Chair of Statistics
Location: Berlin
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Lecture: Einführung in die Bayes-Statistik
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Program leave-one-out posterior predictive checking in R

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## 1 Introduction

### 2 Code

#### 2.1 General code

The Gibbs sampler

```
gibbsSampler <- function(X, Y, R, b = 0, initSigma = 1) {
  B \leftarrow R + b
  # Size of design matrix
  n \leftarrow nrow(X)
  p \leftarrow ncol(X)
  # Variables to store the samples in (initalize sigma with initSigma)
  betas <- matrix(nrow = B, ncol = p)</pre>
  sigma <- c(initSigma, rep(NA, B-1))</pre>
  # Sampling
  for(i in 1:B){
    # OLS of beta
    V \leftarrow solve(t(X)%*%X)
                              # (X^T X) ^-1
    beta_hat <- V%*%t(X)%*%Y # (X^T X)^-1 X^T Y
    # OLS of sigma
    sigma_hat \leftarrow t(Y-X%*\%beta_hat)%*%(Y-X%*\%beta_hat)/(n-p)
    # Sample beta from the full conditional
    betas[i,] <- rmvnorm(1, beta_hat, sigma[i]*V)</pre>
    # Sample sigma from the full conditional
    if(i < B) {
      sigma[i+1] <- 1/rgamma(1, (n-p)/2, (n-p)*sigma_hat/2)
  }
  # Remove burn in
  if(b != 0) {
   betas <- betas[-(1:b),]
    sigma \leftarrow sigma[-(1:b)]
  return(list(betas = betas, sigma = sigma))
```

Cross Validation function

```
crossValidation <- function(X, Y, R, b) {
    # Size of design matrix
    n <- nrow(X)
    p <- ncol(X)
    steps <- length(R)

# Run gibbs sampler to get sampled parameters
    samples <- lapply(1:n, function(i) gibbsSampler(X[-i,], Y[-i], R[steps], b))

# Initalize lists</pre>
```

```
Sigma <- list()
Betas <- list()
Yhati <- list()

# Calculate sigma, betas and Yhati for every step
for(k in 1:steps) {
   Sigma[[k]] <- sapply(samples, function(sample) mean(sample$sigma[1:R[k]]))
   Betas[[k]] <- sapply(samples, function(sample) colMeans(sample$betas[1:R[k],]))
   Yhati[[k]] <- sapply(1:n, function(i) X[i,]%*%Betas[[k]][,i])
}

return(list(Sigma = Sigma, Betas = Betas, Yhati = Yhati))
}</pre>
```

#### 2.2 Model evaluaion

```
bayesModelEvaluation <- function(models, Y, R, b) {</pre>
  # Evaluate multiple models and return results from all models
 n <- length(Y)</pre>
 k <- length(models)
  # Cross validate every model
  results <- lapply(1:k, function(i) crossValidation(models[[i]], Y, R, b))
  # Calculate Mean Squared Errors
  MSEs <- sapply(results, function(el) {
   return((1/(n-nrow(el$Betas[[1]])))*sum((Y-el$Yhati[[1]])^2))
  })
  # Plot MSEs
  par(mfrow = c(1,2))
  barplot(MSEs, xlab = "Models", ylab = "MSE", names.arg = seq(1,k),
          main = "Model evaluation using MSE")
  # Calculate log posterior predictive density (log likelihood)
  # y \sim N(XB, s^2(X^T X)^-1) = N(Yhat, s^2(X^T X)^-1)
  LPPDs <- sapply(results, function(eva) sum(log(dnorm(Y, eva$Yhati[[1]], eva$Sigma[[1]]))))
  # Plot LPPDs
  barplot(-2*LPPDs, xlab = "Models", ylab = "lppd", names.arg = seq(1,k),
          main = "Model evaluation using lppd")
 return(list(MSEs = MSEs, LPPDs = LPPDs))
```

