Applied Machine Learning - Classification

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Outline

- Performance Measures
- OkC Data
- Classification Trees
- More Bagging
- Naive Bayes Models
Load Packages

```r
library(tidymodels)

## --- Attaching packages

## --- Conflicts

## * purrr::discard() masks scales::discard()
## * dplyr::filter() masks stats::filter()
## * dplyr::lag()    masks stats::lag()
## * dplyr::select() masks MASS::select()
## * recipes::step() masks stats::step()

transp <-
  element_rect(fill = "transparent", colour = NA)

thm <- theme_bw() +
  theme(
    panel.background = transp,
    plot.background = transp,
    legend.background = transp,
    legend.key = transp,
    legend.position = "top"
  )

theme_set(thm)
```
Measuring Performance in Classification
**Illustrative Example**

`yardstick` contains another test set example in a data frame called `two_class_example`:

```r
two_class_example %>% head(4)
```

<table>
<thead>
<tr>
<th>#</th>
<th>truth</th>
<th>Class1</th>
<th>Class2</th>
<th>predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Class2</td>
<td>0.00359</td>
<td>0.996</td>
<td>Class2</td>
</tr>
<tr>
<td>2</td>
<td>Class1</td>
<td>0.67862</td>
<td>0.321</td>
<td>Class1</td>
</tr>
<tr>
<td>3</td>
<td>Class2</td>
<td>0.11089</td>
<td>0.889</td>
<td>Class2</td>
</tr>
<tr>
<td>4</td>
<td>Class1</td>
<td>0.73516</td>
<td>0.265</td>
<td>Class1</td>
</tr>
</tbody>
</table>
```

Both `truth` and `predicted` are factors with the same levels. The other two columns represent *class probabilities*.

This reflects that most classification models can generate "hard" and "soft" predictions for models.

The class predictions are usually created by thresholding some numeric output of the model (e.g. a class probability) or by choosing the largest value.
Class Prediction Metrics

With class predictions, a common summary method is to produce a *confusion matrix* which is a simple cross-tabulation between the observed and predicted classes:

```r
two_class_example %>%
  conf_mat(truth = truth, estimate = predicted)
```

<table>
<thead>
<tr>
<th></th>
<th>Truth</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prediction</td>
<td>Class1</td>
</tr>
<tr>
<td>Class1</td>
<td>227</td>
<td>50</td>
</tr>
<tr>
<td>Class2</td>
<td>31</td>
<td>192</td>
</tr>
</tbody>
</table>

These can be visualized using *mosaic plots*.

Accuracy is the most obvious metric for characterizing the performance of models.

