MSiA400_Lab3

Question 1

 \mathbf{a}

b

Question 2

 \mathbf{a}

```
x_i <- c(4, 4, 5, 2, 2, 6)
p_fair <- rep(1/6, 6)
p_unfair <- c(2/13, 2/13, 1/13, 4/13, 2/13, 2/13)
p_fair_all <- 0.5 * p_fair[4]
p_unfair_all <- 0.5 * p_unfair[4]
for (i in seq(2, 6)){
   curr_xi <- x_i[i]
   p_fair_all <- p_fair_all * 0.75 * p_fair[curr_xi]
   p_unfair_all <- p_unfair_all * 0.75 * p_unfair[curr_xi]
}
p_fair_all</pre>
```

[1] 2.543132e-06

```
p_unfair_all
```

[1] 6.292977e-06

All unfair die is more likely to get the given observation.

b

```
s <- c("f")
p_fair_out <- c(0.5 * 1/6)
p_unfair_out <- c(0.5 * p_unfair[4])

for (i in seq(2,6)){
    curr_xi <- x_i[i]
# p(fair | x_i = curr_xi)
    curr <- c("f", "f", "u", "u")</pre>
```

```
curr_p_fair_prev_fair <- 0.75 * p_fair[curr_xi]</pre>
    curr_p_unfair_prev_fair <- 0.25 * p_unfair[curr_xi]</pre>
    curr_p_fair_prev_unfair <- 0.25 * p_fair[curr_xi]</pre>
    curr_p_unfair_prev_unfair <- 0.75 * p_unfair[curr_xi]</pre>
    p_fair_out <- append(p_fair_out, curr_p_fair_prev_fair)</pre>
    p_fair_out <- append(p_fair_out, curr_p_unfair_prev_fair)</pre>
    p_unfair_out <- append(p_unfair_out, curr_p_fair_prev_unfair)</pre>
    p_unfair_out <- append(p_unfair_out, curr_p_unfair_prev_unfair)</pre>
}
p_fair_out
## [1] 0.08333333 0.12500000 0.07692308 0.12500000 0.03846154 0.12500000
## [7] 0.03846154 0.12500000 0.03846154 0.12500000 0.03846154
p_unfair_out
## [1] 0.15384615 0.04166667 0.23076923 0.04166667 0.11538462 0.04166667
## [7] 0.11538462 0.04166667 0.11538462 0.04166667 0.11538462
Question 3
\mathbf{a}
dat <- read_csv("gradAdmit.csv")</pre>
## Rows: 400 Columns: 4
## Delimiter: ","
## dbl (4): admit, gre, gpa, rank
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
set.seed(123)
n <- nrow(dat)</pre>
train_i \leftarrow sample.int(n = n, size = floor(n * 0.8), replace = FALSE)
train <- dat[train_i, ]</pre>
```

test <- dat[-train_i,]</pre>

```
table(train$admit)
##
##
    0
## 216 104
table(test$admit)
##
## 0 1
## 57 23
In the train dataset, 32.5% were admitted. In the test dataset, 28.75% were admitted.
\mathbf{b}
best_m <- svm(factor(admit) ~ ., data = train,</pre>
                             kernel = "polynomial",
                             degree = 4,
                             gamma = 0.01,
                             coef0 = 10,
                             cost = 10)
pred_train <- predict(best_m, newdata = train, type = 'response')</pre>
pred_test <- predict(best_m, newdata = test, type = 'response')</pre>
confusionMatrix(pred_train, factor(train$admit), positive = "1")
## Confusion Matrix and Statistics
##
             Reference
##
## Prediction 0 1
            0 209 84
##
##
            1 7 20
##
##
                   Accuracy : 0.7156
##
                     95% CI: (0.6628, 0.7644)
##
       No Information Rate: 0.675
       P-Value [Acc > NIR] : 0.06664
##
##
##
                      Kappa: 0.1979
##
##
    Mcnemar's Test P-Value : 1.626e-15
##
##
               Sensitivity: 0.19231
##
               Specificity: 0.96759
##
            Pos Pred Value : 0.74074
            Neg Pred Value: 0.71331
##
##
                Prevalence: 0.32500
            Detection Rate: 0.06250
##
```

```
##
      Detection Prevalence: 0.08438
##
         Balanced Accuracy: 0.57995
##
##
          'Positive' Class : 1
confusionMatrix(pred_test, factor(test$admit), positive = "1")
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
##
            0 55 19
            1 2 4
##
##
##
                  Accuracy: 0.7375
##
                    95% CI : (0.6271, 0.8296)
##
       No Information Rate : 0.7125
       P-Value [Acc > NIR] : 0.3613344
##
##
##
                     Kappa : 0.1781
##
##
   Mcnemar's Test P-Value: 0.0004803
##
               Sensitivity: 0.1739
##
               Specificity: 0.9649
##
##
            Pos Pred Value: 0.6667
##
            Neg Pred Value: 0.7432
##
                Prevalence: 0.2875
##
            Detection Rate: 0.0500
##
      Detection Prevalence: 0.0750
##
         Balanced Accuracy: 0.5694
##
##
          'Positive' Class : 1
##
\# precision = TP / (TP + FP)
p_{test} \leftarrow 4 / (4 + 2)
\# recall = TP / (TP + FN)
r_{test} \leftarrow 4 / (4 + 19)
# specificity = TN / (TN + FP)
s_{test} < 55 / (55 + 2)
p_test
## [1] 0.6666667
r_test
```