Run model evaluation with data

```
# Swiss data
dat <- swiss

# Response variable
Y <- dat$Fertility
n <- nrow(dat)

# Design matrices

models <- list(</pre>
```

```
matrix(c(rep(1,n), dat$Education), nrow=n),
matrix(c(rep(1,n), dat$Agriculture), nrow=n),
matrix(c(rep(1,n), dat$Examination), nrow=n),
matrix(c(rep(1,n), dat$Catholic), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Agriculture), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Examination), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Catholic), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Agriculture, dat$Examination), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Agriculture, dat$Catholic), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Examination, dat$Catholic), nrow=n),
matrix(c(rep(1,n), dat$Education, dat$Agriculture, dat$Examination ,dat$Catholic), nrow=n)
)

### Model evaluation ###
criteria <- bayesModelEvaluation(models, Y, R = 50, b = 10) # R = 500, b = 100</pre>
```

#### Model evaluation using MSE

#### Model evaluation using lppd

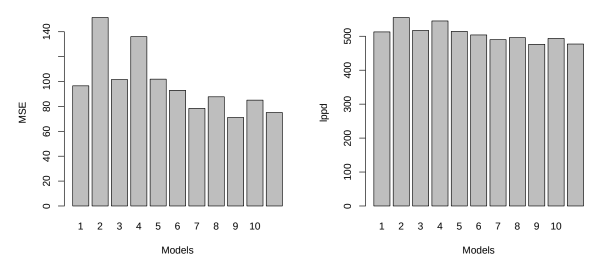


Figure 1: Model evaluation using MSE and lppd

```
# Check proportion
print(criteria$MSEs/(-2*criteria$LPPDs))

## [1] 0.1882963 0.2727617 0.1964855 0.2496199 0.1981878 0.1844312 0.1599963

## [8] 0.1769362 0.1492633 0.1723552 0.1573758

# TODO: Als Tabelle ausgeben? Explain why not fulfilled

# Choose optimal modal
optIdx <- which.min(-2*criteria$LPPDs)

# Check if MSE and LPPD would choose the same
print(paste(which.min(criteria$MSEs), "=", which.min(-2*criteria$LPPDs)))

## [1] "9 = 9"

# TODO den Vergleich einfach nur im Text verwenden?</pre>
```

#### 2.3 Cook's Distance

```
cooksDistance <- function(X, Y, R, b) {
# R is a vector</pre>
```

```
B \leftarrow R + b
  # Prepare projection matrix and number of parameters
  H \leftarrow X%*\%solve(t(X)%*\%X)%*\%t(X)
  p \leftarrow ncol(X)
  steps <- length(R)</pre>
  # Run cross validation with vector R
  cv <- crossValidation(X, Y, R, b)
  # Sample whole model (add R+b samples, remove b later)
  sample <- gibbsSampler(X, Y, R[steps] + b)</pre>
  # Cook's distance: Frequentist appraoch with analytic solution
  betaHat <- solve(t(X)%*%X)%*%t(X)%*%Y
  YhatLinReg <- X%*%betaHat
  E <- Y-YhatLinReg
  cooksLinReg <- (E^2/((1/(n-p))*sum(E^2)*p))*(diag(H)/(1-diag(H))^2)
  cooksBayesCV <- matrix(nrow = n, ncol = steps)</pre>
  for(k in 1:steps) {
    # Estimate posterior mean from betas
    betas <- colMeans(sample$betas[b:B[k],])</pre>
    # Predict values, using posterior mean
    Yhat <- X%*%betas
    # Calculate cook's distance
    dists <- apply(cv$Betas[[k]], 2, function(betas) {</pre>
      return((Yhat - X%*%betas)^2)
    })
    mse \leftarrow (1/(n-p))*sum((Y-Yhat)^2)
    cooksBayesCV[,k] <- colSums(dists)/(p*mse)</pre>
  return(list(cooksBayesCV = cooksBayesCV, cooksLinReg = cooksLinReg, sample = sample))
}
### Cook's Distance ###
# Number of samples
b <- 100
R \leftarrow seq(100,500,50) \# seq(100,5000,50)
# Get optimal model and calculate projection matrix
cooks <- cooksDistance(models[[optIdx]], Y, R, b)</pre>
Traces from bayes regression sampling
Posterior densities from bayes regression sampling
Apply cook's distance measure
```

```
# Initialize matrices
steps <- length(R)
matBayesD <- cooks$cooksBayesCV
matFreqD <- matrix(rep(cooks$cooksLinReg, steps), ncol=steps)
# Calculate differences between Cook's distance methods
cooksMeasure <- colSums((matBayesD-matFreqD)^2)</pre>
```

