# Text Data in Economics Warwick QAPEC Summer School

6. Machine Learning with Text

#### **ML** Essentials

Overview

Regression / Regularization

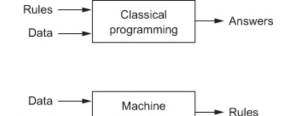
Binary Classification

Multi-Class Models

Ensemble Learning with XGBoos

## What is machine learning?

Answers



learning

- In classical computer programming, humans input the rules and the data, and the computer provides answers.
- ► In machine learning, humans input the data and the answers, and the computer learns the rules.

# What do ML Algorithms do? Fit a function to data points

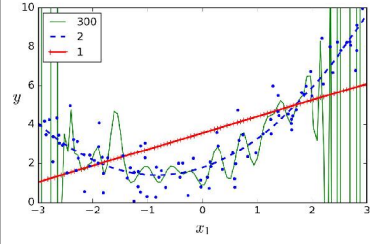


Figure 4-14. High-degree Polynomial Regression

## What do ML Algorithms do? Minimize a cost function

► A typical cost function (or loss function) for regression problems is Mean Squared Error (MSE):

$$MSE(\theta) = \frac{1}{n_D} \sum_{i=1}^{n_D} (h(x_i; \theta) - y_i)^2$$

- $\triangleright$   $n_D$ , the number of rows/observations
- $\triangleright$  x, the matrix of predictors, with row  $x_i$
- $\triangleright$  y, the vector of outcomes, with item  $y_i$
- $h(x_i;\theta) = \hat{y}$  the model prediction (hypothesis)

The **data** (x,y) are taken as given, and the ML algorithm searches for **parameters**  $\theta$  to minimize the cost function.

# Linear Regression is Machine Learning

▶ Ordinary Least Squares Regression (OLS) assumes the functional form  $f(x;\theta) = x_i'\theta$  and minimizes the mean squared error (MSE)

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This minimand has a closed form solution

$$\hat{\theta} = (\mathbf{x}'\mathbf{x})^{-1}\mathbf{x}'\mathbf{y}$$

**most machine learning models do not have a closed form solution**  $\rightarrow$  use numerical optimization instead (gradient descent).

$$MSE(\theta) = \frac{1}{n_D} \sum_{i=1}^{n_D} (h(\theta; \boldsymbol{x}_i) - y_i)^2$$

► The partial derivative for feature *j* is

$$\frac{\partial \mathsf{MSE}}{\partial \theta_j} = \frac{2}{n_D} \sum_{i=1}^{n_D} \left( \underbrace{h(\theta; \mathbf{x}_i) - y_i}_{\text{error for this obs}} \right) \underbrace{\frac{\partial h(\theta; \mathbf{x}_i)}{\partial \theta_j}}_{\text{how } \theta_i \text{ shifts } h(\theta)}$$

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- ightharpoonup estimates how changing  $\theta_i$  would reduce the error across the whole dataset.
- The gradient ∇ gives the vector of these partial derivatives for all features:

$$\nabla_{\theta}\mathsf{MSE} = \begin{bmatrix} \frac{\partial \mathsf{MSE}}{\partial \theta_1} \\ \frac{\partial \mathsf{MSE}}{\partial \theta_2} \\ \vdots \\ \frac{\partial \mathsf{MSE}}{\partial \theta_{n_x}} \end{bmatrix}$$

▶ **Gradient descent** nudges  $\theta$  against the gradient (the direction that reduces MSE):

$$\theta_{t+1} = \theta_t - \eta \nabla_{\theta} \mathsf{MSE}$$

 $ightharpoonup \eta = \text{learning rate}$ 

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- **Each** document *i* has an associated outcome or label  $y_i$  with dimensions  $n_y \ge 1$
- lacktriangle Some documents are labeled and some are unlabeled ightarrow
  - we would like to learn a function  $\hat{y}(d_i)$  based on the labeled data ...
  - ... to machine-classify the unlabeled data.

## First Problem

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- **Each** document is a sequence of symbols  $d_i$ , while (standard) ML algorithms work on numbers.
- ► The solution: all the methods from previous lectures for extracting informative numerical information from documents:
  - style features
  - counts over dictionary patterns
  - tokens
  - n-grams
  - principal components
  - topic shares
  - etc.
- ▶ documents can thus be **featurized** represented as a matrix of vectors x with  $n_x \ge 1$  features.