```r
two_class_example %>%
  accuracy(truth = truth, estimate = predicted)
```

```r
# A tibble: 1 x 3
## .metric .estimator .estimate
## <chr> <chr>      <dbl>
## binary          0.838
```

However, it suffers when there is a *class imbalance*; suppose 95% of the data have a specific class. 95% accuracy can be achieved by predicting samples to be the majority class.
Two Classes

There are a number of specialized metrics that can be used when there are two classes. Usually, one of these classes can be considered the event of interest or the positive class.

One common way to think about performance is to consider false negatives and false positives.

- The sensitivity is the true positive rate (out of all of the actual positives, how many did you get right?).
- The specificity is the rate of correctly predicted negatives, or 1 - false positive rate (out of all the actual negatives, how many did you get right?).

From this, assuming that class1 is the event of interest:

<table>
<thead>
<tr>
<th>##</th>
<th>Truth</th>
</tr>
</thead>
<tbody>
<tr>
<td>## Prediction</td>
<td>Class1</td>
</tr>
<tr>
<td>Class1</td>
<td>227</td>
</tr>
<tr>
<td>Class2</td>
<td>31</td>
</tr>
</tbody>
</table>

sensitivity = $\frac{227}{(227 + 31)} = 0.88$

specificity = $\frac{192}{(192 + 50)} = 0.79$
Conditional and Unconditional Measures

Sensitivity and specificity can be computed from \texttt{sens()} and \texttt{spec()}, respectively.

It should be noted that these are \textit{conditional measures} since we need to know the true outcome.

The event rate is the \textit{prevalence} (or the Bayesian \textit{prior}). Sensitivity and specificity are analogous to the \textit{likelihood values}.

There are \textit{unconditional} analogs to the \textit{posterior values} called the positive predictive values and the negative predicted values.

A variety of other measures are available for two class systems, especially for \textit{information retrieval}.

One thing to consider: what happens if our \textbf{threshold to call a sample an event is not optimal}?
Changing the Probability Threshold

For two classes, the 50% cutoff is customary; if the probability of class #1 is $\geq$ 50%, they would be labelled as class1.

What happens when you change the cutoff?

- Increasing it makes it harder to be called class1 $\Rightarrow$ fewer predicted events, specificity $\uparrow$, sensitivity $\downarrow$
- Decreasing the cutoff makes it easier to be called class1 $\Rightarrow$ more predicted events, specificity $\downarrow$, sensitivity $\uparrow$

With two classes, the Receiver Operating Characteristic (ROC) curve can be used to estimate performance using a combination of sensitivity and specificity.

To create the curve, many alternative cutoffs are evaluated.

For each cutoff, we calculate the sensitivity and specificity.

The ROC curve plots the sensitivity (eg. true positive rate) versus 1 - specificity (eg. the false positive rate).

The area under the ROC curve is a common metric of performance.
The Receiver Operating Characteristic (ROC) Curve

```r
roc_obj <-
  two_class_example %>%
  roc_curve(truth, Class1)

two_class_example %>% roc_auc(truth, Class1)

## # A tibble: 1 x 3
## #  .metric .estimator .estimate
## #  <chr>   <chr>          <dbl>
## 1 roc_auc binary          0.939

autoplot(roc_obj) + thm
```
Changing the Threshold

![Graph showing changing thresholds with sensitivity and specificity values]
The Receiver Operating Characteristic (ROC) Curve

The ROC curve has some major advantages:

- It can allow models to be optimized for performance before a definitive cutoff is determined.
- It is *robust* to class imbalances; no matter the event rate, it does a good job at characterizing model performance.
- The ROC curve can be used to pick an optimal cutoff based on the trade-offs between the types of errors that can occur.

When there are two classes, it is advisable to focus on the area under the ROC curve instead of sensitivity and specificity.

Once an acceptable model is determined, a proper cutoff can be determined.
OkC Data

These data contains several types of fields:

- a number of open text essays related to interests and personal descriptions
- single choice type fields, such as profession, diet, gender, body type, etc.
- multiple choice data, including languages spoken, etc.

We will try to predict whether someone has a profession in the STEM fields (science, technology, engineering, and math) using a random sample of the overall dataset.

The data are included in the workshop's GitHub repo:
Dealing with Class Imbalances

In our data, only 18.2% of the profiles are in STEM professions. This complicates the analysis since many models will overfit to the majority class.

There are two main strategies to deal with this:

- *Cost-sensitive learning* where a higher cost is attached to the minority classes. In this way, the fitting process puts more emphasis on those samples.

- *Sampling procedures* that modify the rows of the data to re-balance the training set.

Cost-sensitive models tend to only produce hard classifications so we will focus on the latter.
Class Imbalance Sampling

There are a variety of methods for subsampling the data. Some exclude or replicate rows in the training set while others try to *synthesize* new data points to balance the classes.

The simplest method for dealing with the problem is to *down-sample* the data to make the number of STEM and non-STEM profiles the same.

While it seems like throwing away most of the data is a bad idea, it tends to produce less pathological distributions of the class probabilities and *might* improve the ROC curve.

It is critical that:

- the sampling should be done *inside of resampling*. Otherwise, the performance estimates can be optimistic.
- these sampling methods take place on the analysis set and not the assessment set.

Note that for a simple logistic regression model, this mainly has the effect of changing the intercept.
Calibration Effect (Test Set Example)
Calibration Effect (Test Set Example)

- **Cut = 0.50 (Sp = 0.691, Sn = 0.679)**
- **Down-Sampled**
- **Cut = 0.50 (Sp = 0.953, Sn = 0.223)**
- **No Sampling**
Resampling and Analysis Strategy

If we down-sample the data, the analysis set will consist of 1458 profiles (equally balanced). We'll again use 10-fold CV to resample the data.

Within each resample, the analysis data are down-sampled and the assessment sets are left alone.

The number of STEM profiles held-out would be about 72 and this should be sufficient to compute sensitivity.

The models will be optimized on the area under the ROC curve.

