Data Mining (Week 1)

# MSc Data Mining

Foundation Topic 05: Classification

Part 01: Introduction to Classification tals

Data Modelling
Advanced

ıle Based

Association Rules

Recommender Systems

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Supervised

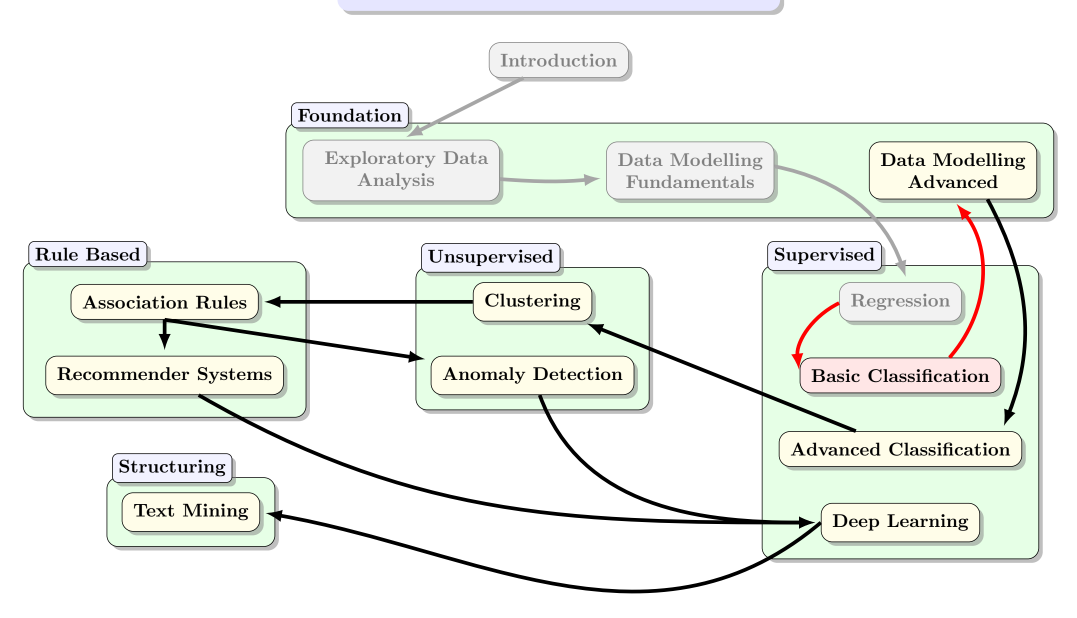
Regression

**Basic Classification** 

#### Outline

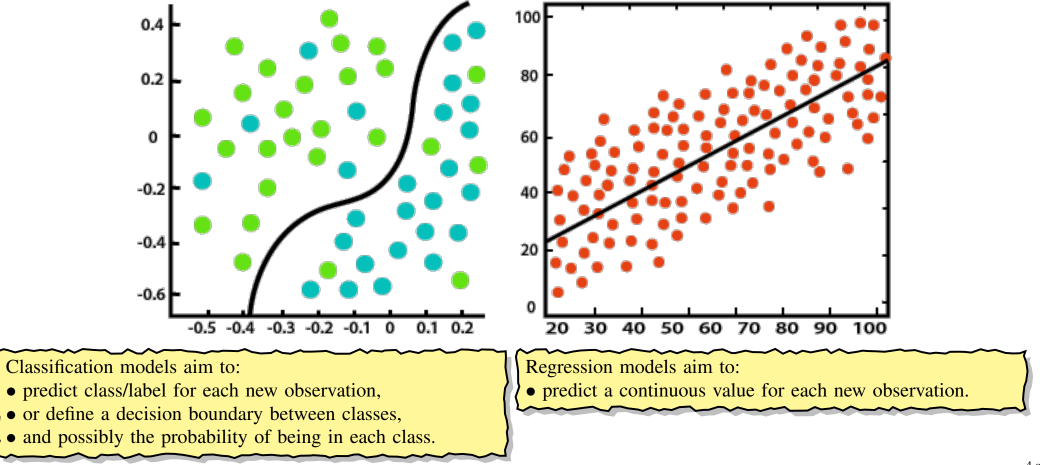
- How classification differs from regression
- Classification metrics
- Lazy vs Eager learners

## Data Mining (Week 5)



## Classification vs Regression

Supervised data models have a target. If target is quantitative (continuous) then have a regression model, if qualitative (categorical) then a classification model.



