 140 iterations, and the estimated parameters are  $\hat{\alpha} = 1.9966463$  and  $\hat{\beta} = 0.7897409$ . Remembering the true values are  $\alpha = 2, \beta = 0.8$ , which means that our algorithm has a high accuracy in estimating the parameters of the model.

```
i <- 1
alg_result <- c()</pre>
for (i in 1:100) {
    set.seed(i)
    i < -i + 1
    a \leftarrow rnorm(1, runif(1, 1, 2), 2) * 0.5 + 0.5
   b \leftarrow rpois(1, runif(1, 1, 2)) * 2 + 2
   startvec \leftarrow c(a,b)
    ans <- ModifiedNewtonRaphson(samples, poissonstuff, startvec)</pre>
    final <- ans[nrow(ans),]</pre>
    final <- append(final, startvec)</pre>
    alg_result <- rbind(alg_result, final)</pre>
}
alg_result <- data.frame(alg_result)</pre>
colnames(alg_result) <- c("number of iterations", "loglik", "alpha_hat", "beta_hat", "alpha_start", "be
alg_result %>% head() %>% knitr::kable(digit =4)
```

	$number\ of\ iterations$	log lik	$alpha\_hat$	$beta\_hat$	$alpha\_start$	$beta\_start$
final	14	-1171.789	1.9966	0.7897	0.8065	4
final.1	30	-1171.789	1.9966	0.7897	1.6237	8
final.2	15	-1171.789	1.9966	0.7897	1.9528	4
final.3	20	-1171.789	1.9966	0.7897	-1.0750	6
final.4	8	-1171.789	1.9966	0.7897	1.5824	2
final.5	24	-1171.789	1.9966	0.7897	2.8384	6

boxplot(alg\_result[,3:4])



I repeated the algorithm for 100 times from different starting values, all of them can reach a close estimate.