```r
library(caret)
ctrl <- trainControl(
  method = "cv",
  # Also predict the probabilities
classProbs = TRUE,
  # Compute the ROC AUC as well as the sens and spec from the default 50% cutoff. The
  # function `twoClassSummary` will produce those.
summaryFunction = twoClassSummary,
savePredictions = "final",
sampling = "down"
)
```
Classification Trees
Classification Trees

Tree-based classifiers conduct searches of the predictors to find the best split of the data to create two subsets.

"Best", in most cases, means that the class distribution is as pure as possible in the subsets (i.e. is mostly one class).

There are many different types of classification trees, including conditional inference trees, C5.0, Bayesian additive regression trees, globally optimal trees, CART, and others.

We'll focus on CART via the rpart package for these notes.
Two Possible Splits - Which One is Better?

The data have been down-sampled so that classes have equal frequencies.
A Better Split

The two sets were derived by the model and are not listed here due to their sizes.

"Set 1" includes

## [1] "college_university"
## [2] "dropped_out_of_college_university"
## [3] "dropped_out_of_high_school"
## [4] "dropped_out_of_ph_d_program"
## [5] "graduated_from_college_university"
## [6] "graduated_from_masters_program"
## [7] "graduated_from_ph_d_program"
## [8] "graduated_from_space_camp"
## [9] "masters_program"
## [10] "working_on_space_camp"
The Official First Split
The Next Two splits

Once the initial split is made, the model will then split the resulting two data sets using new searches in those leaves.

The process continues until there are not enough data points left to accurately split or a predefined split limit has been reached.

This is the tree growing process and, once complete, most tree-based models begin to prune the trees using some method that balances complexity with performance.
Classification Trees

caret contains multiple method for training CART models.

We'll go with method = "rpart2" which uses maxdepth as the tuning parameter.

Also, we'll use the non-formula interface so that the factor predictors are not converted to indicator variables.

```r
set.seed(5515)
cart_mod <- train(
    x = okc_train[, names(okc_train) != "Class"],
    y = okc_train$Class,
    method = "rpart2",
    metric = "ROC",
    tuneGrid = data.frame(maxdepth = 1:20),
    trControl = ctrl
)
```
Note that there are 10 terminal nodes and thus 10 possible probability values.
CART Resampling Profile

ROC (Cross-Validation)

Max Tree Depth
Classification Tree Average ROC Curve

ROC curves will be created for each model so a convenience function will be used repeatedly to create an approximate curve:

```r
approx_roc_curve <- function(x, label) {
  x %>%
    pluck("pred") %>%
    roc_curve(obs, stem) %>%
    mutate(model = label)
}
approx_roc_curve(cart_mod, "CART") %>%
ggplot(aes(x = 1 - specificity, y = sensitivity)) +
  geom_path() +
  geom_abline(col = "red", alpha = .5)
```
Resampled Confusion Matrix

Since there are 10 different assessment sets, separate confusion matrices can be constructed for each.

train() has its own confusionMatrix() method that can show aggregations of these matrices.

By default, the overall rates are shown in each cell.

