## Introduction to Ensemble methods

- · We have looked at different machine learning models now
  - K-Nearest Neighbours
  - Classification & Regression Trees (CART)
  - Logistic Regression
  - Neural Networks
  - Support Vector Machines
- · Question: Given a dataset, which ML algorithm should we pick, and how do you know which technique will perform the best?
- · Unfortunately, there is no good answer to this question.
  - It is mostly a process of trial-and-error
  - Each kind of ML algorithm yields a different model/hypothesis - But there is no perfect model/hypothesis in practice

· So you may ask could we combine several imperfect models into a better model?

- · Analogies of combining multiple models in our society
  - Elections combine voter's choices to pick a "good" candidate
  - Committees combine several experts' apinion to make better decisions
- . Intuition behind combining multiple models/hypotheses
  - Individuals (or individual models) often make mistakes, but the "majority" is less likely to make mistakes
  - Individuals often have partial knowledge, but a committee can pool expertise to make better decisions
- · Ensemble learning can combine an ensemble of
  - Different types of base models (e.g. Neural networks, CART and SVM)
  - Same base model trained slightly differently V We are going to follow this approach

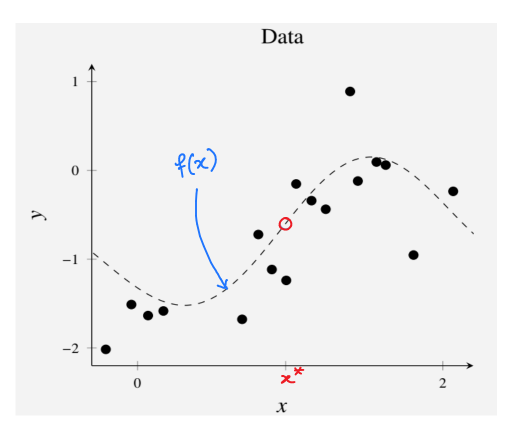
Bagging (or Bootstrap Aggregating)

· A central concept in ML is the bias-variance tradeoff

- The more flexible a model is, the lower its bias will be

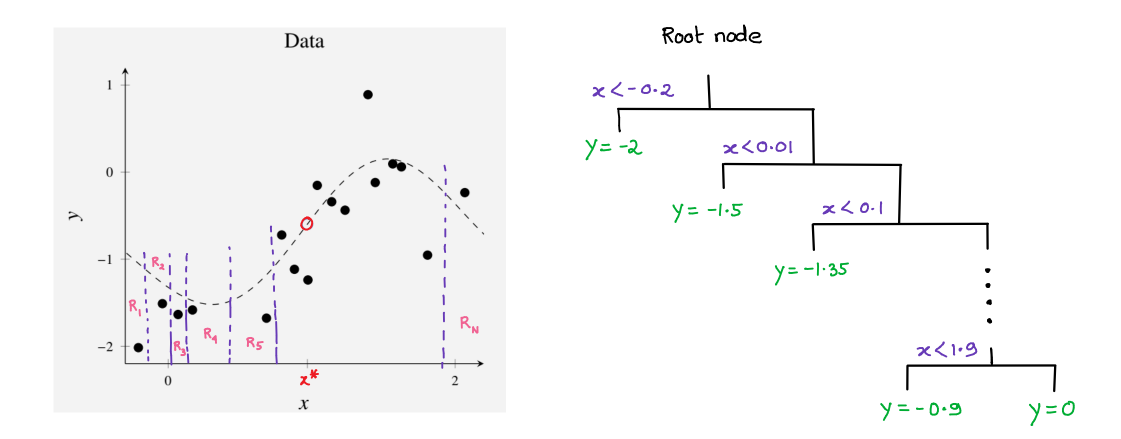
- Examples of highly flexible models that can represent complicated input-output relationships are K-Nearest Neighbours, CART, NNs, etc.
- . The downside of such highly flexible models is the risk of overfitting
- · Overfitted models lead to unwanted high variance in predictions
- By using bagging, we can reduce the variance of the base model without increasing its bias
- · Lets take an example of regression trees with bagging

