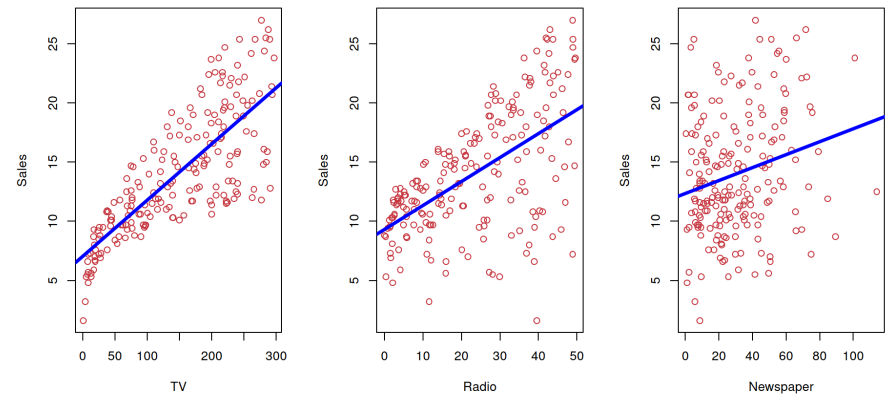
# **Statistical Learning**

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y=某產品在200家商店的銷售量

### What is Statistical Learning?



Shown are *Sales* vs. *TV*, *Radio* and *Newspaper*, with a blue linear-regression line fit separately to each

Can we predict Sales using these three? Perhaps we can do better using a model Sales ≈ f(TV, Radio, Newspaper)

2

## Notation

- Here, *Sales* is a response, dependent variable, or target that we wish to predict.
  We generically refer to the response as Y
- *TV* is a feature, independent variable, input, or predictor; we name it  $X_1$ . Likewise, name *Radio* as  $X_2$ , and so on
  - We can refer to the input vector collectively as

 $X = (X_1, X_2, X_3)$ 

Now, we write our model as

 $Y = f(X) + \epsilon$ 

where  $\epsilon$  captures measurement errors and other discrepancies and has a mean of zero

## Notation

Vectors are represented as a column vector

$$X_1 = \begin{pmatrix} X_{11} \\ X_{21} \\ \vdots \\ X_{n1} \end{pmatrix}$$

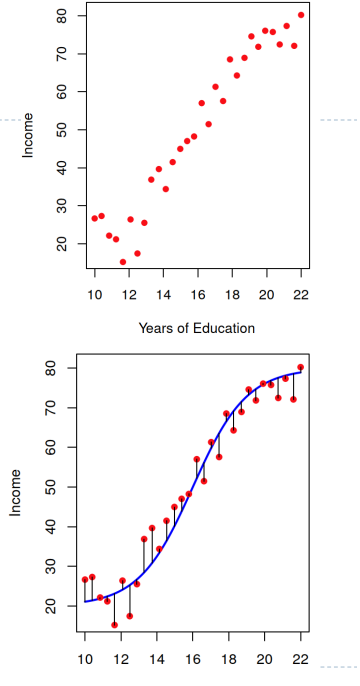
- We will use *n* to represent the number of distinct data points or observations
- We will let *p* denote the number of variables that are available for predictions
  - A general *design matrix* or input matrix can be written as an  $n \times p$  matrix

$$\begin{pmatrix} x_{11} & \cdots & x_{1p} \\ \vdots & \ddots & \vdots \\ x_{n1} & \cdots & x_{np} \end{pmatrix}$$

▶ *Y* is usually a scalar in our example; if we have *n* observations, it can be written as

## What is f(X) good for?

- With a good *f*, we can make predictions of *Y* at new points X = x
  - We can understand which components of  $X = (X_1, X_2, ..., X_p)$  are <u>important</u> in explaining Y and which are irrelevant. e.g., *Seniority* and *Years of Education* have a big impact on *Income*, but *Marital Status* typically does not
  - Depending on the complexity of *f*, we may be able to understand how each component X<sub>i</sub> of X affects Y
- In essence, statistical learning refers to <u>a set of</u> <u>approaches for estimating f</u>