## Classification vs Regression

- Unlike regression, statistical distributions play a limited role in evaluating a classifier:
  - Scope for hypothesis testing is limited (there is no equivalent of the statsmodels diagnostic output (covered by Bernard, in week 4).
  - Depend on empirical metrics accuracy, precision, recall, f1-score, auc, ...
- Classification metrics tend to be easier to use/understand than those in regression classification metrics are based on counts of correct (or incorrect) cases divided by a subset of cases.
- Central concept in classification model is the confusion matrix:

		Predicted				
		Negative	Positive			
ual	Negative	True Negative (TN)	Type I error False Positive (FP)	N		
Actual	Positive	Type II error False Negative (FN)	True Positive ( <i>TP</i> )	P		
•		$\hat{N}$	$\hat{P}$	T		

## **Unbalanced Classification Datasets**

Practical classification datasets are often unbalanced — where the frequency of the classes in the target are very uneven:

• Telecommunication customer churn datasets.

Churn rate of 2%–10%.

Credit Card Fraud Detection

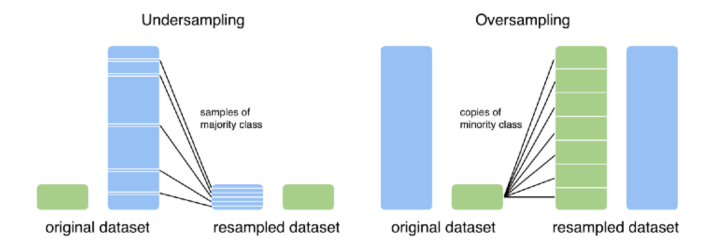
0.172% (492 frauds / 284,807 transactions).

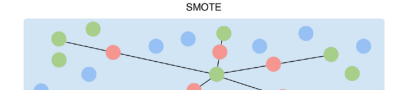
National Institutes of Health Chest X-Ray Dataset

14 cases in 5,606 cases

#### > Solutions >

Use metrics suitable for unbalanced datasets and/or techniques such as SMOTE for over/under sampling





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## **Summary of Classification Models**

	Data Pre-proc	essing*	Impact :	from	
Model	Normalisation	Scaling	Collinearity	Outliers	Summary
Logistic Regression	V	~	<b>✓</b>	<b>✓</b>	<ul> <li>Descriptive with good accuracy</li> <li>Linear relationship between features</li> <li>Reasonable computational requirements</li> </ul>
Naïve Bayes	NA	NA	<b>✓</b>	X	<ul><li>Works with categorical features only</li><li>Suitable for small train datasets</li></ul>
KNN	<b>✓</b>	<b>✓</b>	<b>✓</b>	×	<ul> <li>Local approximation, lazy learner</li> <li>Heavy computational requirements in prediction</li> </ul>
Random Forest (Week 8)	×	×	×	×	<ul> <li>High prediction accuracy</li> <li>Limited explainability</li> <li>Works with both continuous and categorical features</li> </ul>
Support Vector Classi (Week 8)	fier 🗶	<b>✓</b>	<b>X</b> / <b>V</b>	<b>✓</b>	<ul> <li>High prediction accuracy</li> <li>Explainability depends on kernel</li> <li>Computational effort depends on kernel</li> </ul>
Neural Networks (Week 12?)	×	~	<b>✓</b>	<b>✓</b>	<ul> <li>High prediction accuracy</li> <li>Self-extract features</li> <li>Heavy computational requirements</li> </ul>

<sup>\*</sup>Normalize (changing shape) using transformations, scale (change location/spread) via MinMaxScaler, StandardScaler, or RobustScaler if have outliers.