- Consider the data obtained as  $y = f(x) + \epsilon$  noise
- We would like to train an ML model using this data, so as to be able to predict new data points well
- A good prediction would mean that the brained model should predict f(x) shown by the dotted line well at  $x^*$

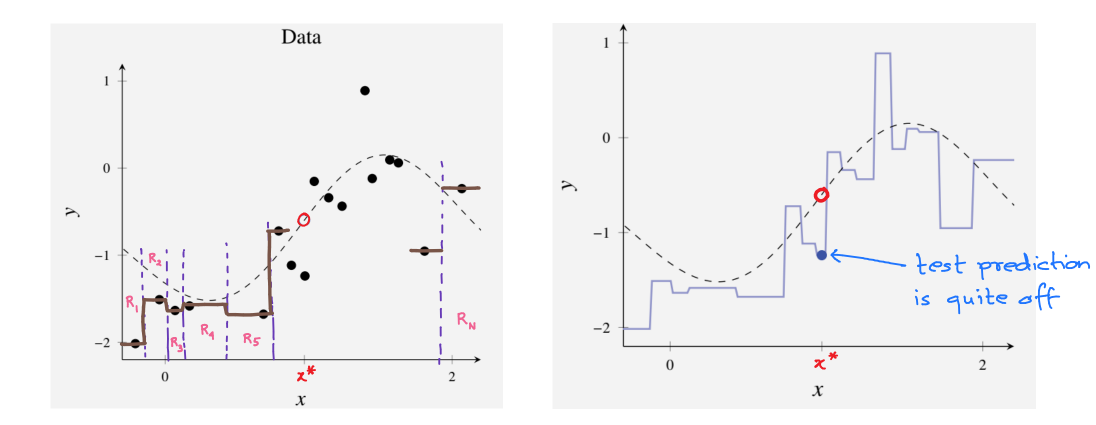


• For this problem, let us use Regression Trees as the chosen ML method, since they are non-parametric methods and are very flexible

- Recall that in Classification and Regression trees, we partition the input space using box-shaped decision boundaries
- Lets consider a Regression tree which is grown until each leaf node has only one data point

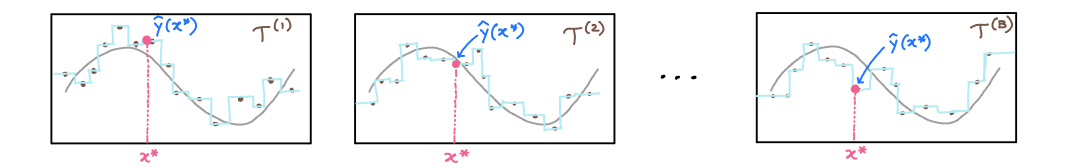


• On fitting a regression tree (with one data point in each leaf), we get an OVERFITTED Regression Tree



- Due to overfitting, the resulting regression tree is a low-bias-high-variance
   model
  - high variance means the trained model is very sensitive to the training data; if the training data changes, the predictions change a lot

- Because of the noise in training data, we can think of the prediction  $\hat{y}(x^*)$  from the trained model as a random variable It means that if we had multiple datasets and we trained different RTs
  - on them, each of their predictions  $\widehat{y}(x^*)$  would be different



• So if we assumed that we had access to B independent datasets  $T^{(1)}, T^{(2)}, \dots, T^{(B)}$ , then we could brain a separate tree for each dataset and obtain separate predictions  $\hat{y}_{b}(\underline{x}^{*})$ ,  $b=1, 2, \dots, B$ , then:

- Each 
$$y_{b}(\underline{x}^{*})$$
 would have low bias and high variance  
- By averaging  $\hat{y}(\underline{x}^{*}) = \frac{1}{B} \sum_{b=1}^{B} \hat{y}_{b}(\underline{x}^{*})$ , the bias is kept  
small, but the variance is reduced by a factor of B! (Proof?)