Years of Education

## Why estimating f

• <u>Prediction</u>: In many situations, a set of inputs *X* are readily available, but the output *Y* cannot be easily obtained; we can then use  $\hat{f}$  as follows

$$\hat{Y} = \hat{f}(X)$$

- In this setting,  $\hat{f}(X)$  is often treated as a black box
- There will be reducible and irreducible error
  - *Reducible error* can be potentially improved by using the most appropriate statistical learning technique to estimate *f*
  - *Irreducible error* may contain unmeasured variables that are useful in predicting *Y*: since we don't measure them, *f* cannot use them for its prediction. It may also include unmeasurable variation
- We will focus on the part of the <u>reducible error</u>

## Why estimating f

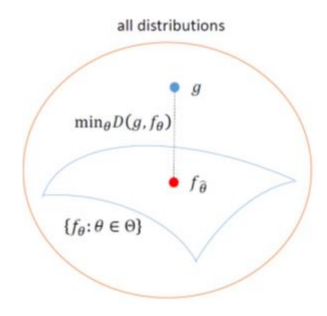
- Inference: We are often interested in understanding the association between Y and X<sub>1</sub>, ..., X<sub>p</sub>. In this situation, we wish to estimate f, but our goal is not necessarily to make predictions for Y
  - Which predictors are associated with the response?
  - What is the relationship between the response and each predictor?
  - Can the relationship between Y and each predictor be adequately summarized using a linear equation, or is the relationship more complicated?
- We will see a number of examples that fall into the prediction setting, the inference setting, or a combination of the two

# How to estimating f

- g is the distribution of data that is unknown
  - We have training set  $\{(x_1, y_1), \dots, (x_n, y_n)\}$
- 1. Choose a model  $f_{\theta}$ 
  - Parametric

8

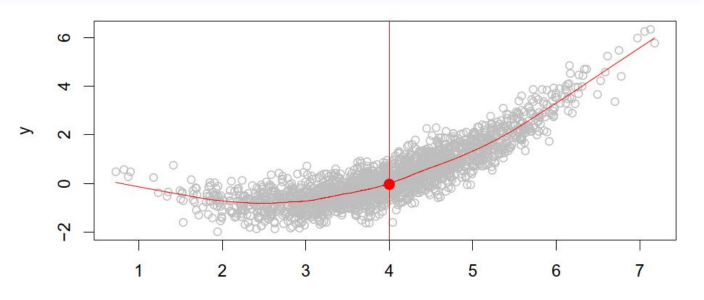
- Explicit assumption
- Estimating a fixed set of parameters by *fitting* or *training*
- Non-parametric
  - No explicit assumption
  - Need a large number of observations
- 2. Choose a quality measure (objective function) for fitting
  - Mean square error (Likelihood)...
- 3. Optimization (fitting) to choose the best  $\theta$ 
  - Calculus to find close form solution, gradient descent, expectation-maximization...



## Supervised vs Unsupervised learning

- Supervised Learning problem
  - In the <u>regression problem</u>, *Y* is quantitative (e.g., price, blood pressure)
  - In the <u>classification problem</u>, *Y* takes values in a finite, unordered set (survived/died, digit 0-9, cancer class of tissue sample)
  - We have training data (x<sub>1</sub>, y<sub>1</sub>), ..., (x<sub>n</sub>, y<sub>n</sub>). These are observations (examples, instances) of these measurements
- Unsupervised Learning problem
  - No outcome variable, just a set of predictors (features) measured on a set of samples
  - Objective is fuzzier find groups of samples that behave similarly, find features that behave similarly, find linear combinations of features with the most variation
- Semi-supervised learning problem
  - Only for *m* of the observations (m < n) that we have the response

## The regression problem



Is there an ideal f(X)? In particular, what is a good value for f(X) at any selected value of X, say X = 4? There can be many Y values at X = 4. A good value is

$$f(4) = E(Y|X=4)$$

E(Y|X = 4) means the expected value (average) of Y given X = 4. This ideal f(x) =
 E(Y|X = x) is called the regression function
 10

The regression function f(x)

#### • Also defined for vector X; e.g. $f(x) = f(x_1x_2, x_2) = E(Y | X_1 = x_1, X_2 = x_2, X_3 = x_3)$

- The ideal or optimal predictor of Y with regard to mean-squared prediction error: f(x) = E(Y|X = x) is the function that minimizes  $E[(Y f(X))^2|X = x]$  over all functions f at all points X = x
- $\epsilon = Y f(x)$  is the <u>irreducible</u> error i.e., even if we knew f(x), we would still make errors in prediction, since at each X = x, there is typically a distribution of possible Y values

