DATA VISUALISATION

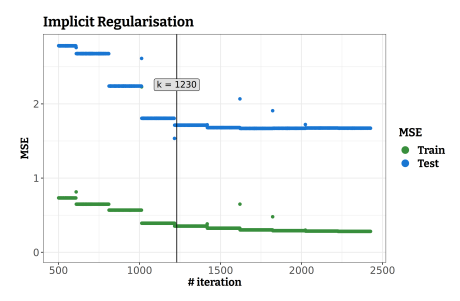
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Statement of Contribution

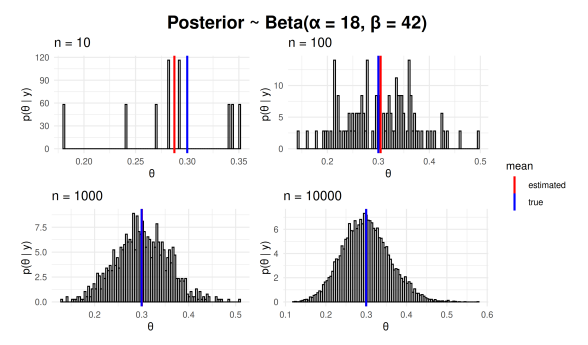
The plots for Convergence to the True Mean and MCMC Parameter Simulation for Poisson Regression were created with collaboration with Farid Musayev.

Implicit Regularisation



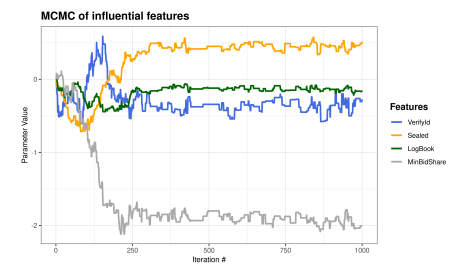
Interpretation: This plot shows how the mean squared error (MSE) changes with the number of iterations of gradient descent. The parameters are optimized implicitly. We see that the test MSE does not decrease significantly after 1230 iterations. Thus, the early stopping criterion can be used to avoid wasting computational resources.

Convergence to the True Mean



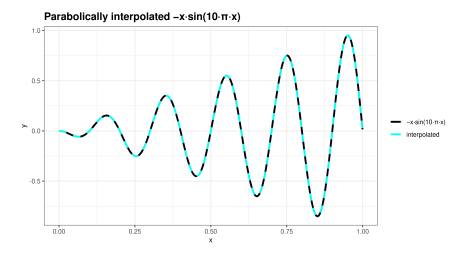
Interpretation: We sample data from the binomial distribution. With increasing number of sampled data points, the mean of the sampled data points converges to the true calculated mean of the binomial distribution. When n = 1000, it is already difficult to distinguish between the true and estimated means.

MCMC Parameter Simulation for Poisson Regression



Interpretation: We use the Metropolis Hastings algorithm to simulate the parameters of Poisson Regression. The plot above demonstrates how four parameters corresponding to four features converge after about 250 iterations.

Parabolic Approximation



Interpretation: In the plot above, one can observe how good is parabolic approximation using the gradient descent algorithm. The black line shows the real function, and the dashed cyan line shows the approximated values.

