# CompStatLab2

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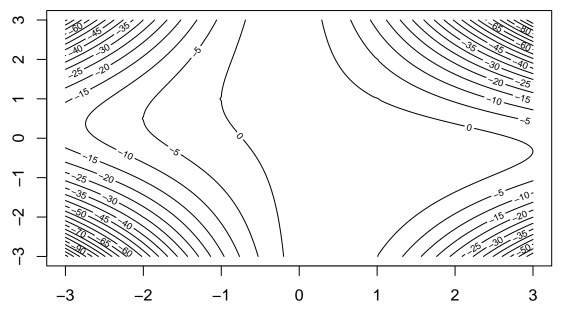
Computational Statistics 732A90 | Computer Lab 1

## Question 1: Optimisation of a two-dimensional function

Consider the function  $g(x,y) = -x^2 - x^2y^2 - 2xy + 2x + 2$  It is desired to determine the point (x, y), (x, y), (x, y), (x, y), (x, y), where the function is maximized.

a. Derive the gradient and the Hessian matrix in dependence of x, y. Produce a contour plot of the function g.

Answer: Please reference Appendix two\_dimensional.R, and below are plot of function g.



b. Write an own algorithm based on the Newton method in order to find a local maximum of g

## Answer:

Please reference Appendix two\_dimensional.R

c. Use different starting values: use the three points (x, y) = (2, 0), (-1, -2), (0, 1) and a fourth point of your choice. Describe what happens when you run your algorithm for each of those starting values. If your algorithm converges to points (x, y), compute the gradient and the Hessian matrix at these points and decide about local maximum, minimum, saddle point, or neither of it. Did you find a global maximum for x, y = [-3, 3]?

### Answer:

Please see below R results:

```
## 1) using starting points: (x,y) = (2,0), the newton method coverges to point:
##
## [1,] 1.0000256
## [2,] -0.9999341
## the corresponding gradient is:
## [1] 2.926312e-05 -8.049134e-05
## the corresponding hessian matrix is:
          [,1]
                   [,2]
## [1,] -3.999737 1.999839
## [2,] 1.999839 -2.000102
## This Hessian Matrix is negative definite.
## 2) using starting points: (x,y) = (-1,-2)
## [1,] 1.166791e-11
## [2,] 1.000000e+00
## the corresponding gradient is:
## [1] -2.939771e-08 -2.333581e-11
## the corresponding hessian matrix is:
      [,1]
## [1,] -4 -2.0000e+00
## [2,] -2 -2.7228e-22
## This Hessian Matrix is neither positive nor negative definite.
## 3) using starting points: (x,y) = (0,1)
     [,1]
## [1,]
## [2,]
         1
## the corresponding gradient is:
## [1] 0 0
## the corresponding hessian matrix is:
      [,1] [,2]
##
## [1,]
      -4 -2
## [2,]
      -2
## This Hessian Matrix is neither positive nor negative definite.
## 4) using starting points: (x,y) = (1,-1)
     [,1]
## [1,]
## [2,] -1
```

```
## the corresponding gradient is:
## [1] 0 0
## the corresponding hessian matrix is:
## [,1] [,2]
## [1,] -4 2
## [2,] 2 -2
## This Hessian Matrix is negative definite
```

## This Hessian Matrix is negative definite.

According to above R results for 4 different starting points, it's easily to get the conclusion:

- 1) when starting point is (2,0), the converaged point is (1.0000256,-0.9999341) which is local maximum point since the Hessian matrix at that point is negative definite.
- 2) when starting point is (-1,-2), the converaged point is (1.166791e-11,1.000000e+00) which is saddle point since the Hessian matrix at that point is neither positive or negative definite.
- 3) when starting point is (0,1), the converaged point is (0,1) which is saddle point since the Hessian matrix at that point is neither positive or negative definite.
- 4) when starting point is (1,-1), the converaged point is (1,-1) which is local maximum since the Hessian matrix at that point is negative definite.

d. What would be the advantages and disadvantages when you would run a steepest ascent algorithm instead of the Newton algorithm?

#### Answer:

## Question 2

a. Write a function for an ML-estimator for (0, 1) using the steepest ascent method with a step-size reducing line search (back-tracking). For this, you can use and modify the code for the steepest ascent example from the lecture. The function should count the number of function and gradient evaluations.

#### Answer:

b. Compute the ML-estimator with the function from a. for the data (xi, yi) above. Use a stopping criterion such that you can trust five digits of both parameter estimates for 0 and 1. Use the starting value (0, 1) = (-0.2, 1). The exact way to use backtracking can be varied. Try two variants and compare number of function and gradient evaluation done until convergence.