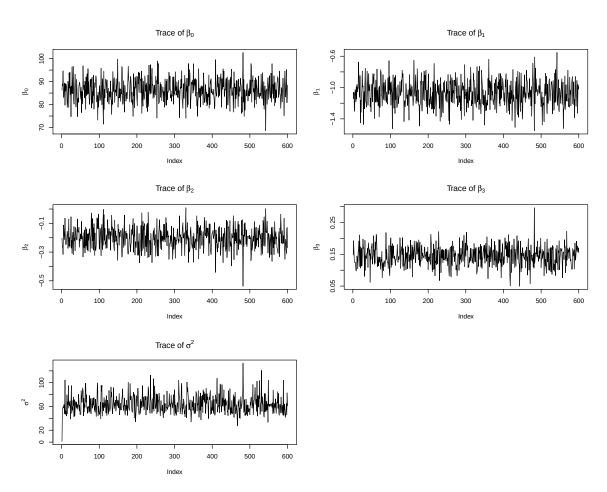


Figure 2: Traces

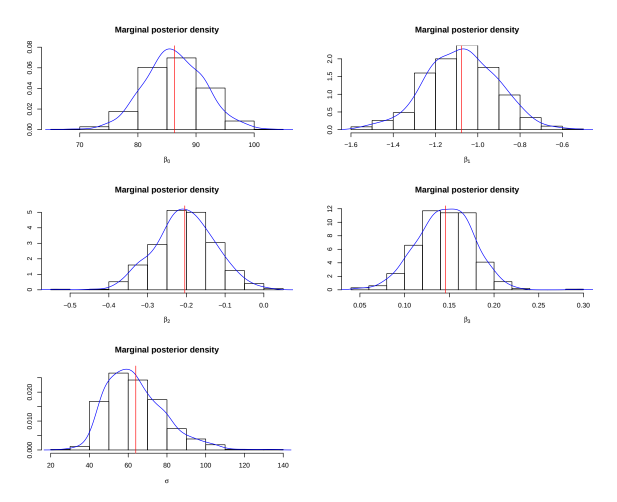


Figure 3: Densities

### Cook's distance measure

## Square distances: Cook's cross validation vs. Cook's analytic

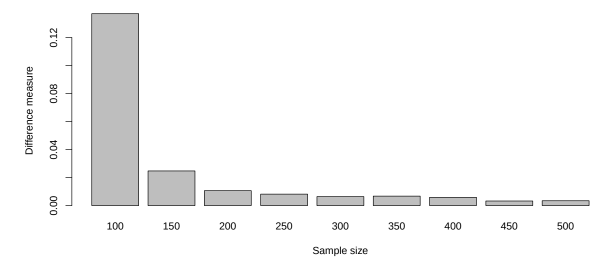


Figure 4: Cook's distance measure

Cook's distance, comarison between bayes and frequentist

## 3 References

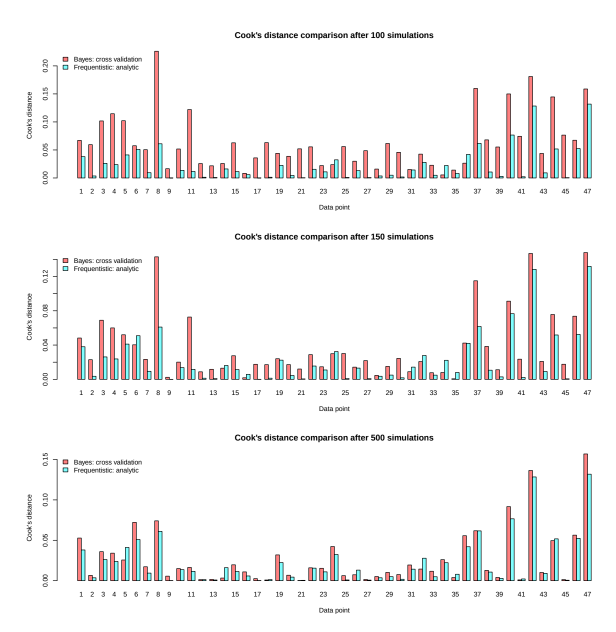


Figure 5: Pointwise cook's distance after three different simulation sizes, compared to analytic cook's distance from frequentist linear regression model using least squares.