# MA8510: Introduction to Data Mining

Collaborate Session 2: Naive Bayes

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## Housekeeping

- Collaborate 1 = Wednesdays 6-7pm (Martha)
- Collaborate 2 = Thursdays 7-8pm (Hongbin)

For my Collaborate Sessions, you can get the **slides & R code** for each week here:

https://github.com/MarthaCooper/MA8510



## Subject: MA8510 Intro to Data Mining

#### MA8510 Learning Outcomes

- 1. Overview of Data Mining and Examples
- 2. Unsupervised data mining methods e.g. clustering and outlier detection;
- 3. Unsupervised and supervised techniques for dimensionality reduction;
- 4. Supervised data mining methods for pattern classification (Today = Naive Bayes);
- 5. Apply these concepts to real data sets using R (Today).

## Today's Goals

- Understand the background behind Naive Bayes Classifiers
- Apply Naive Bayes Classifiers to real datasets using R
- Understand the pros and cons of Naive Bayes Classifiers

## Naive Bayes Classifier

- Supervised, probabilistic classifier based on Bayes Theorem
- Strong independence assumptions
- A famous use case is spam filtering of emails
- Factor variable Multinomial Naive Bayes Classifier
- Continuous variables Gaussian Naive Bayes Classifier
- Given these features, does this sample belong to class X or Y?

#### Data

- We are going to use Class, Sex and Age data to predict Survival on the Titanic
- We first need to get the data and do a bit of wrangling

```
data(Titanic)
str(Titanic)
#function to convert to long format (one passenger per row)
counts to cases <- function(x){
  #how many time to repeat each row
  inds <- rep.int(seq len(nrow(x)), times = x[["Freq"]])</pre>
  #remove frequency column
  x <- select(x, -Freq)
  #get rows from x
  x[inds,]
case titanic <- counts to cases(as.data.frame(Titanic))</pre>
#head(case titanic, 3)
#dim(case titanic)
```

#### Split into test and training

- We will train our Naive Bayes Classifier on 80% of the data
- We can use the caret package to split the data

```
library(caret, warn.conflicts = F, quietly = T)
set.seed(1234)
split <- createDataPartition(case_titanic$Survived, p = 0.8, list = FAL$
train <- case_titanic[split, ]
test <- case_titanic[-split, ]
c(nrow(train), nrow(test)) # print number of observations in test vs. to
table(train$Survived) %>% prop.table() # Proportions of Survived Yes and
```

#### Naive Bayes: An Overview

- Bayes Theorem, by Reverend Thomas Bayes, is about conditional probability; the probability of A given that B occurred [deonoted as P(A|B)]
- In our Titanic example, the probability of a passenger having survived  $S_k$  (where  $S_{yes}$  = survived and  $S_{no}$  = did not survive), given that it's predictor values are  $x_1, x_2 \dots x_p$  [denoted as  $P(S_k | x_1, \dots x_p)$ ].

#### Naive Bayes: An Overview

Bayes Theorem looks like this:

$$P(S_k|X) = rac{P(X|S_k) \cdot P(S_k)}{P(X)}$$

#### where:

- $P(S_k)$  The prior probability of the outcome based on the training data, what is the probability of a person surviving or not? Our prior probability of survival is ~32% and the probability of not surviving is ~68%.
- P(X) The probability of observing the predictor variables.
- $P(X|S_k)$  The conditional probability or likelihood, For each class (i.e. survived and no survived), what is the probability of observing the predictors.
- $P(S_k|X)$  The posterior probability. We update our prior probabilities with our observed information to find the posterior probability that an observation has class  $S_k$ .

#### Naive Bayes: An Overview

$$P(S_{yes}|X) < P(S_{no}|X)$$

$$P(S_{yes}|X) = P(S_{no}|X)$$

$$P(S_{yes}|X) > P(S_{no}|X)$$

- Because we are comparing between classes, we don't need the denominator P(X) as it will be the same in both classes.
- ullet So what we actually end up calculating is proportional (not equal to) to  $P(S_k|X)$