#### **ML** Essentials

#### Overview

Regression / Regularization Binary Classification Multi-Class Models

Ensemble Learning with XGBoost

# Three Types of (Standard) Machine Learning Problems

Determined by the data type of the outcome variable (or label):

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  - e.g., guilty or innocent
- ▶ **Regression**: a one-dimensional, continuous, real-valued outcome.
  - e.g., number of days of prison assigned
- Multinomial Classification: Three or more discrete, un-ordered outcomes.
  - e.g., predict what judge is assigned to a case: Alito, Breyer, or Cardozo

## Loss functions, more generally

- ▶ The loss function  $L(\hat{y}, y)$  assigns a score based on prediction and truth:
  - ▶ Should be bounded from below, with the minimum attained only for cases where the prediction is correct.
- ► The average loss for the test set is

$$\mathcal{L}(\theta) = \frac{1}{n_D} \sum_{i=1}^{n_D} L(h(\mathbf{x}_i; \theta), \mathbf{y}_i)$$

▶ The estimated parameter matrix  $\theta$  solves

$$\hat{ heta} = rg \min_{ heta} \mathcal{L}( heta)$$

 $\hookrightarrow$  optimizes over parameter space; treats the data as constants.

#### Gradient Descent

- even when cost function is not convex (eg neural nets), gradient descent often gets decent results.
- ▶ **Stochastic** gradient descent (SGD) computes the gradient for a single randomly sampled instance (at each iteration).
  - ► Much faster, still works well.

## Data Prep for Machine Learning

- ▶ Data Pre-Processing: See Geron Chapter 2 for pandas and sklearn syntax:
  - imputing missing values.
  - feature scaling (often helpful/necessary for ML models to work well)
    - ▶ if predictors are sparse (e.g. bag-of-words), use StandardScaler(with\_mean=False).
  - encoding categorical variables.
  - Best practice: reproducible data pipeline.

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- ► Train/Test Split:
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  - standard approach: randomly sample 80% training dataset to learn parameters, form predictions in 20% testing dataset for evaluating performance.

# Use Cross-Validation During Model Training

- Within the training set:
  - Use cross-validation with grid search to get model performance metrics across subsets of data using different hyperparameter specs.
  - Find the best hyperparameters for out-of-fold prediction in the training set.
- ▶ Then evaluate model performance in the test set using these hyperparameters.

## Model Evaluation in Test Set

Evaluating a "good" model is context-dependent. Here are some basics.

## Regression:

- mean squared error (MSE)
- ▶ R-squared (same ranking as MSE, but units are more interpretable)
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#### Classification:

- ▶ more complicated, but accuracy is a good baseline: accuracy = (# correct test-set predictions) / (# of test-set observations)
- ▶ What if one of the outcomes is over-represented e.g., 19 out of 20? Then I can guess the modal class and get 95% accuracy.
  - ▶ Some alternative classifier metrics designed to address class imbalance (more below).

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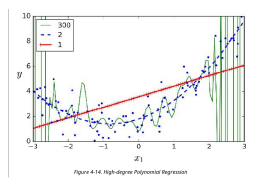
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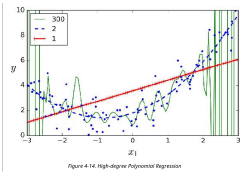
## Regression models ↔ Continuous outcome

- If the outcome is continuous (e.g., Y = tax revenues collected, or criminal sentence imposed in months of prison):
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▶ *Regularization*: model training methods designed to reduce/prevent over-fitting.

$$\hat{\boldsymbol{\theta}} = \arg\min_{\boldsymbol{\theta}} \frac{1}{n_D} \sum_{i=1}^{n_D} L(h(\boldsymbol{x}_i; \boldsymbol{\theta}), \boldsymbol{y}_i) + \lambda R(\boldsymbol{\theta})$$

- $ightharpoonup R(\theta)$  is a "regularization function" or "regularizer", designed to reduce over-fitting.
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"Lasso" (or L1) penalty:

$$R_1 = \|\theta\|_1 = \sum_{j=1}^{\infty} |\theta_j|$$

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shrinks coefficients toward zero and helps select between collinear predictors.