```r
confusionMatrix(cart_mod)
```

## Cross-Validated (10 fold) Confusion Matrix

## (entries are percentual average cell counts across runs)

<table>
<thead>
<tr>
<th></th>
<th>Reference</th>
<th>Prediction</th>
<th>stem</th>
<th>other</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>13.2</td>
<td>30.1</td>
<td>13.2</td>
<td>30.1</td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>51.6</td>
<td>5.0</td>
<td>51.6</td>
</tr>
</tbody>
</table>

## Accuracy (average) : 0.6487
Variable Importance Scores

CART tracks the decrease in impurity in the nodes when splits are made.

These can be aggregated over the tree to produce another general importance score.

rpart can also keep track of splits not used in the model (e.g. surrogate, competing) that can be used when there are missing predictor values. The \texttt{varImp()} method has options to turn these on/off.

Turning these off gives only the splits uses in the official tree.
Hands-On: I Don't Want to be a Dummy!

Take the previous code and use the formula method to create the model. For `train()`, the formula method always creates dummy variables for predictors that are factors.

Is there any difference in performance?

Is the final model affected? How?

Take 15 min to answer these questions.
Bagging (Again)
Instability of Trees

Many types of trees are *unstable* in that they have high variance; if the data are slightly changed, a large impact can be seen on the structure of the model.

This is generally bad and might make you question the interpretability of trees.

For example, what happens if we were to build our model on bootstrap samples of the training set?

In the three following plots, the maximum tree depth was capped at 5 to make the trees easier to visualize.
Bootstrap Sample #1

```
vn=abcdefghjklmnorwACEFGJKOPQR
- stem
- other

male>=0.5

education=cdhknpr
- stem
- awesome>=0.5
  - stem
  - other
```

```
technology>=0.5
```
Bootstrap Sample #3
Lemonade from Lemons

The *nice* thing about this instability is that it makes trees great candidates for ensemble methods.

Since ensembles use multiple models, they are only effective when the constituent models are *diverse*; otherwise the same predictions are averaged.

Let's bag the CART trees by using bootstrap samples of the training set and growing the largest possible tree.

caret wraps ipred::bagging() using method = "treebag". We will use the default ensemble size of 25 models.
Bagging CART Trees

```r
set.seed(5515)
cart_bag <- train(
  x = okc_train[, names(okc_train) != "Class"],
  y = okc_train$Class,
  method = "treebag",
  metric = "ROC",
  trControl = ctrl
)
```
Bagged CART Results

cart_bag

## Bagged CART
## 4000 samples
## 90 predictor
## 2 classes: 'stem', 'other'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 3600, 3601, 3599, 3600, 3600, 3600, ...
## Additional sampling using down-sampling
##
## Resampling results:
##
## ROC  Sens  Spec
## 0.746  0.708  0.67
Resampled Confusion Matrix

As measured by the default cutoffs, there is some increase in accuracy that is achieved by improving the specificity (27% versus CART's 30.1%).

```
confusionMatrix(cart_bag)