## Lazy vs Eager Learners

#### Lazy learner

Stores training data (or only minor processing) and uses this to compute prediction when given test data.

- Does not generalise until after training
- Does not produce a standalone model
- Training data must be kept for prediction
- Local approximations
- Often based on search
- New data is just added to the training data and model adapts, it can respond more easily to changing conditions

#### Eager learner

Builds a model from the train set, before receiving new data for prediction

- Training has an extra goal: to generalise from the data
- Training has an extra output: standalone model
- Training data can be discarded after use
- Local and/or global approximations
- Based on *computation*
- Models *drift* with time, so not suited to highly dynamic contexts, as it needs retraining

Usually an (eager) model requires much less memory than a (lazy) training set.

## A Non-perfect Test — Type I and Type II Errors

#### Consider an imperfect test with two outcomes, there are four possible outcomes:

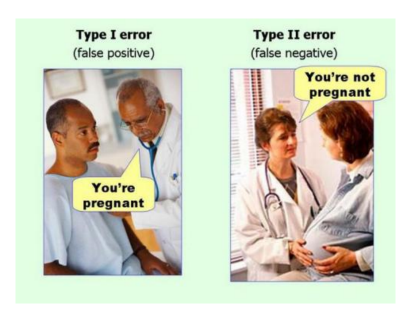
#### Confusion Matrix

# Predicted Negative Positive Negative Type I errorTrue Negative (TN) False Positive (FP)Positive $\hat{N}$ True Positive (TP) $\hat{N}$ $\hat{P}$ T

- If the test is applied to  $T = P + N = \hat{P} + \hat{N}$  observations / subjects / instances then we have four independent quantities TP, TN, FP, and FN.
- How do we combines these quantities into a single metric?
- The fraction of correct results seems like a good idea

$$accuracy = \frac{TN + TP}{N + P}$$

But what happens, if we are testing for a rare event? Maximising accuracy will result in the test always returning negative.



- Ideally we want the probability of either error to be zero but that may not be possible.
- Depending on the conditions we often modify the test to reduce probability of the type of error we don't want at the expense of increasing the probability of the other — think court case vs medical condition.

# Confusion matrix (Contingency table) Metrics

#### Accuracy — how often is the classifier correct?

Accuracy = 
$$\frac{TP + TN}{P + N}$$
(How often is the classifier correct?)

• False negative rate (FNR) =  $\frac{FN}{P} = 1 - TPR$ 

Predicted

Negative Positive

Negative Type I error

True Negative (TN) False Positive (FP)Positive Type II error

False Negative (FN) True Positive (TP)  $\hat{N}$   $\hat{P}$  T

- Sensitivity = Recall = True positive rate (TPR) =  $\frac{TP}{P}$  = 1 FNR (Of actual positive cases, how many were predicted positive?)
- Specificity =  $\frac{TN}{N}$  = 1 FPR (When it's actually no, how often does we predict no?) (Of actual negative cases, how many were predicted negative?)
- False positive rate (FPR) = false acceptance =  $\frac{FP}{N}$  = 1 Specificity
- **Precision** = positive predictive value (PPV) =  $\frac{TP}{\hat{P}} = \frac{TP}{TP + FP}$  (Of predicted positive cases, how many were actualy positive?)