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- Elastic Net:  $R_{\text{enet}} = \lambda_1 R_1 + \lambda_2 R_2$

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# Binary Outcome ↔ Binary Classification

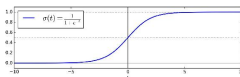
- ▶ Binary classifiers try to match a boolean outcome  $y \in \{0,1\}$ .
  - The standard approach is to apply a transformation (e.g. sigmoid/logit) to normalize  $\hat{y} \in [0,1]$ .
  - ▶ Prediction rule is 0 for  $\hat{y} < .5$  and 1 otherwise.

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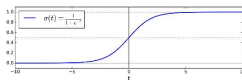
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- ► The binary cross-entropy (or log loss) is:

$$L(\theta) = \underbrace{-\frac{1}{n_D} \sum_{i=1}^{n_D} \left[ \underbrace{y_i}_{y_i=1} \underbrace{\log(\hat{y}_i)}_{\log \text{ prob} y_i=1} + \underbrace{(1-y_i) \underbrace{\log(1-\hat{y}_i)}_{\log \text{ prob} y_i=0} \right]}_{\log \text{ prob} y_i=0}$$

$$\hat{y} = \operatorname{sigmoid}(\mathbf{x} \cdot \theta) = \frac{1}{1 + \exp(-\mathbf{x} \cdot \theta)}$$



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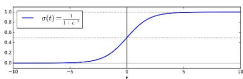


▶ Plugging into the binary-cross entropy loss gives the logistic regression cost objective:

$$\min_{\theta} \sum_{i=1}^{n_D} -y_i \log(\operatorname{sigmoid}(\boldsymbol{x}_i \cdot \theta)) - [1 - y_i] \log(1 - \operatorname{sigmoid}(\boldsymbol{x}_i \cdot \theta))$$

does not have a closed form solution, but it is convex (guaranteeing that gradient descent will find the global minimum).

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Like linear regression, logistic regression can be regularized with L1 or L2 penalties.

		Predicted Class		
		Negative	Positive	
True Class	Negative	# True Negatives	# False Positives	
	Positive	# False Negatives	# True Positives	

► Cell values give counts in the test set.

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Precision decreases with false positives. "When I guess this outcome, I tend to guesses correctly."

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$$Precision (for positive class) = \frac{True Positives}{True Positives + False Positives}$$

Precision decreases with false positives. "When I guess this outcome, I tend to guesses correctly."

Recall (for positive class) = 
$$\frac{\text{True Positives}}{\text{True Positives} + \text{False Negatives}}$$

Recall decreases with false negatives. "When this outcome occurs, I don't miss it."

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**Balanced accuracy** = the average recall in both classes:

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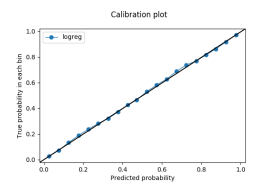
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#### **AUC-ROC** = Area Under the Receiver Operating Characteristic Curve

- provides an aggregate measure of performance across all possible classification thresholds.
- ▶ Interpretation: randomly sample one positive and one negative example. AUC = probability that the model correctly guesses which is which.

# **Evaluating Classification Models: Calibration Curves**



- ► Plotting the binned fraction in a category (Y axis) against the predicted probability in a category (X axis):
- Provides evidence of whether the classifer is replicating the conditional distribution of the outcome.

```
from seaborn import regplot
regplot(y_test, y_pred, x_bins=20)
```

Andrew Peterson and Arthur Spirling, "Classification accuracy as a substantive quantity of interest: Measuring polarization in Westminster systems," *Political Analysis* (2018).