## Cross-Validated (10 fold) Confusion Matrix
##
## (entries are percentual average cell counts across resamples)
##
##             Reference
## Prediction stem other
##    stem  12.9  27.0
##    other  5.3  54.8
##
## Accuracy (average): 0.677
```
Bagged Classification Tree Average ROC Curve

Although the bagged model's curve is uniformly better than the single tree, their performance is very similar for the cutoffs closest to upper left-hand corner.
When bagging, the CART importances scores for each tree are aggregated across trees.

Many more predictors are used in this model.
Hands-On: How Many Trees?

As previously mentioned, caret uses `ipred::bagging()` to create the model.

Look at the help function to determine which `bagging()` argument controls the number of bootstraps.

How can we make `train()` use a different value? (hint: `?train`)

Does changing this affect the area under the ROC curve?

Take another 10 mins.
Other Ensemble Methods

There are a variety of other methods for creating ensembles

- **Random forests** are just like bagging but the trees are made more diverse by randomly sampling a subset of predictors to be used in each split. (*ranger*)

- **Boosting** fits a *sequence* of trees and modifies the case-weights of each data point to increase diversity. (*xgboost*, *C50*)

- **Rotation forests** is PCA signal extraction on a random subset of the data prior to creating the trees. (*rotationForest*)

- Regression **committees** adjust the outcome data over a sequence of models. (*Cubist*)

- **Stacking** is an ensemble method where different types of models can be blended together through averaging. (*caretEnsemble*)

R has multiple implementations of these methods.
Bayes' Rule
Naive Bayes Models

This classification model is motivated directly from statistical theory based on Bayes' Rule:

\[
Pr[Class|Predictors] = \frac{Pr[Class] \times Pr[Predictors|Class]}{Pr[Predictors]} = \frac{Prior \times Likelihood}{Evidence}
\]

In English:

Given our predictor data, what is the probability of each class?

The prior is the prevalence that was mentioned earlier (e.g. the rate of STEM profiles). This can be estimated or set.

Most of the action is in \( Pr[Predictors|Class] \), which is based on the observed training set.

Predictions are based on a blend of the training data and our prior belief about the outcome...
So Why is it Naive?

Determining $Pr[\text{Predictors}|\text{Class}]$ can be very difficult without strong assumptions because it measures the joint probability of all of the predictors.

- For example, what is the correlation between a person's essay length and their religion?

To resolve this, naive Bayes assumes that all of the predictors are independent and that their probabilities can be estimated separately.

The joint probability is then the product of all of the individual probabilities (an example follows soon).

This assumption is almost certainly bogus but the model tends to do well despite this.
The Effect of Independence

The probability contours assume multivariate normality with different assumptions.

Suppose the red dot is a new sample.

Probability of the red point: 0.0000066 (accurate) and 0.013 (inaccurate).
Conditional Densities for Each Class

\[ Pr[Essay Length|Class] \]
Conditional Values for Numeric Predictors

\[ Pr[\text{Essay Length} = 8 | \text{Class}] \]
Conditional Probabilities for Categorical Predictors

\[ Pr[\text{Religion}|\text{Class}] \]
Combining Predictor Scores with the Prior

For an Atheist who wrote $e^8$ words, their likelihood values were:

- $Pr[Essay\ Length = 8|STEM] \times Pr[Religion = Atheism|STEM] = 0.518 \times 0.207 = 0.107$

- $Pr[Essay\ Length = 8|Other] \times Pr[Religion = Atheism|Other] = 0.428 \times 0.103 = 0.044$

However, when these are combined with the prior probability for each class, the relative probabilities show:

- $Pr[Predictors|STEM] \times Pr[STEM] = 0.107 \times 0.182 = 0.02$

- $Pr[Predictors|Other] \times Pr[Other] = 0.044 \times 0.818 = 0.036$

We don't need to compute the evidence; we can just normalize these values to add up to 1.

The results is that the posterior probability that this person is in a STEM field is 35.1%.
"Shrinkage" Properties of Bayesian Models

When our observed data is of high quality (top panel), the likelihood is narrow and dominates the equation.

However, when our observed information is poor, the likelihood can be very wide and diffuse.

When this occurs, Bayes' rule relies more on our prior belief than on the data in-hand.

This is generally called "shrinkage".
**Pros and Cons**

**Good:**

- This model can be very quickly trained (and theoretically in parallel).
- Once trained, the prediction is basically a look-up table (i.e. fast).
- Nonlinear class boundaries can be generated.

**Bad:**

- Linearly diagonal boundaries can be difficult.
- With many predictors, the class probabilities become poorly calibrated and U-shaped with most values near zero or one.
U-Shaped Class Probability Distributions

A completely non-informative data set was simulated using the naive assumption.

The training set has 500 data points over two classes and 450 predictors.

When a model is fit with 10 predictors, the distribution of the class probabilities gives us shapes that we would expect.

What happens when the number of predictors becomes larger?
U-Shaped Class Probability Distributions

n = 250 per class, Predictors: 10

- Truly Class A
- Truly Class B
Naive Bayes Data Preparations

There is a step specifically designed for converting binary dummy variables into factors.

First, we identify them, then use quasiquotation (via `!!!(x)`) to pass them to the step function.