# For any estimate $\hat{f}(x)$ of f(x), we have $E\left[\left(Y - \hat{f}(x)\right)^2 \middle| X = x\right] = E[f(x) + \epsilon - \hat{f}(x)]^2 = [f(x) - \hat{f}(x)]^2 + Var(\epsilon)$

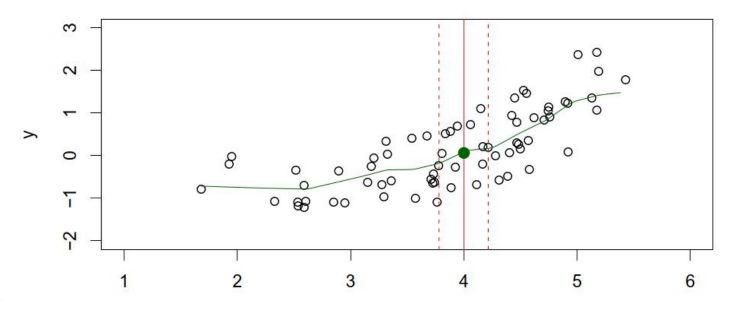
#### How to estimate f

• Typically, we have few if any data points with X = 4 exactly!

- So that we cannot compute E(Y|X = x)!
- Relax the definition and let

 $\hat{f}(x) = Ave(Y | X \in N(x))$ 

where N(x) is some <u>neighborhood</u> of x.

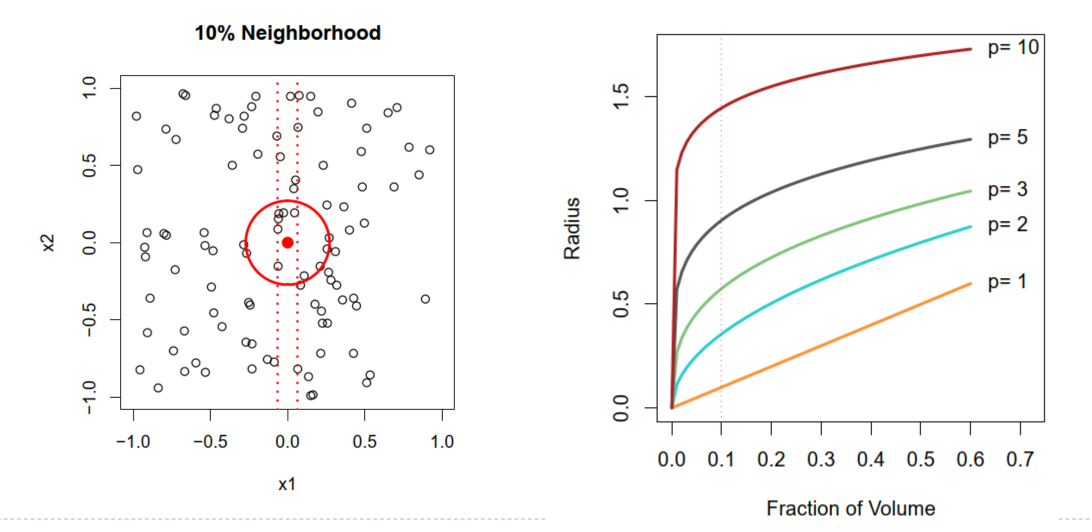


## The curse of dimensionality...

• Nearest neighbor averaging can be good for small  $p (p \le 4)$  and large n

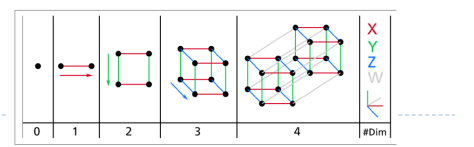
- We will discuss smoother versions, such as kernel and spline smoothing later in the course
- Nearest neighbor methods can be lousy when p is large. Reason: the <u>curse of</u> <u>dimensionality</u>. Nearest neighbors tend to be far away in high dimensions.
  - We need to get a reasonable fraction of the n values of y<sub>i</sub> to average to bring the variance down e.g., 10%
  - A 10% neighborhood in high dimensions need no longer be local, so we lose the spirit of estimating E(Y|X = x) by local averaging