Appendix

Code for implicit regularisation

Code for convergence to the true mean

```
# Set seed
set.seed(12345)
# Given:
s = 13
n = 50
f = n - s
alpha 0 = 5
beta 0 = 5
# Beta Posterior distribution
beta posterior \leftarrow function(n, s = 13, f = 37){
  alpha 0 = 5
  beta 0 = 5
  rbeta(n = n, shape1 = alpha 0 + s, shape2 = beta 0 + f)
}
# True expected value and standard deviation
expected value \leftarrow (alpha 0 + s)/ ((alpha 0 + s) + (beta 0 + f))
variance <- (alpha 0 + s) * (beta 0 + f)/(((alpha 0 + s) + (beta 0 + f))**2*
                                               (alpha 0 + s + beta 0 + f + 1))
std <- variance**0.5</pre>
# Theta values generated from beta posterior for different n
```

```
df <- data.frame(theta 10 = beta posterior(n = 10),</pre>
                 theta 100 = \text{beta posterior}(n = 100),
                 theta 1000 = \text{beta posterior}(n = 1000),
                 theta 10000 = \text{beta posterior}(n = 10000))
# Plots for different n values
\# n = 10
pl1 <- ggplot(df) +
  geom histogram(aes(x = theta 10, y = ..density..., ),
                 fill = "lightgrey", bins = 100,
                 color = "black", alpha = 0.2) +
  geom vline(aes(xintercept = mean(theta 10), color = "estimated"),
             size = 1) +
  geom vline(aes(xintercept = expected value, color = "true"),
             size = 1) +
  scale color manual(name = "mean",
                     values = c(estimated = "red", true = "blue")) +
  labs(title = "n = 10", x = "\u03B8", y = "p(\u03B8 | y)") +
  theme minimal() + theme(legend.position = "none")
# n = 100
pl2 <- ggplot(df) +
  geom histogram(aes(x = theta 100, y = ..density..., ),
                 fill = "lightgrey", bins = 100,
                 color = "black", alpha = 0.2) +
  geom vline(aes(xintercept = mean(theta 100), color = "estimated"),
             size = 1) +
  geom vline(aes(xintercept = expected value, color = "true"),
             size = 1) +
  scale color manual(name = "mean",
                     values = c(estimated = "red", true = "blue")) +
  labs(title = "n = 100", x = "\u03B8", y = "p(\u03B8 | y)") +
  theme minimal() + theme(legend.position = "none")
# n = 1000
pl3 <- ggplot(df) +
  geom histogram(aes(x = theta 1000, y = ..density..., ),
                 fill = "lightgrey", bins = 100,
                 color = "black", alpha = 0.2) +
  geom vline(aes(xintercept = mean(theta 1000), color = "estimated"),
             size = 1) +
  geom vline(aes(xintercept = expected value, color = "true"),
             size = 1) +
  scale color manual(name = "mean",
                     values = c(estimated = "red", true = "blue")) +
  labs(title = "n = 1000", x = "\u03B8", y = "p(\u03B8 | y)") +
  theme minimal() + theme(legend.position = "none")
# n = 10000
pl4 \leftarrow qaplot(df) +
  geom histogram(aes(x = theta 10000, y = ..density..., ),
                 fill = "lightgrey", bins = 100,
```

```
color = "black", alpha = 0.2) +
  geom vline(aes(xintercept = mean(theta 10000), color = "estimated"),
             size = 1) +
  geom vline(aes(xintercept = expected value, color = "true"),
             size = 1) +
  scale color manual(name = "mean",
                     values = c(estimated = "red", true = "blue")) +
  labs(title = "n = 10000", x = "\u03B8", y = "p(\u03B8 | y)") +
  theme minimal()
# Extract the legend
g legend<-function(a.gplot){</pre>
  tmp <- ggplot gtable(qqplot build(a.qplot))</pre>
  leg <- which(sapply(tmp$grobs, function(x) x$name) == "guide-box")</pre>
  legend <- tmp$grobs[[leg]]</pre>
  return(legend)
}
shared legend <- g legend(pl4)
grid.arrange(arrangeGrob(pl1, pl2, pl3, pl4 + theme(legend.position = "none"),
                          ncol = 2),
             top = textGrob("Posterior ~ Beta(\u03B1 = 18, \u03B2 = 42)",
                             gp = gpar(fontsize = 18, fontface = "bold")),
             shared legend,
             heights = c(10, 0),
             widths = c(8.5, 1.5)
```