Probability detour - Variance reduction by averaging

Let  $z_1, z_2, ..., z_B$  be a collection of identically distributed but possibly dependent random variables, with

Mean : 
$$\mathbb{E}[z_b] = \mu$$
  
Variance:  $Var(z_b) = \sigma^2$  for  $b = 1, 2, ..., B$ 

Correlation: Corr 
$$(z_i, z_j) = P$$
  $i \neq j$ ,  $i, j = 1, 2, ..., B$ 

Then one can show that the mean and variance of the average  $\frac{1}{B} \sum_{b=1}^{B} z_{b}$  are: (assuming P>0)

$$\mathbb{E}\left[\frac{1}{B}\sum_{b=1}^{B}z_{b}\right] = \mu, \quad Var\left[\frac{1}{B}\sum_{b=1}^{B}z_{b}\right] = \frac{1-\rho}{B}\sigma^{2} + \rho\sigma^{2}$$
  
small for large B

- · Problem: We only have access to one training dataset
- · Solution : Bootstrap the data!
- Bootstrap is a method of artificially creating multiple datasets (of size N) out of one dataset (also of size N)

- Sample N times with replacement from the original training data 
$$T = \{ \underline{X}^{(i)}, y^{(i)} \}_{i=1}^{N}$$

- Repeat B times to generate B "bootstrapped" training datasets 
$$\tilde{T}^{(1)}, \tilde{T}^{(2)}, \dots, \tilde{T}^{(B)}$$

- BAGGING
  - For each bootstrapped dataset  $\tilde{T}^{(b)}$ , we train a tree (basemodel) Averaging them,  $\hat{Y}_{bag} = \frac{1}{B} \sum_{b=1}^{B} \tilde{y}^{b}(\underline{x})$

