CLASSIFICATIONS

Order from Chaos



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- 1 Variables
 - Categorical Variables
 - Continuous Variables
 - Converting Variable Types
- 2 Classifications
 - Logistic Regression
 - K-Means
 - Hierarchies
 - Random Forests
 - Networks

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Types of Variables

Variables can be classed into a multitude of types. The most common classification system knows:

Categorical Variables

- also known as Qualitative Variables
- Scales can be either
 - Nominal
 - Ordinal

Continuous Variables

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 - Discrete
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Categorical Variables

Categorical variables are those variables which **establish and fall into distinct groups and classes**.

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- assign each unit of the population to one of a finite number of groups
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In R, categorical variables usually come up as object type factor or character.

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Categorical Variables (Examples)

Examples of categorical variables:

- Biome Classifications (e.g. "Boreal Forest", "Tundra", etc.)
- Sex (e.g. "Male", "Female")
- Hierarchy Position (e.g. " α -Individual", " β -Individual", etc.)
- Soil Type (e.g. "Sandy", "Mud", "Permafrost", etc.)
- Leaf Type (e.g. "Compound", "Single Blade", etc.
- Sexual Reproductive Stage (e.g. "Juvenile", "Mature", etc.)
- Species Membership
- Family Group Membership
- ...

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Examples of categorical variables:

- Temperature
- Precipitation
- Weight
- pH
- Altitude
- Group Size
- Vegetation Indices
- Time
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Continuous variables can be converted into categorical variables via a method called **binning**:

Given a variable range, one can establish however many "bins" as one wants. For example:

- Given a temperature range of 271K 291K, there may be 4 bins of equa size:
 - Bin A: 271K < X < 276K
 - Bin B: 276K < X < 281K
 - Bin C: $281K < X \le 286K$
 - Bin D: $286K < X \le 291K$

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Confusion Of Units



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Theory

Logistic Regression

glm(..., family=binomial(link='logit')) in base R

Purpose: Understand how certain variables drive distinct outcomes.

- Down to Study-Design:
 - Variable values are **independent** (not paired)
 - Binary logistic regression: response variable is binary
 - Ordinal logistic regression: response variable is ordina
- Need for Post-Hoc Tests
 - Absence of influential outliers
 - Absence of multi-collinearity
 - Predictor Variables and log odds are related in a linear fashion

Assumntions

Theory

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Assumptions:

- Absence of influential outliers
 - Absence of influential outliers
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Example - The Data

```
library(titanic)
train_df <- na.omit(titanic_train) # remove NA rows
test_df <- train_df[c(1:50),c(2,3,5,6)] # 50 data for testing
train_df <- train_df[c(-1:-50),c(2,3,5,6)] # remaining data for training
head(train_df)
## Survived Pclass Sex Age</pre>
```

```
## 68 0 3 male 19
## 69 1 3 female 17
## 70 0 3 male 26
## 71 0 2 male 32
## 72 0 3 female 16
## 73 0 2 male 21
```

Can we explain **Survival** ('Survived') based on *Passenger class* ('Pclass'), *sex* ('Sex'), and *age* ('Age'). Was it really "Women and children first"?

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Example - The Model

Age

Pclass -1.29567 0.143615 -9.022 1.850e-19 ## Sexmale -2.45459 0.214837 -11.425 3.123e-30

```
Logistic Regression Coefficients can't be interpreted the same way as regular linear model coefficients since we are interested in survival probabilities between 0 and 1.
```

-0.03867 0.007937 -4.872 1.105e-06

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Example - Explanation & Prediction

Clearly, women of a young age in first class had the highest survival rate.

How do we know this? As *class* increases (from 1 to 3), survival probability decreases (-1.2957). Furthermore, men (*sexmale*) had, on average, a much lower survival rate than women (-2.4546). Lastly, increasing *age* negatively affected survival chances (-0.0387).

But how sure can we be of our model accuracy? We can test it by **predicting** some new data and **validating** our predictions:

```
# predict on test data
fitted <- predict(Logistic_Mod, newdata=test_df, type='response')
# if predicted survival probability above .5 assume survival
fitted <- ifelse(fitted > 0.5 , 1, 0)
# compare actual data with predictions --> ERROR RATE
mean(fitted != test_df$Survived)
## [1] 0.2
```

In reality, one would fine-tune the probability at which to assume survivorship!

Theory

K-Means Clustering

Mclust() in mclust package

Purpose: Identify a number of *k* clusters in our data.

Variance of the distribution of each variable is spherical

Issumptions: ■ All variables have the same variance

 \blacksquare Prior probability for all k clusters are the same

'mclust' is capable of identifying the statistically most appropriate number of clusters for the data set.