Code for MCMC parameter simulation for Poisson regression

```
# Metropolis Random Walk for Poisson regression
library(ggplot2)
library(mvtnorm)

data <- read.table("eBayNumberOfBidderData.dat", header = TRUE)
names(data)[1] <- "target"
X <- as.matrix(data[, -1])
y <- as.matrix(data[, 1])

Bayesian analysis of the Poisson regression

# Prior: beta ~ N(0, 100*inv((t(X)*X))), where X is the n x p covariate matrix
Sigma <- 100 * solve(t(X) %*% X)
mean = rep(0, 9)
betas_init = rep(0, 9)

# Posterior is assumed multivariate normal: N(beta_mode, inv(J_x(beta_mode))))
# LogPosterior:
logPost <- function(betas, mean, Sigma, X, y){</pre>
```

```
logPrior <- dmvnorm(x = betas, mean = mean, sigma = Sigma, log = TRUE)</pre>
  # unknown parameters in regression are betas.
  # This is why we write prior for them
 linPred <- X %*% betas</pre>
  logLik <- sum(linPred * y - exp(linPred)) #LogLik for the Poisson Model</pre>
  return(logPrior + logLik)
}
mode optim <- optim(betas init, logPost, gr=NULL, mean, Sigma, X, y,
                     method=c("BFGS"), control=list(fnscale=-1), hessian=TRUE)
# Name the coefficient by covariates
names(mode optim$par) <- names(as.data.frame(X))</pre>
# Compute approximate standard deviations.
approxPostStd <- sqrt(diag(-solve(mode optim$hessian)))</pre>
# Name the coefficient by covariates
names(approxPostStd) <- names(as.data.frame(X))</pre>
post mean = mode optim$par
print('The posterior mode is')
print(post mean)
print('The approximate posterior standard deviation is')
print(approxPostStd)
post cov mat <- -solve(mode optim$hessian)</pre>
print('The posterior covariance matrix is')
print(post cov mat)
N < -1000
random walk metropolis <- function(logPost, c = 1){
  theta \leftarrow matrix(NA, ncol = 9, nrow = N)
  theta[1, ] <- rep(0, 9)
  accept rate <- 1
  for (i in 2:N){
    proposed <- as.vector(rmvnorm(1, mean = theta[i-1, ],</pre>
                                    sigma=c*post cov mat))
    post prev <-logPost(betas = theta[i-1, ], mean, Sigma, X, y)
    post new <- logPost(betas = proposed, mean, Sigma, X, y)</pre>
    accept pr <- min(1, exp(post new - post prev))</pre>
    u <- runif(1)
    if (u <= accept pr){</pre>
      theta[i, ] <- proposed</pre>
      accept rate <- accept rate + 1
    else{
      theta[i, ] <- theta[i-1, ]</pre>
    }
  }
  print("Acceptance rate is")
  print(accept rate/N)
  return(theta)
}
```

```
theta <- random walk metropolis(logPost)</pre>
df plot <- data.frame(x = 1:N, VerifyId = theta[, 3], Sealed = theta[, 4],</pre>
                     Logbook = theta[, 8], MinBidShare = theta[, 9])
plot <- ggplot(df plot, aes(x = x)) +
 theme bw() + labs(title = "MCMC of influential features") +
 geom line(aes(y = VerifyId, color = "1"), size = 1) +
 geom_line(aes(y = Sealed, color = "2"), size = 1) +
 geom line(aes(y = Logbook, color = "3"), size = 1) +
 geom line(aes(y = MinBidShare, color = "4"), size = 1) +
 xlab("Iteration #") + ylab("Parameter Value") +
 "LogBook", "MinBidShare"),
                    name = "Features") +
 theme(title = element text(size = 12, face = 'bold'),
       axis.title = element text(size = 10, face = "plain"))
print(plot)
```

Code for parabolic approximation

```
library(ggplot2)
f <- function(n, fun, funName){</pre>
  #Required for later plot drawing
  x < -c()
  a0 < - c()
  a1 < - c()
  a2 < - c()
  #Interval partition
  lenInt <- 1/n</pre>
  largestPoint <- 0</pre>
  lowestPoint <- 0
  for (k in 1:n){
    #Theree points allocation
    largestPoint <- largestPoint + lenInt</pre>
    lowestPoint <- largestPoint - lenInt</pre>
    middlePoint <- (largestPoint + lowestPoint)/2</pre>
    #data to insert into optim
    data <- data.frame(x = c(lowestPoint, middlePoint, largestPoint),</pre>
                         v = c(fun(lowestPoint),
                                fun(middlePoint),
                                fun(largestPoint)))
    #print(data)
    optimized \leftarrow optim(fn = squaredError, par = c(0, 0, 0), data = data,
                         method = "BFGS") #The TA said to use BFGS
```

```
#these vectors are required for final plotting
    x <- c(x, middlePoint)</pre>
    a0 <- c(a0, optimized$par[1])</pre>
    a1 <- c(a1, optimized$par[2])</pre>
    a2 <- c(a2, optimized$par[3])</pre>
  }
  #The final data.frame for plotting
  dfPlot < -cbind.data.frame(x = x, a0 = a0, a1 = a1, a2 = a2)
  plot <- ggplot(dfPlot, aes(x = x)) +
    theme bw() +
    stat function(fun = fun, aes(color = "1"), n = 1000, size = 1.2) +
    stat function(fun = function(x) dfPlot$a0 + dfPlot$a1*x + dfPlot$a2*x^2,
                  aes(color = "2"), n = 1000,
                  size = 1.2,
                   linetype = "dashed") +
    scale color manual(name = NULL,
                       labels = c("\u2212x\u00B7sin(10\u00B7\u03C0\u00B7x)",
                                            "interpolated"),
                       values = c("black", "cyan")) +
    labs(title = paste0("Parabolically interpolated "
                         "\u2212x\u00B7sin(10\u00B7\u03C0\u00B7x)")) +
    xlab("x") +
    ylab("y") +
    theme(title = element text(size = 12, face = 'bold'),
          axis.title = element text(size = 10, face = "plain"))
  print(plot)
#To minimize
squaredError <- function(data, par) {</pre>
  MSE <- with(data, sum(par[1] + par[2]*x + par[3]*x*x - y)^2)
  #print(MSE)
  return(MSE)
}
f(n = 1000, fun = function(x))
  result <- -x*sin(10*pi*x)
  return(result)
}, funName = "-x*sin(10*pi*x)")
#As we can see, the interpolated function coincides with the real functions.
```