### Answer:

```
## $coefficients
## beta0 beta1
## -0.007299539 1.254971983
##
## $counts
## func_count gradien_count
## 14 21
```

c. Use now the function optim with both the BFGS and the Nelder-Mead algorithm. Do you obtain the same results compared with b.? Is there any difference in the precision of the result? Compare the number of function and gradient evaluations which are given in the standard output of optim.

#### Answer:

```
## $par
## [1] -0.009356126 1.262812832
```

```
##
## $value
## [1] -6.484279
##
## $counts
## function gradient
##
        12
##
## $convergence
## [1] 0
## $message
## NULL
## $par
## [1] -0.009423433 1.262738266
## $value
## [1] -6.484279
##
## $counts
## function gradient
##
         47
                  NA
##
## $convergence
## [1] 0
##
## $message
## NULL
d. Use the function glm in R to obtain an ML-solution and compare it with your results before.
Answer:
##
## Call:
## glm(formula = y \sim x, family = binomial(link = "logit"))
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -1.6780 -1.1734
                     0.7491
                                1.0251
                                         1.1814
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.00936
                            0.87086 -0.011
                                                0.991
## x
                1.26282
                            1.86663
                                      0.677
                                                0.499
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 13.460 on 9 degrees of freedom
## Residual deviance: 12.969 on 8 degrees of freedom
## AIC: 16.969
##
## Number of Fisher Scoring iterations: 4
```