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Assumptions: All variables have the same variance

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Example - The Data I

```
data("iris")
head(iris)
##
     Sepal.Length Sepal.Width Petal.Length Petal.Width Species
## 1
             5.1
                         3.5
                                     1.4
                                                 0.2
                                                      setosa
## 2.
             4.9
                         3.0
                                     1.4
                                                 0.2 setosa
             4.7
                                     1.3
## 3
                       3.2
                                                 0.2 setosa
             4.6
                      3.1
                                     1.5
                                                 0.2 setosa
## 4
                                    1.4
## 5
            5.0
                       3.6
                                                0.2 setosa
## 6
           5.4
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                                     1.7
                                                 0.4 setosa
```

Can we accurately identify the 'Species' contained within the data set by clustering according to 'Sepal.Length', 'Sepal.Width', 'Petal.Length', and 'Petal.Width'?

Here, we decide to limit the number of clusters to the number of species present so we can test how well the prediction went.

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Example - The Data II

When building a *training* and *test* data set for identification of discrete values, we need to identify data of each group in both data sets. We do so via **stratified sampling**.

Doing this assures that we have data for each group to build a classifier as well as test the validity of our grouping.

Example - The Model I

```
library(mclust)
Mclust_mod <- Mclust (training_df[,-5], # data for the cluster model
                     G = length(unique(training_df[,5]))) # group number
plot (Mclust_mod, what = "uncertainty")
        Sepal.Length
                           Sepal.Width
                                              Petal.Length
                                                                 Petal Width
```

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Example - The Model II

Looking at the cluster centres and/or spreads can help with some **biological** interpretation.

```
Mclust_means <- Mclust_mod[["parameters"]][["mean"]] # extract means
colnames(Mclust_means) <- unique(training_df$Species) # set columns
Mclust_means
## setosa versicolor virginica
## Sepal.Length 4.9907 6.052 6.696
## Sepal.Width 3.4302 2.811 2.974
## Petal.Length 1.4628 4.539 5.759</pre>
```

I prefer a visualization as seen on the previous slide.

Petal.Width 0.2535 1.521 2.024

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Example - Explanation & Prediction

Clearly, Petal.Length, and Petal.Width are extremely good separators for our different clusters with the green and red clusters (versicolor and virginica) overlapping a lot in Sepal.Length and Sepal.Width space.

But how sure can we be of our model accuracy? We can test it by **predicting** the cluster membership and **validating** our predictions against the real data:

```
Mclust_pred <- predict.Mclust(Mclust_mod, iris[,-5]) # prediction
fitted <- Mclust_pred$classification # predicted species number
# compare actual data with predictions --> ERROR RATE
mean(fitted != as.numeric(iris$Species))
```

```
## [1] 0.1067
```

Theory

Hierarchical Clustering

hclust() in base R or rpart() in rpart package and many others

Purpose: Build a decision tree for classification of our data.

very easy to explain and inte

Easy to visualize.

- Easily handle qualitative predictors without the need to create dummy variables.
- Very sensitive to the choice of linkage.
- Generally do not have the same level of predictive accuracy as some of the other regression and classification approaches.
- Irees can be very non-robust

Disadvantages

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```

Again, let's see if we can accurately identify the 'Species' contained within the data set by clustering according to 'Sepal.Length', 'Sepal.Width', 'Petal.Length', and 'Petal.Width'.

Example - The Data II & Model I

'hclust()' can only handle distance matrices.

We a distance matrix between the numeric components of our data like so:

```
dist_mat <- dist(iris[, -5])
```

A distance matrix stores information about the dissimilarity of different observations.

Now, we can build our initial model:

```
clusters <- hclust(dist_mat, method = "complete")</pre>
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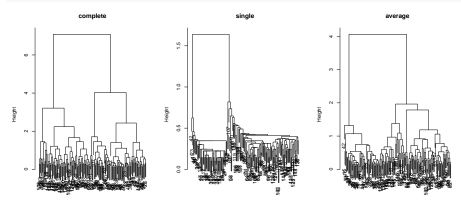
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```

Example - The Model II

```
par(mfrow = c(1,3))
plot(clusters, main = "complete")
plot(hclust(dist_mat, method = "single"), main = "single")
plot(hclust(dist_mat, method = "average"), main = "average")
```



Example - Explanation & Prediction

Hierarchical clustering recognises as many groups as there are observation and we may wish to **prune** the decision tree to a meaningful split level.

We know that we have three species in our data, so we may want to cut the complete tree at a height of 3 (not because that's the number of species, but because the tree just so happens to recognize three clusters at that level of decision-making).