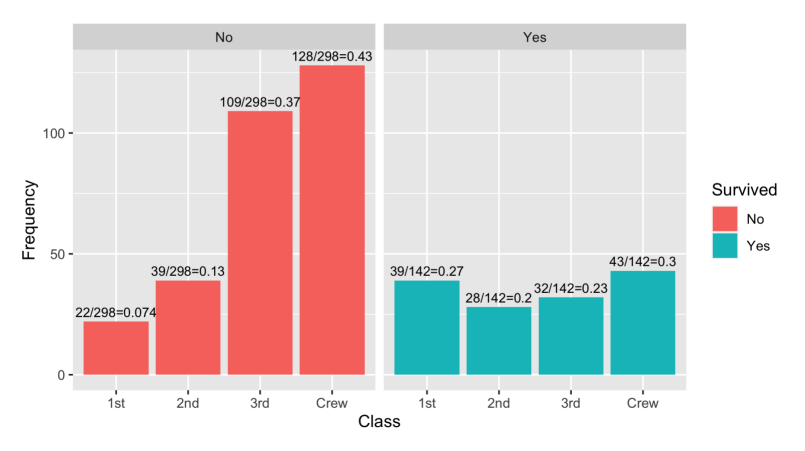
### Naive Bayes: Why Naive?

- Naive Bayes makes the assumption that the predictor variables are all conditionally independent of each other
- So  $P(S_k|X)$  is proportional to  $P(S_k)$  times the product of conditional probabilities of all predictors in the class.

$$P(S_k|X) \propto P(S_k) \cdot \prod_{i=1}^n P(x_i|S_k)$$

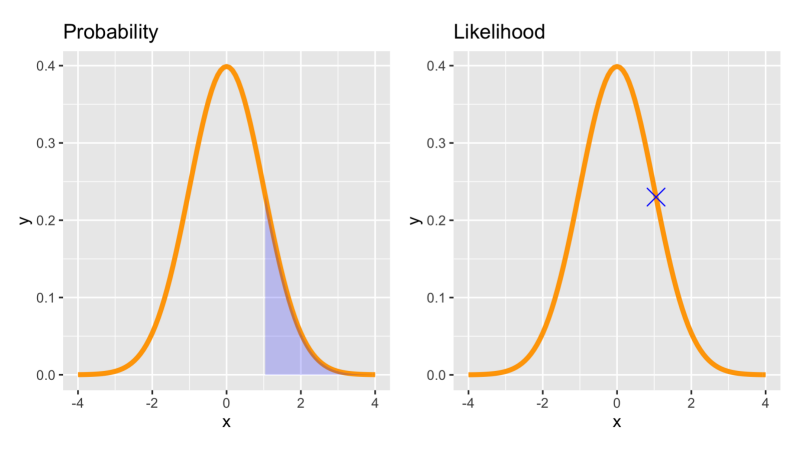
## Calculating Likelihoods

- Factor Variables use frequencies from the training data to calculate probabilities of the each predictor in each class (like our Titanic data).
- For factor variables, probabilities = likelihoods.



## Calculating Likelihoods

- Continuous Variables use a probability density function (PDF), e.g. Gaussian, to calculate the likelihood of each predictor in a class.
- n.b. For continuous variable, probabilities and likelihoods are different



## Naive Bayes Classifier with factor variables - Titanic Survival

There are lots of R packages to apply naive bayes (e1071, klaR, naivebayes etc...) but I like caret - common framework for many algorithms.

```
predictors <- names(train)[1:3] #Create response and predictor data
x <- train[,predictors] #predictors
y <- train$Survived #response</pre>
train control <- trainControl(method = "cv", number = 10) #set up 10 fol
survival mod1 <- train( #train the model
 X = X
 y = y,
 method = "nb",
  trControl = train control
confusionMatrix(survival mod1) #results of model on training data
pred <- predict(survival mod1, newdata = test) #make predictions</pre>
confusionMatrix(pred, test$Survived) #assessing the classifier
```