#### The curse of dimensionality



14

## The curse of dimensionality



https://www.oreilly.com/library/view/hands-on-machine-learning/9781492032632/

p	1	2	3	4	5	6
(a) <u>Ball with</u> <u>radius R</u>	2R	$\pi R^2$	$\frac{4}{3}\pi R^3$	$\frac{\pi^2}{2}R^4$	$\frac{8\pi^2}{15}R^5$	$\frac{\pi^3}{6}R^6$
(b) Volume of hypercube 2 <sup>p</sup>	2	4	8	16	32	64
r = (a)/(b)	R	$\frac{\pi R^2}{4}$	$\frac{\pi R^3}{6}$	$\frac{\pi^2 R^4}{32}$	$\frac{\pi^2 R^5}{60}$	$\frac{\pi^3 R^6}{384}$

 $r = \frac{\pi^{\frac{p}{2}}}{2^{p}\Gamma(\frac{p}{2}+1)}R^{p}$ , it turns out that if we want to cover a fraction of r of the hypercube, we will need a ball with a radius  $\frac{2}{\pi^{\frac{1}{2}}}[r\Gamma(\frac{p}{2}+1)]^{\frac{1}{p}}$  (note that  $\Gamma(\frac{p}{2}+1) \sim \sqrt{\pi p} \left(\frac{p}{2e}\right)^{\frac{p}{2}}$ ) See chapter 2 of Foundations of Data Science

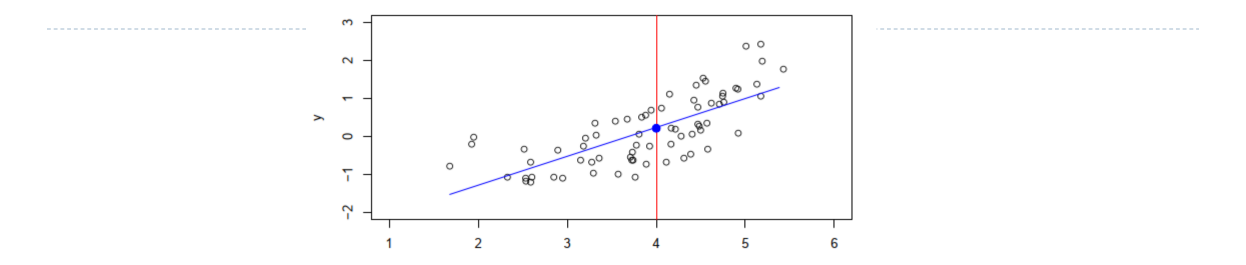
## Parametric and structured models

• The linear model is an important example of a parametric model:

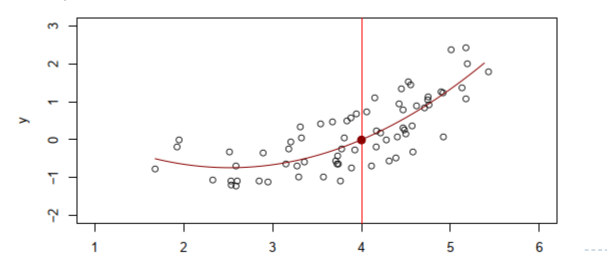
 $f_L(X) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_p X_p$ 