```
clusterCut <- cutree(clusters, 3) # cut tree
table(clusterCut, iris$Species) # assess fit
##
## clusterCut setosa versicolor virginica</pre>
```

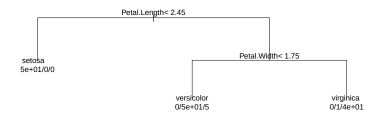
1 50 0 0 ## 2 0 23 49 ## 3 0 27 1

As we can see here, our decision tree has had no issue identifying *setosa* and *versicolor* into clusters 1 and 2 respectively. However, it is struggling with placing the species *virginica*.

Example - Decisions

So far we weren't able to tell the actual decision rules of how to cluster our data. Let's do this:

```
library(rpart)
fit <- rpart(Species ~. , data = iris)
plot(fit, margin = .1); text(fit, use.n = TRUE)</pre>
```



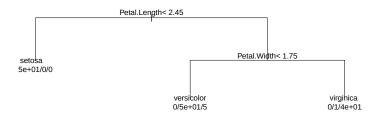
We can tell that our decisions for assigning species membership build on Petal. Length and Petal. Width in this example (remember the K-mean clustering)

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Purpose:

Random Forests

tuneRF() in randomForest package

Identify which variables to use for clustering our data and

build a tree.

vantages: Extremely powerful.

Very robust.

Easy to interpret

■ A black box algorithm.

Computationally expensive

Diodavarragoo

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```

One final time, we ask whether we can accurately identify the 'Species' contained within the data set by clustering according to 'Sepal.Length', 'Sepal.Width', 'Petal.Length', and 'Petal Width'

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Example - The Model

```
library(randomForest)
set.seed(42) # set seed because the process is random
RF_Mod <- tuneRF(x = iris[,-5], # variables which to use for clustering
                y = iris[,5], # correct cluster assignment
                strata = iris[,5], # stratified sampling
                doBest = TRUE, # run the best overall tree
                ntreeTry = 20000, # consider this number of trees
                improve = 0.0001, # improvement if this is exceeded
                trace = FALSE, plot = FALSE)
## -0.1429 0.0001
## 0 0.0001
RF Mod[["confusion"]]
##
            setosa versicolor virginica class.error
## setosa
                 50
                                                0.00
## versicolor 0
                            47
                                                0.06
## virginica
                                      47
                                                0.06
```

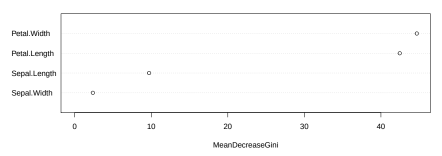
Example - Explanation

That is one stunningly accurate classification!

Let's see which variables where actually the most useful when making our clustering decisions:

varImpPlot (RF_Mod)





Network Clustering

cluster_optimal(), etc. in igraph package and many others

Purpose:

Identify compartments which are strongly connected within, but not between each other.

Advantages:

- Highly flexible approaches.
- Network analyses offer much more than clustering.
- Allow for clustering of very different data and identification relationships than other approaches.
- Steep learning curve

Disadvantages.

- Tricky in formatting data correctly.
- Choices can become overwhelming

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Example - The Data

Here, we take a foodweb contained within the foodwebs data collection of the igraphdata package. We are using the Middle Chesapeake Bay in Summer foodweb (Hagy, J.D. (2002) Eutrophication, hypoxia and trophic transfer efficiency in Chesa-peake Bay PhD Dissertation, University of Maryland at College Park (USA), 446 pp.).

```
library(igraph)
library(igraphdata)
data("foodwebs")
Foodweb_ig <- foodwebs[[2]]</pre>
```

Let's see what kind of network-internal clusters we can make out.

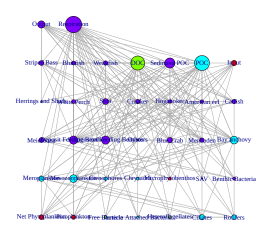
Example - A Directed Network

A directed network is one in which we know which node/vertex is acting one which other node/vertex.

We identify the clusters as follows:

```
Clusters <- cluster_optimal(Foodweb_ig)
Colours <- Clusters$membership
Colours <- rainbow(max(Colours))[Colours]
plot(Foodweb_ig,
    vertex.color = Colours,
    vertex.size = degree(Foodweb_ig) *0.5,
    layout-layout.grid, edge.arrow.size=0.001)</pre>
```

This identifies sub-networks/clusters by optimizing the modularity score of the overall network (i.e. optimizing connections within vs. between clusters).



Example - An Undirected Network

An **undirected network** is one in which we don't know which node/vertex is acting one which other node/vertex.

We identify the clusters as follows (there are more options):

```
Foodweb_ig <- as.undirected(Foodweb_ig)
Clusters <- cluster_fast_greedy(Foodweb_ig)
Colours <- Clusters$membership
Colours <- rainbow(max(Colours))[Colours]
plot(Foodweb_ig,
    vertex.color = Colours,
    vertex.size = degree(Foodweb_ig) *0.5,
    layout=layout.grid, edge.arrow.size=0.001)
```

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