[1] 0.173913

```
s_test
## [1] 0.9649123
There are 320 samples in the train set. There are 104 in admit, 216 in reject. Thus, we need 216 - 104 =
112 to obtain the most balanced dataset
perc <- 100 * (112-104) / 104
train$admit <- factor(train$admit)</pre>
train_df <- as.data.frame(train)</pre>
new_train <- SMOTE(form = admit~., train_df, perc.over = perc, perc.under = 0)</pre>
# table(new_train$admit)
train_full <- rbind(train_df, new_train)</pre>
table(train_full$admit)
##
##
    0
## 216 216
\mathbf{d}
best_m <- svm(factor(admit) ~ ., data = train_full,</pre>
                              kernel = "polynomial",
                              degree = 4,
                              gamma = 0.01,
                              coef0 = 10,
                              cost = 10)
pred_test <- predict(best_m, newdata = test, type = 'response')</pre>
confusionMatrix(pred_test, factor(test$admit), positive = "1")
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 40 11
##
##
            1 17 12
##
##
                   Accuracy: 0.65
                     95% CI: (0.5352, 0.7533)
##
##
       No Information Rate : 0.7125
##
       P-Value [Acc > NIR] : 0.9109
##
##
                      Kappa: 0.2074
##
##
    Mcnemar's Test P-Value: 0.3447
##
##
                Sensitivity: 0.5217
                Specificity: 0.7018
##
```

```
##
            Pos Pred Value: 0.4138
##
            Neg Pred Value: 0.7843
##
                Prevalence: 0.2875
##
            Detection Rate : 0.1500
##
      Detection Prevalence: 0.3625
         Balanced Accuracy: 0.6117
##
##
          'Positive' Class : 1
##
##
# precision = TP / (TP + FP)
p_test_smote <- 12 / (12 + 19)
\# recall = TP / (TP + FN)
r_test_smote <- 12 / (12 + 11)
\# specificity = TN / (TN + FP)
s_{test_{smote}} < 38 / (38 + 19)
p_test_smote
## [1] 0.3870968
r_test_smote
## [1] 0.5217391
s_test_smote
```

[1] 0.6666667

The precision and specificity are decreased, but the recall is increased compared to the unbalanced dataset.

Problem 4

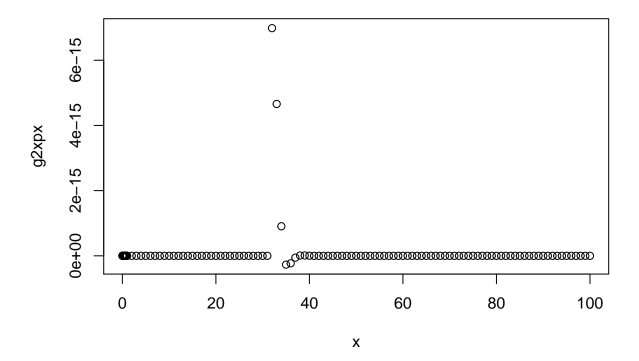
 \mathbf{a}

```
set.seed(123)
lambda <- 1
n <- ceiling((1 / lambda^2) / (10^(-6) * 0.01))
curr_i <- numeric(3)
real_i <- 1/(1+lambda^2)

curr_n <- n
x <- runif(curr_n, 0, 1)
y <- -log(x) / lambda
curr_i <- 1/curr_n * sum(sin(y) / lambda)
curr_i</pre>
```

[1] 0.5000296

```
crit \leftarrow exp(-10 * pi)
sum(y > crit) / length(y)
## [1] 1
\mathbf{b}
set.seed(123)
lambda <- 1
n \leftarrow ceiling((1 / lambda^2) / (10^(-6) * 0.01))
curr_i <- numeric(3)</pre>
real_i <- 1/(1+lambda^2)</pre>
curr_n <- n
x \leftarrow runif(curr_n, 0, 1)
y \leftarrow -log(x) / lambda
est <- 1/curr_n * sum(sin(y) / lambda)</pre>
est
## [1] 0.5000296
real <-1 / (1 + 1^2)
real
## [1] 0.5
\mathbf{c}
We can plot g^2(x)p(x) to find the best p^*(x)
x \leftarrow c(seq(0.01, 0.1, 0.01), seq(0.1, 1, 0.1), seq(1, 100))
g2xpx \leftarrow exp(-x) * ifelse(x >= 10*pi, sin(x), 0)
plot(x, g2xpx)
```



Here, we can see that when $g^2(x)p(x)$ is small, it is equal to 0. Note the asymptotic behavior of the curve. Thus, we must ensure that when $x < 10\pi$, $p^*(x) < p(x)$. By same logic, when $x > 10\pi$, $p^*(x) > p(x)$.

 \mathbf{d}

```
set.seed(123)
lambda <- 1
n <- ceiling((1 / lambda^2) / (10^(-6) * 0.01))
curr_i <- numeric(3)
real_i <- 1/(1+lambda^2)

curr_n <- n
x <- runif(curr_n, 0, 1)
y <- -log(x) / lambda
est <- 1/curr_n * sum(sin(y) / lambda)
est</pre>
```

```
## [1] 0.5000296
```

```
real <- 1 / (1 + 1^2)
real
```

```
## [1] 0.5
```