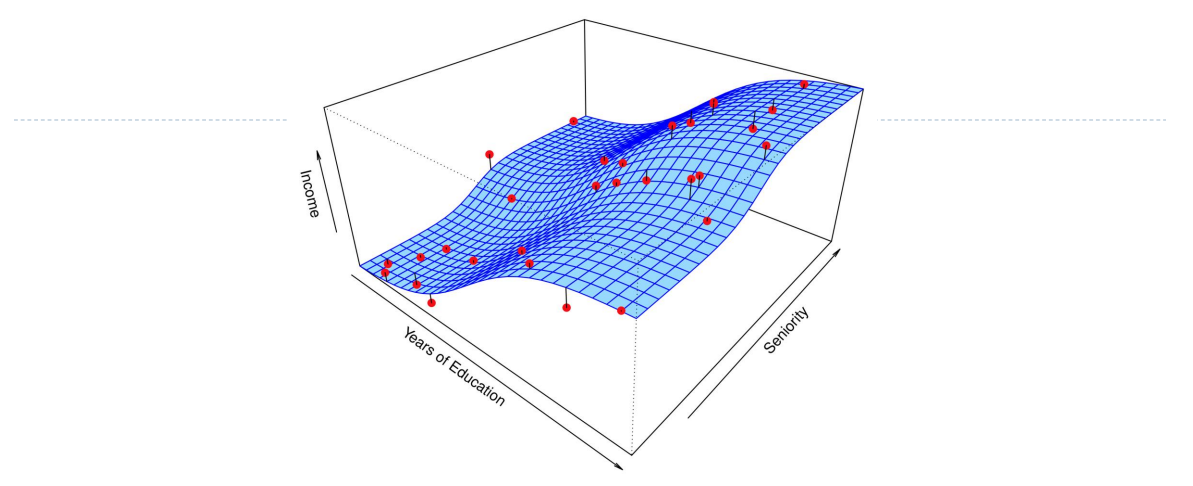
- A linear model is specified in terms of p + 1 parameters  $\beta_0, \beta_1, \dots, \beta_p$
- We estimate the parameters by <u>fitting</u> the model to training data
- Although it is almost never correct, a linear model often serves as a good and <u>interpretable</u> approximation to the unknown true function f(X)

A linear model  $f_L(X) = \beta_0 + \beta_1 X$  gives a reasonable fit here



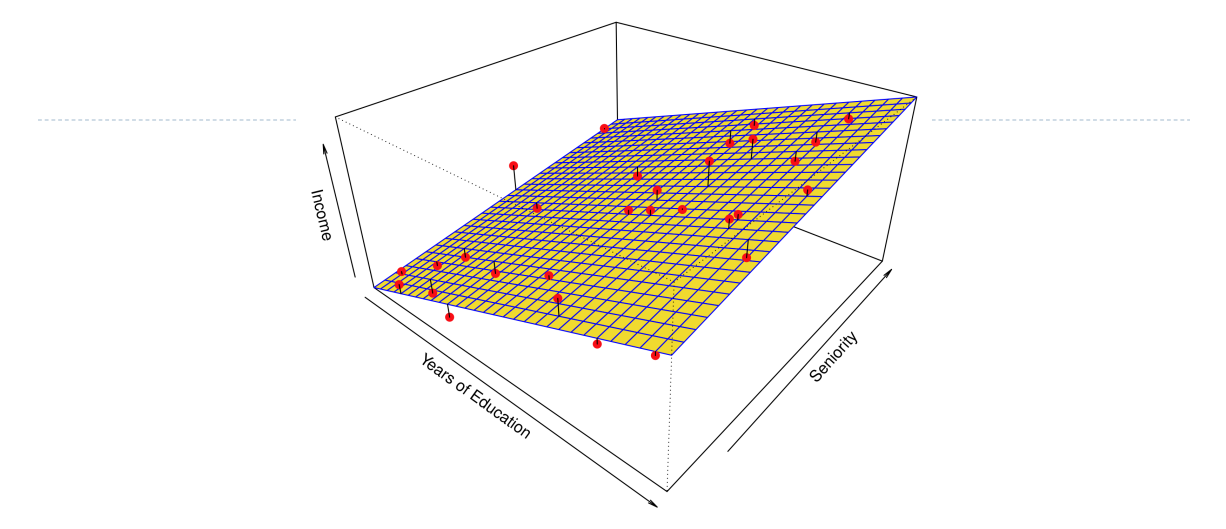
• A quadratic model  $f_Q(X) = \beta_0 + \beta_1 X + \beta_2 X^2$  fits slightly better



