

Applied Data Mining Homework 2

Xun Zhao, xz2827

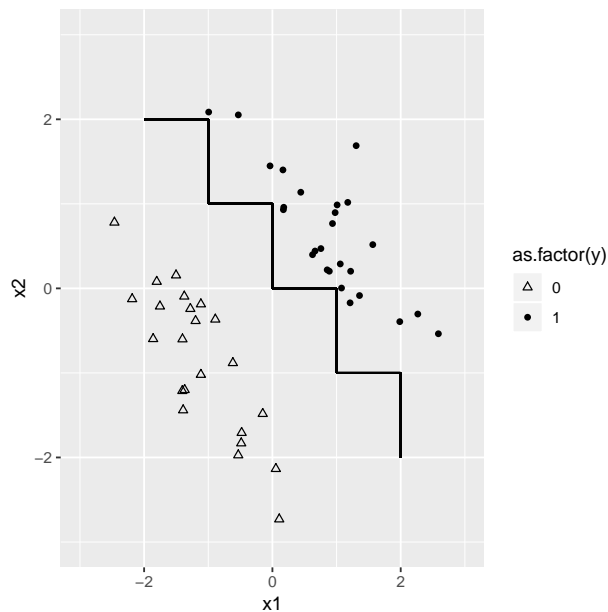
Problem 1: Trees

1.

Yes. But it needs the tree to be more complex.

As the tree classifier boundary is made of some line segments that are parallel with x-axis or y-axis, we can draw a zigzag curve along the linear boundary. Thus, the sloping linear boundary becomes many segments, and every segment is a decision in the tree classifier.

`## Warning: Removed 1 rows containing missing values (geom_point).`



2.

No, the Bayes-Optimal is defined as:

$$f(\vec{x}) = \underset{y}{\operatorname{argmax}} P(\mathbf{Y} = y | \mathbf{X} = \vec{x})$$

And the risk is defined as:

$$R(f) = \sum_{+1, -1} \int L(y, f(\vec{x})) P(\vec{x}, y) d\vec{x}$$

Thus, the risk $R(f)$ depends on the possible data distribution instead of only existing data points. Moreover, the Bayes-optimal classifier is defined by the possible distribution, so it can always minimize the risk under certain data model. However, the tree classifier cannot fit the linear boundary, which causes the misclassification compared with Bayes-optimal. As the Bayes-optimal has the lowest risk, the risk of tree classifier will always differ from Bayes-optimal, unless the Bayes-optimal's boundary is parallel to x or y-axis.

3.

$$g(\vec{x}) = \begin{cases} f_1(\vec{x}) > 0 & \begin{cases} f_2(\vec{x}) > 0 \rightarrow \text{class A} \\ f_2(\vec{x}) < 0 \rightarrow \text{class B} \end{cases} \\ f_1(\vec{x}) < 0 & \begin{cases} f_3(\vec{x}) > 0 \rightarrow \text{class B} \\ f_3(\vec{x}) < 0 \rightarrow \text{class C} \end{cases} \end{cases}$$

Problem 2: 10-fold Cross Validation

1.

10 folds means that we devide the data set χ in to 10 parts with the same size $\frac{||\chi||}{10}$.

$$\chi = \bigcup_{i=1}^{10} \chi_i$$

$$\text{if } i \neq j, \chi_i \cap \chi_j = \emptyset$$

$$||\chi_i|| = \frac{||\chi||}{10}$$

2.

For each new $i \in \{1, 2, \dots, 10\}$, namely, different cross validation set $\chi_{cv} = \chi_i$, we learn the model from training set

$$\chi_{\text{training}} = \chi - \chi_i = \bigcup_{j \neq i} \chi_j$$

Next, we apply the classification model f on $\chi_{cv} = \chi_i$ to calculate the error on cross validation set. Then, repeat this process for each i .

3.

As the risk function $R(k)$ given below, we compare the risk (for all folds and for all cross validation data points).

$$R(k) = \frac{1}{10} \sum_{i=1}^{10} \frac{1}{||\chi_i||} \sum_{j=1}^{||\chi_i||} l[y_{ij}, f(\vec{x}_{ij}; k, \chi - \chi_i)]$$

4.

We can simply choose k that minimize the risk $R(k)$.

$$k = \underset{k \in \{1,3,5,7,9\}}{\operatorname{argmin}} R(k)$$

5.

The most obvious disadvantage of k-NN algorithm is that it has to store all the training data, and iterate all the training data when classifying a new data point. Thus, it is time-consuming and occupy a lot of storage.

Problem 3: Cross validating a nearest neighbor classifier

1. Read data from files

```
pixels = as.matrix(read.table('../Data/uspsdata.txt'))
labels = as.matrix(read.table('../Data/uspscl.txt'))
nums = nrow(pixels)
```

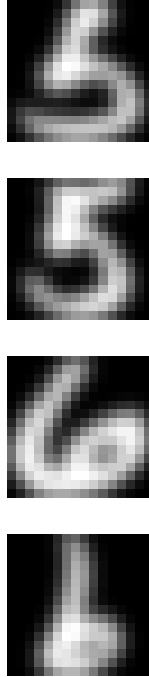
2. Plot first 4 digits

```
image.print <- function(x){
  x.matrix <- matrix(x, 16, 16, byrow = FALSE)
  x.matrix.rotated <- t(apply(x.matrix, 1, rev))
  image(x.matrix.rotated,
        axes = FALSE,
        col = grey(seq(0, 1, length.out = 256))
  )
}
```

```

}
for(i in 1:4){
  image.print(pixels[i,])
}

```



3. Devide data into 3 parts

```

# Data set is random already. So split without shuffling
p.tr = pixels[1:round(nums * 0.6),]
l.tr = labels[1:round(nums * 0.6),]
p.cv = pixels[(round(nums * 0.6) + 1):round(nums * 0.8),]
l.cv = labels[(round(nums * 0.6) + 1):round(nums * 0.8),]
p.ts = pixels[(round(nums * 0.8) + 1):nums,]
l.ts = labels[(round(nums * 0.8) + 1):nums,]

```

4. Train with $k = 1$

```

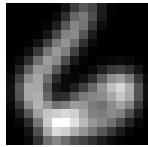
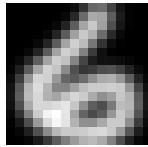
library(class)
estim = knn(p.tr, p.ts, cl = l.tr, k = 1)
err.rate = sum(estim != l.ts) / length(estim)

```

The test error is:

```
## [1] "err.rate = 0.05"
```

The misclassified digits are:



5. Optimize k

```
k.seq = seq(1, 13, 2)
err.rates = sapply(
  k.seq,
  function(k){
    estim = knn(p.tr, p.ts, cl = l.tr, k = k)
    sum(estim != l.ts) / length(estim)
  })
print(rbind(k.seq, err.rates))
```

```
##           [,1] [,2] [,3] [,4] [,5] [,6] [,7]
## k.seq      1.00 3.000    5 7.000 9.000 11.000 13.000
## err.rates 0.05 0.025    0 0.025 0.025 0.025 0.025
```

To choose the best hyperparameter k , we need to choose the k with least cross validation error:

```
## [1] "k.optim = 5, min(err.rates) = 0"
```

```
estim = knn(rbind(p.tr, p.ts), p.cv, cl = c(l.tr, l.ts), k = k.optim)
err.rate = sum(estim != l.cv) / length(estim)
```

Trained with training set and test set, the model's error rate on validation set is:

```
## [1] "err.rate = 0"
```