Recall — important when the costs of false negatives are high

Precision — important when the costs of false positives are high

# Confusion matrix (Contingency table) Metrics

## $F_1$ Score

The F-measure or balanced F-score ( $F_1$  score, is a special case of the  $F_\beta$  score) is the harmonic mean of precision and recall:

$$F_1 = 2 \left[ \frac{1}{\frac{1}{\text{precision}} + \frac{1}{\text{recall}}} \right] = 2 \left[ \frac{\text{precision} \times \text{recall}}{\text{precision} + \text{recall}} \right]$$

#### A word of Caution ... >

Consider the three binary classifiers A, B and C

	A		A B			$\mathbb{C}$
	F	T	F	T	F	T
F	0	0.1	0.1	0	0.1 0.12	0
T	0	0.9	0.1	0.8	0.12	0.78

Metric	A	В	C	(best)
Accuracy	0.9	0.9	0.88	AB
Precision	0.9	1.0	1.0	BC
Recall	1.0	0.888	0.8667	A
F-score	0.947	0.941	0.9286	A

Clearly classifier A is useless since it always predicts label  $\tau$  regardless of the input. Also, B is slightly better than C (lower off- diagonal total).

Yet look at the performance metrics – B is never the clear winner.

We use some metrics because they are easy to understand, and not because they always give the "correct" result.

The mutual information between predicted and actual label (case) is defined

$$I(\hat{y}, y) = \sum_{\hat{y} = \{0,1\}} \sum_{y = \{0,1\}} p(\hat{y}, y) \log \frac{p(\hat{y}, y)}{p(\hat{y})p(y)}$$

where  $p(\hat{y}, y)$  is the joint probability distribution function.

This gives the intuitively correct rankings B > C > A

Metric	A	В	$\mathbf{C}$
Accuracy	0.9	0.9	0.88
Precision	0.9	1.0	1.0
Recall	1.0	0.888	0.8667
F-score	0.947	0.941	0.9286
<b>Mutual information</b>	0	0.1865	0.1735

# Multiclass Classifier — Micro Average vs Macro Average Performance

In a multi-class classifier we have more than two classes. To combine the metrics for individual classes to get an overall system metrics, we can apply either

#### Micro-Average Method

Sum up the individual true positives, false positives, and false negatives of the system for different classes and then apply totals to get the statistics.

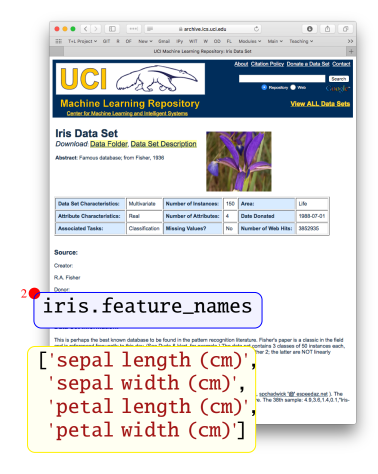
#### Macro-average Method

Average the precision and recall of the system on different classes.

See classification\_report from sklearn.metrics (Example: IRIS dataset)

<del></del>		` .	-	
	precision	recall	f1-score	support
setosa	1.00	0.95	0.97	19
versicolor		0.74	0.77	23
virginica	0.71	0.83	0.77	18
accuracy			0.83	60
macro avg	0.84	0.84	0.84	60
weighted avg	0.84	0.83	0.84	60

## Example: IRIS Dataset — Load



```
from sklearn import datasets
iris = datasets.load_iris()

df = pd.DataFrame(iris.data)
df.columns = iris.feature_names
df['target'] = iris.target_names[iris.target]
df.sample(4)
```

	sepal length (cm)	sepal width (cm)	petal length (cm)	petal width (cm)	target
1	<u>7</u> 5.1	3.5	1.4	0.3	setosa
8	<u>0</u> 5.5	2.4	3.8	1.1	versicolor
9	<del>7</del> 6.2	2.9	4.3	1.3	versicolor
9	<b>9</b> 5.7	2.8	4.1	1.3	versicolor

The data set contains, four numeric features, 3 classes of 50 instances each, where each class refers to a type of iris plant. One class is linearly separable from the other 2; the latter are NOT linearly separable from each other.