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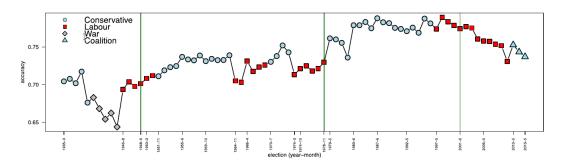
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In years that classifier is more accurate, speech is more polarized:



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#### Multiple Classes: Setup

▶ The outcome is  $y_i \in \{1,...,k,...,n_y\}$  output classes, which can also be represented as a one-hot vector

$$\mathbf{y}_i = \{\mathbf{1}[y_i = 1], ..., \mathbf{1}[y_i = n_y]\}$$

#### Multiple Classes: Setup

▶ The outcome is  $y_i \in \{1,...,k,...,n_y\}$  output classes, which can also be represented as a one-hot vector

$$\mathbf{y}_i = {\mathbf{1}[y_i = 1], ..., \mathbf{1}[y_i = n_y]}$$

▶ We want to learn a vector function

$$\mathbf{y} = \mathbf{h}(\mathbf{x}, \theta)$$

taking text features x as inputs and outputing a vector of probabilities across outcome classes:

$$\hat{\mathbf{y}} = {\{\hat{y}^1, ..., \hat{y}^{n_y}\}}, \sum_{k=1}^{n_y} \hat{y}^k = 1, \hat{y}^k \ge 0 \ \forall k$$

for prediction step, can select the highest-probability class:

$$\tilde{y} = \arg\max_{k} \hat{y}_{[k]}$$

# Categorical Cross Entropy

► The standard loss function in multinomial classification is **categorical cross entropy**:

$$L(\theta) = -\sum_{k=1}^{n_y} \mathbf{y}^k \log(\hat{y}^k(\mathbf{x}, \theta))$$

measures dissimilarity between the true label distribution y and the predicted label distribution  $\hat{y}$ .

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- measures dissimilarity between the true label distribution y and the predicted label distribution  $\hat{y}$ .
- Since there is just one true class  $(y = 1 \text{ for one class } k^*, \text{ and zero for others}),$  simplifies to

$$L(\theta) = -\log(\hat{y}^{k^*}(\boldsymbol{x}, \theta))$$

- Rewards putting higher probability on the true class, ignores distribution of probabilities on other classes.
- function is convex  $\rightarrow$  gradient descent will find the optimum.

#### Multinomial Logistic Regression

Multinomial logistic regression computes probabilities for each class k using the softmax transformation

$$\hat{y}_k(\mathbf{x}_i) = \Pr(y_i = k) = \frac{\exp(\theta'_k \mathbf{x}_i)}{\sum_{l=1}^{n_y} \exp(\theta'_l \mathbf{x}_i)}$$

- ightharpoonup softmax is the multiclass generalization of sigmoid ightharpoonup can then interpret  $\hat{y}$  as probabilities.
- ▶  $n_x$  features and  $n_y$  output classes  $\rightarrow$  there is a  $n_y \times n_x$  parameter matrix  $\Theta$ , where the parameters for each class  $\theta_k$  are stored as rows.

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#### The **L2-penalized logistic regression** has loss function

$$\mathcal{L}(\theta) = -\frac{1}{n_D} \sum_{i=1}^{n_D} \log \frac{\exp(\theta'_{k^*} \mathbf{x}_i)}{\sum_{j=1}^{n_y} \exp(\theta'_j \mathbf{x}_i)} + \lambda \sum_{j=1}^{n_x} \sum_{k=1}^{n_y} (\theta_{[j,k]})^2$$

- $\lambda$  = strength of L2 penalty (could also add lasso penalty)
  - as before, predictors should be scaled to the same variance.

		Predicted Class		
		Class A	Class B	Class C
	Class A	Correct A	A, classed as B	A, classed as C
True Class	Class B	B, classed as A	Correct B	B, classed as C
	Class C	C, classed as A	C, classed as B	Correct C

More generally, with **multi-class confusion matrix** M with items  $M_{ij}$  (row i, column j):

Precision for 
$$k = \frac{\text{True Positives for } k}{\text{True Positives for } k + \text{False Positives for } k} = \frac{M_{kk}}{\sum_{l} M_{lk}}$$
Recall for  $k = \frac{\text{True Positives for } k}{\text{True Positives for } k + \text{False Negatives for } k} = \frac{M_{kk}}{\sum_{l} M_{kl}}$ 

$$F_1(k) = 2 \times \frac{\operatorname{precision}(k) \times \operatorname{recall}(k)}{\operatorname{precision}(k) + \operatorname{recall}(k)}$$

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Can average these metrics across classes to get aggregate metrics.