Bagging example with regression trees as basemodel

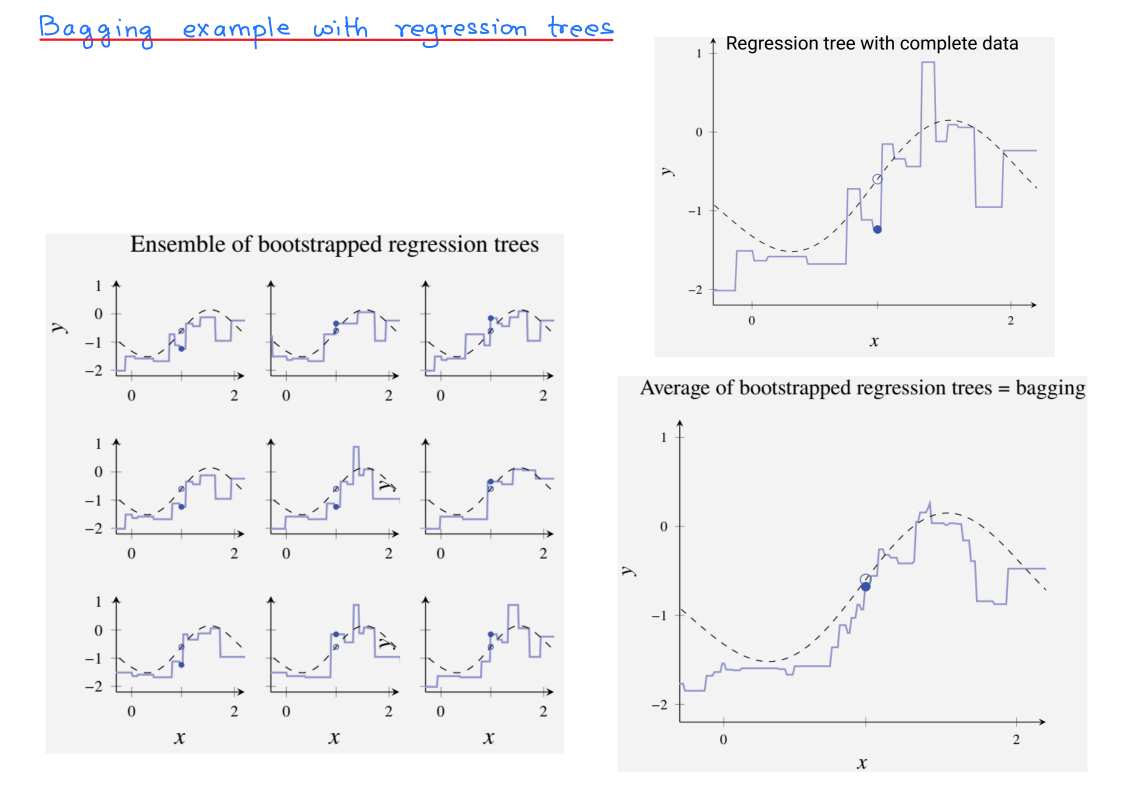
Assume that we have a training set

$$\mathcal{T} = \left\{ \left( \underline{\mathbf{x}}^{(1)}, y^{(1)} \right), \left( \underline{\mathbf{x}}^{(2)}, y^{(2)} \right), \left( \underline{\mathbf{x}}^{(3)}, y^{(3)} \right), \cdots, \left( \underline{\mathbf{x}}^{(n)}, y^{(n)} \right) \right\}$$

• We generate, say, B = 9 datasets by bootstrapping:

$$\begin{split} \widetilde{\tau}^{(1)} &= \left\{ \left( \underline{x}^{(1)}, \gamma^{(1)} \right), \left( \underline{x}^{(2)}, \gamma^{(2)} \right), \left( \underline{x}^{(3)}, \gamma^{(3)} \right), \cdots, \left( \underline{x}^{(3)}, \gamma^{(3)} \right) \right\} \\ \widetilde{\tau}^{(2)} &= \left\{ \left( \underline{x}^{(1)}, \gamma^{(1)} \right), \left( \underline{x}^{(n)}, \gamma^{(n)} \right), \left( \underline{x}^{(n)}, \gamma^{(n)} \right), \cdots, \left( \underline{x}^{(n)}, \gamma^{(n)} \right) \right\} \\ \vdots \\ \widetilde{\tau}^{(9)} &= \left\{ \left( \underline{x}^{(1)}, \gamma^{(1)} \right), \left( \underline{x}^{(1)}, \gamma^{(1)} \right), \left( \underline{x}^{(2)}, \gamma^{(2)} \right), \cdots, \left( \underline{x}^{(3)}, \gamma^{(3)} \right) \right\} \end{split}$$

• We compute B = 9 (deep) regression trees  $\tilde{y}^{(1)}(\underline{x}), \tilde{y}^{(2)}(\underline{x}), ..., \tilde{y}^{(9)}(\underline{x})$ one for each dataset  $\tilde{\tau}^{(1)}, \tilde{\tau}^{(2)}, ..., \tilde{\tau}^{(9)}$ , and average  $\tilde{y}_{bag} = \frac{1}{9} \sum_{b=1}^{9} \tilde{y}^{(b)}(\underline{x})$ 



## Bagging algorithm

• Training : Learn all base models Data: Training dataset  $T = \{ \underline{x}^{(i)}, y^{(i)} \}_{i=1}^{N}$ The choice of 'B' is mainly guided by Result: B' base models computational constraints for b = 1, ..., B do - Generate a bootstrap dataset  $\widetilde{T}^{(b)} = \{ \widetilde{\mathbb{Z}}^{(i)}, \widetilde{y}^{(i)} \}_{\dots}^{N}$ - Learn a base model from T(b) end Obtain  $\hat{y}_{bag}(\underline{x})$  by averaging:  $\hat{y}_{bag}(\underline{x}) = \frac{1}{R} \sum_{i=1}^{\infty} \tilde{y}^{(b)}(\underline{x})$ · Prediction with the base models Data: 'B' base models and test input x\* Result: A prediction  $\hat{y}_{bag}(\underline{x}^*)$ Use same formula