```r
is_dummy <- function(x) length(unique(x)) == 2 & is.numeric(x)
dummy_or_not <- map_lgl(okc_train, is_dummy)
dummies <- names(dummy_or_not)[dummy_or_not]
head(dummies)
```

```
[1] "male" "essay_link" "tech" "technology" "math" "computer"
```

```r
no_dummies <- recipe(Class ~ ., data = okc_train) %>%
  step_bin2factor((!!!(dummies)) %>%
  step_zv(all_predictors())
```

We will use basic defaults for the tuning parameters:

```r
nb_grid <- expand.grid(usekernel = TRUE, fL = 0, adjust = 1)
```
Naive Bayes Training

```r
code
set.seed(5515)
nb_mod <- train(
  no_dummies,
  data = okc_train,
  method = "nb",
  metric = "ROC",
  tuneGrid = nb_grid,
  trControl = ctrl
)
```

Some warnings: ## Warning in FUN(X[[i]], ...): Numerical 0 probability for all classes with observation 1

This is due to the poorly calibrated probabilities although the warning is a bit misleading. This issue does not generally affect performance and can be ignored.
Naive Bayes Resampling Profile

```
## Naive Bayes
##
## 4000 samples
## 90 predictor
## 2 classes: 'stem', 'other'
##
## Recipe steps: bin2factor, zv
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 3600, 3601, 3599, 3600, 3600, 3600, ...
## Additional sampling using down-sampling
##
## Resampling results:
##
##  ROC  Sens  Spec
## 0.793  0.726  0.715
##
## Tuning parameter 'fL' was held constant at a value of 0
## Tuning parameter 'usekernel' was held constant at
## a value of TRUE
## Tuning parameter 'adjust' was held constant at a value of 1
```
Three ROC Curves

```r
all_curves <-
  all_curves %>%
  bind_rows(approx_roc_curve(nb_mod, "Naive Bayes"))

ggplot(all_curves) +
  aes(x = 1 - specificity, y = sensitivity,
      group = model, col = model) +
  geom_path() +
  geom_abline(col = "red", alpha = .5)
```
Test Set Results

test_res <- okc_test %>%
  dplyr::select(Class) %>%
  mutate(
    prob = predict(nb_mod, okc_test, type = "prob"), ["stem"],
    pred = predict(nb_mod, okc_test)
  )
roc_auc(test_res, Class, prob)

## # A tibble: 1 x 3
## #  .metric .estimator .estimate
## # <chr>   <chr>       <dbl>
## 1 roc_auc binary       0.791

getTrainPerf(nb_mod)

## # TrainROC TrainSens TrainSpec method
## # 1 0.793 0.726 0.715 nb
Test Set ROC Curve

two_class <- metric_set(sens, spec, accuracy)
two_class(test_res, truth = Class, estimate = pred)

## # A tibble: 3 x 3
## # <chr> <chr>       <dbl>
## 1 sens  binary     0.71
## 2 spec  binary     0.759
## 3 accura binary    0.749

test_roc <- roc_curve(test_res, Class, prob)

ggplot(test_roc) +
aes(x = 1 - specificity, y = sensitivity) +
geom_path() +
geom_abline(col = "red", alpha = .5)
Test Set Class Probabilities

```r
ggplot(test_res, aes(x = prob)) + geom_histogram(binwidth = .04) + facet_wrap( ~ Class)
```