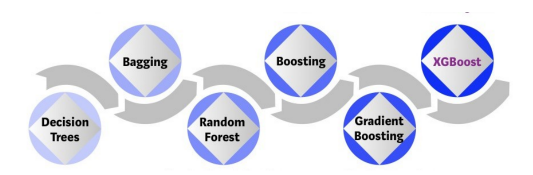
- e.g., balanced accuracy = unweighted average of recalls across classes.
- can weight classes by their frequency in dataset

#### ML Essentials

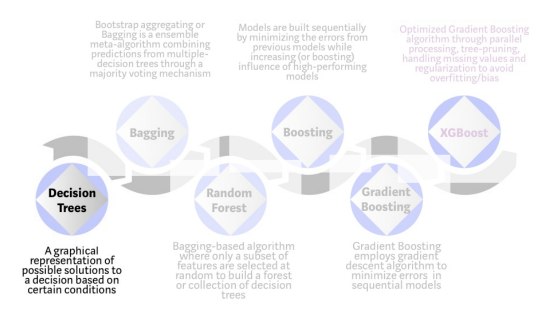
Overview
Regression / Regularization
Binary Classification
Multi-Class Models

Ensemble Learning with XGBoost

#### XGBoost: Overview

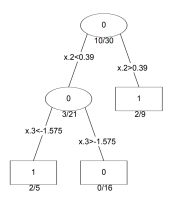


#### XGBoost Ingredients: Decision Trees



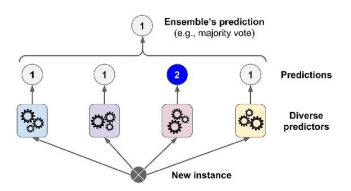
#### **Decision Trees**

#### Classification Tree



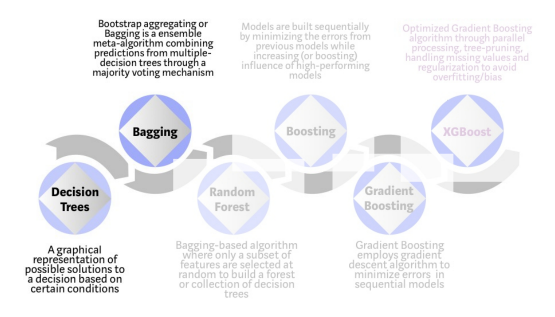
- ▶ Decision trees learn a series of binary splits in the data based on hard thresholds.
  - if yes, go right; if no, go left.
- Can have additional splits as you move through the tree.
- fast and interpretable, but performance is often poor.

#### **Voting Classifiers**



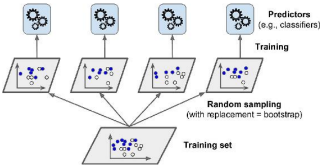
- voting classifiers (ensembles of different models that vote on the prediction) generally out-perform the best classifier in the ensemble.
  - more diverse algorithms will make different types of errors, and improve your ensemble's robustness.

#### XGBoost Ingredients: Bootstrapping



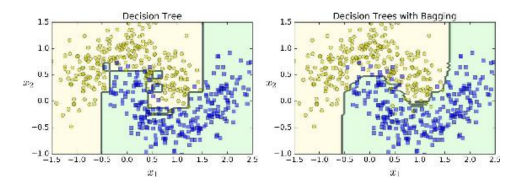
#### Bootstrapping

▶ Rather than use the same data on different classifiers, one can use different subsets of the data on the same classifier:



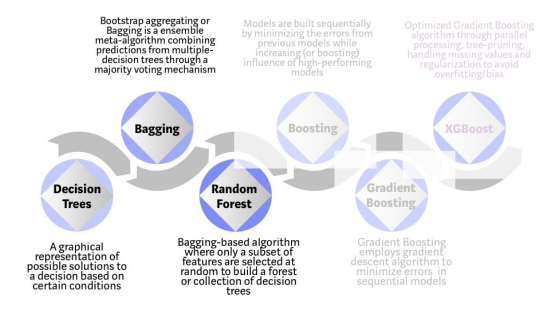
can also use different subsets of features across subclassifiers.