## Appendix:

 $two\_dimensional.R$ 

```
# function g to be maximized
g <- function(x,y) {</pre>
     return(-x^2 - x^2 * y^2 - 2 * x * y + 2 * x + 2)
# partial derivative x for function g
deriv_x <- function(x, y) {</pre>
     return(-2 * x - 2 * x * y^2 - 2 * y + 2)
# partial derivative y for function g
deriv_y <- function(x, y) {</pre>
     return(-2 * x^2 * y - 2 * x)
# gradient for function g
gradient <- function (x, y) {</pre>
     return(c(deriv_x(x,y), deriv_y(x,y)))
# second partial derivative x for function g
deriv_x_11 <- function(x, y) {</pre>
     return(-2 - 2 * y^2)
\# second partial derivative x/y for function g
deriv_x_12 <- function(x, y) {</pre>
     return(-4 * x * y - 2)
}
\# second partial derivative y for function g
deriv_y_22 <- function(x, y) {</pre>
     return(-2 * x^2)
}
# hessian matrix for function g
hessian_g <- function(x, y) {
     return(matrix(c(deriv_x_11(x,y), deriv_x_12(x,y), deriv_x_12(x,y), deriv_y_22(x,y)), nrow = 2, ncol = 1, ncol = 1,
# produce a contour plot
xgrid \leftarrow seq(-3,3, by=0.05)
ygrid \leftarrow seq(-3,3, by=0.05)
length_x <- length(xgrid)</pre>
length_y <- length(ygrid)</pre>
dxy <- length_x * length_y</pre>
gxy <- matrix(rep(NA,dxy), nrow = length_x)</pre>
for ( i in 1:length_x) {
     for ( j in 1:length_y) {
           gxy[i, j] <- g(xgrid[i], ygrid[j])</pre>
```

```
mgxy <- matrix(gxy, nrow = length_x, ncol = length_y)</pre>
contour(xgrid, ygrid, mgxy, nlevels=40)
# newton function
newton_g \leftarrow function(x,y, eps=0.0001) {
 xt \leftarrow c(x,y)
 xt1 < -c(x,y) + 2
 while( t(xt1 - xt) %% (xt1 - xt) > eps) {
   xt1 <- xt
   xt <- xt1 - solve(hessian_g(xt[1] , xt[2])) %*% gradient(xt[1],xt[2])</pre>
 return(xt)
}
# check definiteness of a hessian matrix
check_definiteness <- function(matrix) {</pre>
 eigenvalues <- eigen(matrix)$values</pre>
 if (all(eigenvalues > 0)) {
   cat("This Hessian Matrix is positive definite.\n")
 } else if (all(eigenvalues < 0)) {</pre>
   cat("This Hessian Matrix is negative definite.\n")
 } else {
   cat("This Hessian Matrix is neither positive nor negative definite.\n")
 }
}
# using different starting points for newton method
converges_point1 <- newton_g(2,0)</pre>
cat("1) using starting points: (x,y) = (2,0), the newton method coverges to point: \n")
converges_point1
grad_point1 <- gradient(converges_point1[1],converges_point1[2])</pre>
cat("the corresponding gradient is: \n")
grad_point1
hessian_point1 <- hessian_g(converges_point1[1],converges_point1[2])
cat("the corresponding hessian matrix is: \n")
hessian_point1
check_definiteness(hessian_point1)
cat("-----")
converges_point2 <- newton_g(-1,-2)</pre>
cat("2) using starting points: (x,y) = (-1,-2) \n"
converges_point2
grad_point2 <- gradient(converges_point2[1],converges_point2[2])</pre>
cat("the corresponding gradient is: \n")
grad_point2
```

```
hessian_point2 <- hessian_g(converges_point2[1],converges_point2[2])</pre>
cat("the corresponding hessian matrix is: \n")
hessian point2
check_definiteness(hessian_point2)
cat("========"")
converges_point3 <- newton_g(0,1)</pre>
cat("3) using starting points: (x,y) = (0,1) \n"
converges_point3
grad_point3 <- gradient(converges_point3[1],converges_point3[2])</pre>
cat("the corresponding gradient is: \n")
grad_point3
hessian_point3 <- hessian_g(converges_point3[1],converges_point3[2])
cat("the corresponding hessian matrix is: \n")
hessian_point3
check definiteness(hessian point3)
cat("-----")
converges_point4 <- newton_g(1,-1)</pre>
cat("4) using starting points: (x,y) = (1,-1) \ n")
converges_point4
grad_point4 <- gradient(converges_point4[1],converges_point4[2])</pre>
cat("the corresponding gradient is: \n")
grad_point4
hessian_point4 <- hessian_g(converges_point4[1],converges_point4[2])</pre>
cat("the corresponding hessian matrix is: \n")
hessian_point4
check definiteness(hessian point4)
g <- function(x,y) {</pre>
ml\_estimator.R
# Define the log-likelihood function
log likelihood <- function(beta, x, y) {</pre>
 p \leftarrow 1 / (1 + exp(-beta[1] - beta[2]*x))
 sum(y * log(p) + (1 - y) * log(1 - p))
# Define the gradient of the log-likelihood function
gradient <- function(beta, x, y) {</pre>
 p <- 1 / (1 + exp(-beta[1] - beta[2]*x))</pre>
 db0 \leftarrow sum(y - p)
```

```
db1 \leftarrow sum((y - p) * x)
  c(db0, db1)
# Steepest ascent function
steepest_ascent <- function(beta_start, x, y, tol=1e-5, alpha0=1) {</pre>
  beta <- beta_start</pre>
  log_likelihood_count <- 0</pre>
  gradient_count <- 0</pre>
  conv <- 999
  while (conv > tol) {
    alpha <- alpha0
    beta old <- beta
    beta <- beta_old + alpha * gradient(beta_old, x, y)</pre>
    gradient_count <- gradient_count + 1</pre>
    while (log_likelihood(beta, x , y) < log_likelihood(beta_old, x , y)) {</pre>
      alpha <- alpha/2
      beta <- beta_old + alpha * gradient(beta_old, x , y)</pre>
      gradient_count <- gradient_count + 1</pre>
      log_likelihood_count <- log_likelihood_count + 2</pre>
    conv <- sum((beta - beta_old) * (beta - beta_old))</pre>
  result <- list(coefficients=c(beta0=beta[1],beta1=beta[2]), counts=c(func_count=log_likelihood_count,
  return(result)
}
# drug and placebo data
x \leftarrow c(0, 0, 0, 0.1, 0.1, 0.3, 0.3, 0.9, 0.9, 0.9)
y \leftarrow c(0, 0, 1, 0, 1, 1, 1, 0, 1, 1)
# Initial values for beta
beta_start \leftarrow c(-0.2, 1)
# Compute the ML estimator
ml_estimator_result <- steepest_ascent(beta_start, x, y)</pre>
# Print the results
print(ml_estimator_result)
# BFGS
result_bfgs <- optim(beta_start, log_likelihood, gradient, x=x, y=y, method="BFGS", control = list(fnsc
print(result_bfgs)
# Nelder-Mead
result_nelder <- optim(beta_start, log_likelihood, gradient, x=x, y=y, method="Nelder-Mead", control =
print(result_nelder)
# Fit the model
fit <- glm(y ~ x, family=binomial(link="logit"))</pre>
# Print the results
print(summary(fit))
```