## Example: IRIS Dataset — Preprocess Data

We will cover some classifiers in a moment, but for now just treat the classifiers (LogisticRegression) as a black box and focus on the general process:

```
Extract the data (features and target)
```

The IRIS dataset has 4 features, but to simplify visualisation we are only going to use the first two<sup>†</sup> ('sepal length' and 'sepal width'):

```
dataset_name = "IRIS"
X, y, target_names = iris.data[:,:2], iris.target, iris.target_names
```

## Split dataset into train and test

We will keep 40% of the data for testing. Setting the parameter random\_state to a value means that we will get a random — but still reproducible — split.

```
from sklearn.model_selection import train_test_split
X_train, X_test, y_train, y_test = train_test_split(X, y, train_size=0.6, random_state=666)
```

<sup>†</sup>Python for Data Science — Cheat Sheet Numpy Basics

## Example: IRIS Dataset — Fit Model and Predict

## > Select classifier >

Scikit-learn supports a large set of classifiers, and aims to have a consistent interface to all. First import classifier and create instance . . .

from sklearn.linear\_model import LogisticRegression
model = LogisticRegression(max\_iter=500)

#### Train model

Then we train (fit) the classifier/model using only the features (X\_train) and targets (y\_train) from the train dataset ...

model.fit(X\_train, y\_train)

LogisticRegression(max\_iter=500)

#### Predict

Now that model is trained, we can use it to generate predictions, using the features (X\_test) from the test dataset ...

```
y_pred = model.predict(X_test)
```

## Example: IRIS Dataset — Evaluate

## Scoring and confusion matrix

We could just compute the score using whatever metric we have picked ...

```
from sklearn.metrics import accuracy_score
accuracy_score(y_test, y_pred)
```

0.8333333333333334

But this needs context, and even if "good" it can hide critical flaws. Lets look at the confusion matrix ...

```
from sklearn.metrics import confusion_matrix
cm = confusion_matrix(y_test,y_pred)
cm
```

```
array([[18, 1, 0],
[ 0, 17, 6],
[ 0, 3, 15]])
```

or, to get a nicer output, convert to a DataFrame ...

```
df_cm = pd.crosstab(target_names[y_test], target_names[y_pred])
  df_cm.index.name = 'Actual'
  df_cm.columns.name = 'Predicted'
  df_cm
```

Predicted	setosa	versicolor	virginica
Actual			
setosa	18	1	0
versicolor	0	17	6
virginica	0	3	15

## Example: IRIS Dataset — Evaluate

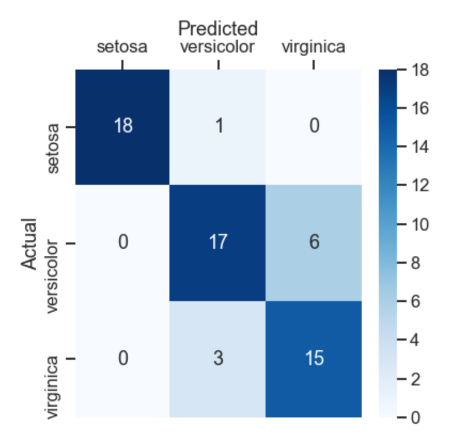
The confusion matrix is fundamental in evaluating a classifier, so find a presentation/visualisation that you like and use it. Here I have a heat map representation that I tend to use.

#### Predicted setosa versicolor virginica

Actual			,
setosa	18	1	0
versicolor	0	17	6
virginica	0	3	15

The first class **setosa** was only misclassified once, while the classifier had more difficulty between the second two classes.

plt.figure(figsize=(6,6))
g = sns.heatmap(df\_cm, annot=True, cmap="Blues")
g.xaxis.set\_ticks\_position("top")
g.xaxis.set\_label\_position('top')



## Example: IRIS Dataset — Evaluate

-0

The classification report, constructed from the confusion matrix, summaries the most common metrics per class and for overall averages ...

> from sklearn.metrics import classification\_report print(classification\_report(y\_test, y\_pred, target\_names=target\_names))