#### **Bootstrapping Benefits**



- ▶ A bootstraped ensemble generally has a similar bias but lower variance than a single predictor trained on all the data.
- ▶ Predictors can be trained in parallel using separate CPU cores.

## XGBoost Ingredients: Random Forests



Random Forests are optimized ensembles of bootstrapped decision trees:

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from sklearn.ensemble import RandomForestClassifier
rfc = RandomForestClassifier()
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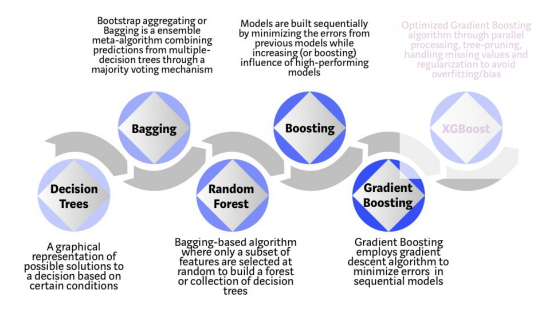
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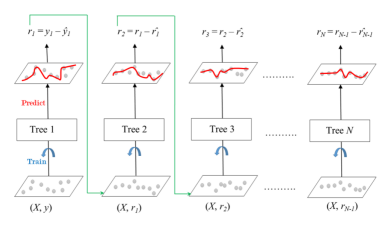
- 1. Each voting tree gets its own sample of data.
- 2. At each tree split, a random sample of features is drawn, only those features are considered for splitting.
- 3. For each tree, error rate is computed using data outside its bootstrap sample.

# XGBoost Ingredients: Gradient Boosting



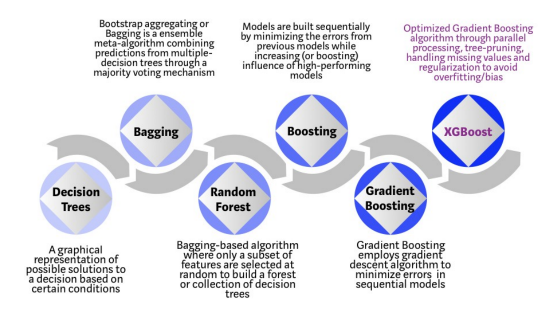
# **Gradient Boosting Machines**

Gradient boosting refers to an additive ensemble of trees:



Adds additional layers of trees to fit the residuals of the first layers

## XGBoost Ingredients



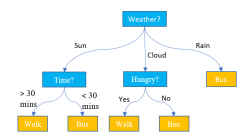
#### **XGBoost**

- ► Feurer et al (2018) find that XGBoost beats a sophisticated AutoML procedure with grid search over 15 classifiers and 18 data preprocessors.
- A good starting point for any machine learning task.
- easy to use
- actively developed
- efficient / parallelizable
- provides model explanations
- takes sparse matrices as input

from xgboost import XGBClassifier

### Tree Ensembles are Black Boxes

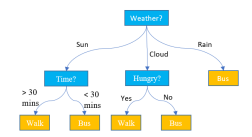
Small decision trees have the advantage of being highly interpretable.



#### Tree Ensembles are Black Boxes

Small decision trees have the advantage of being highly interpretable.





- Larger trees and ensembles (e.g. XGBoost) lose this nice feature.
- Best-performing ML models are hard to interpret because they use lots of features and exploit non-linearities and interactions.

### Interpreting Tree Ensembles

#### XGBoost's Feature Importance Metric:

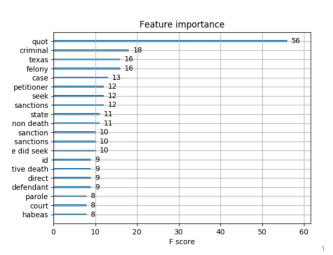
- ► At each decision node, compute **information gain** for feature *j* **(change in predicted probability**).
- Average across all nodes for each j.

Ranks predictors by their relative contributions.

```
from xgboost import plot_importance
plot_importance(xgb_reg, max_num_features=10)
```

### Feature Importance

```
from xgboost import plot_importance
plot_importance(xgb_reg, max_num_features=20)
<IPython.core.display.Javascript object>
```



➤ XGBoost provides a metric of feature importance that summarizes how well each feature contributes to predictive accuracy.

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- 6. Answer the research question!