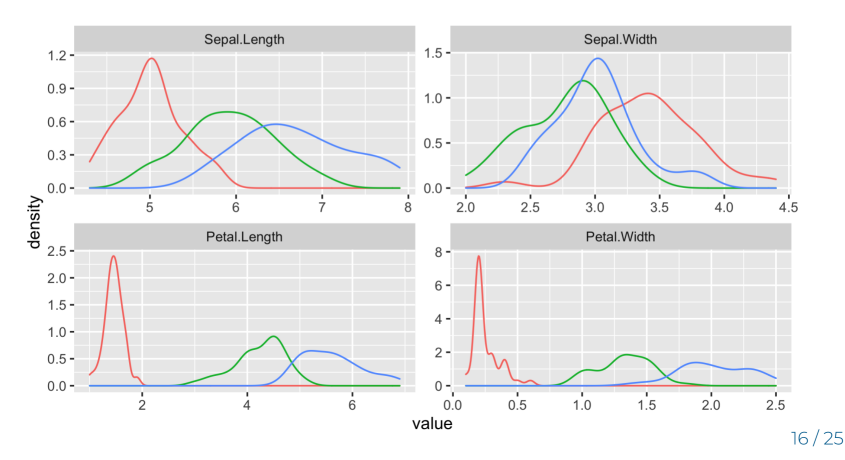
• Predict Species based on sepal length, sepal width, petal length and petal width.

```
set.seed(1234)
split <- createDataPartition(iris$Species, p = 0.8, list = FALSE)

train <- iris[split, ]
test <- iris[-split, ]

c(nrow(train), nrow(test)) # print number of observations in test vs. train(train$Species) %>% prop.table() # Proportions of Species
```

- Checking assumption of gaussian distribution
- Are data normally distributed? Do we need to transform? Or should we use non-parametric kernel density distribution instead? (Actually we will try both by tuning the model)



• First, fit a model with 10 fold cross validation

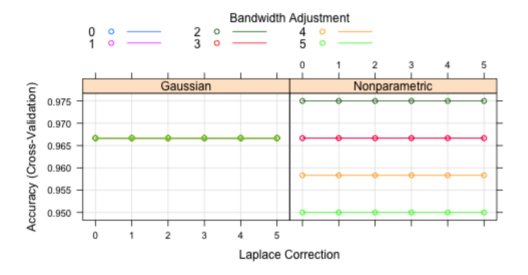
```
predictors <- names(train)[1:4] #Create response and predictor data
x <- train[,predictors] #predictors</pre>
y <- train$Species #response
train control <- trainControl(method = "cv", number = 10) #set up 10 fo.
species mod1 <- train( #train the model
 X = X
  y = y,
 method = "nb",
  trControl = train control
confusionMatrix(species_mod1) #results of model on training data
```

- Can we do any better?
- Tuning parameters = usekernal, adjust, fL

```
tune_params <- expand.grid( #define tuning parameters
  usekernel = c(TRUE, FALSE),
  fL = 0:5,
  adjust = 0:5
)

# train model
species_mod2 <- train(
  x = x,
  y = y,
  method = "nb",
  trControl = train_control,
  tuneGrid = tune_params
)</pre>
```

plot(species\_mod2)



- Laplace smoother makes no difference
- Non parametric Kernel with Bandwidth = 2 is best

Results for best model

### Overcoming randomness of splitting data

• Split data into test and training 10 times, and perform tuning & cross validation within each one

### Overcoming randomness of splitting data

• Split data into test and training 10 times, and perform tuning & cross validation within each one

```
all mods <- lapply(seq_len(ncol(splits)), function(i){</pre>
  train <- iris[splits[,i], ] #split into i'th test and train</pre>
  test <- iris[-splits[,i], ]
  predictors <- names(train)[1:4] #Create response and predictor data
  x <- train[,predictors] #predictors</pre>
  y <- train$Species #response
  species mod <- train( #fit models with tuning and cv
      X = X
      y = y,
      method = "nb",
      trControl = train control,
      tuneGrid = tune_params
  pred <- predict(species mod, newdata = test) #make predictions</pre>
  confusionMatrix(pred, test$Species)$overall["Accuracy"] #assessing the
})
```

## Overcoming randomness of splitting data

• Split data into test and training 10 times, and perform tuning & cross validation within each one

```
mean(unlist(all_mods))
sd(unlist(all_mods))
```

### Naive Bayes Pros and Cons

#### Pros

- Simple
- Fast
- Scales well

#### Cons

- Assumes independence of variables
- Assumes all variables are equally important
- Not as accurate as other methods e.g. Random Forests

## Extra reading/listening

#### Confused? Go here:

• A great conceptual overview StatQuest

#### Feeling confident? Try this:

- Use the caret *preProc* argument in **train()** to see if preprocessing the iris dataset improves model accuracy
- Try using the h2o package instead

#### Just for fun:

Listen to this Data Skeptic podcast

#### Important!

 Accuracy is not the best or only test for algorithm performance, explore more!

#### References

- UC Business Analytics Tutorial
- Zhang, 2016