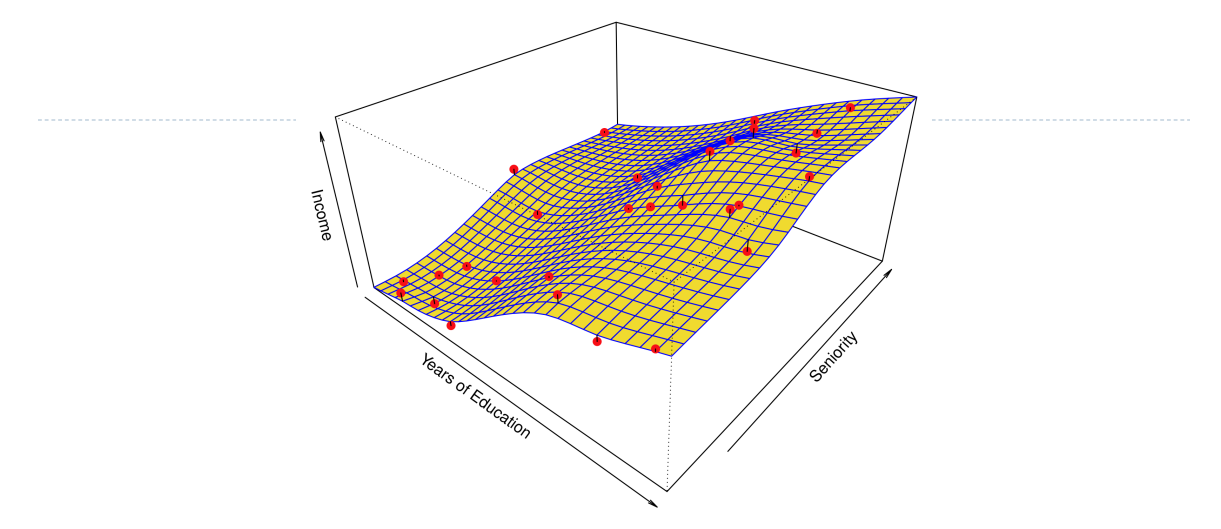


Simulated example. Red points are simulated values for income from the model

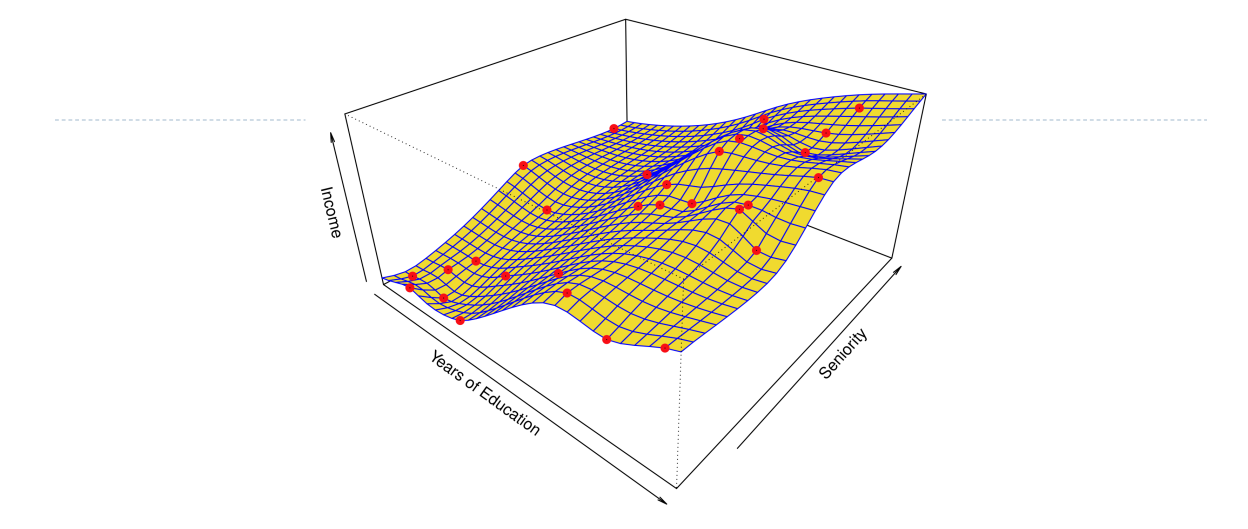
 $income = f(education, seniority) + \epsilon$ f is the blue surface



• Linear regression model fit to the simulated data  $\hat{f}_L(education, seniority) = \hat{\beta}_0 + \hat{\beta}_1 \times education + \hat{\beta}_2 \times seniority$ 



• More flexible regression model  $\hat{f}_s(education, seniority)$  fit to the simulated data. Here, we use a technique called a <u>thin-plate spline</u> to fit a flexible surface. We control the roughness of the fit (chapter 7)

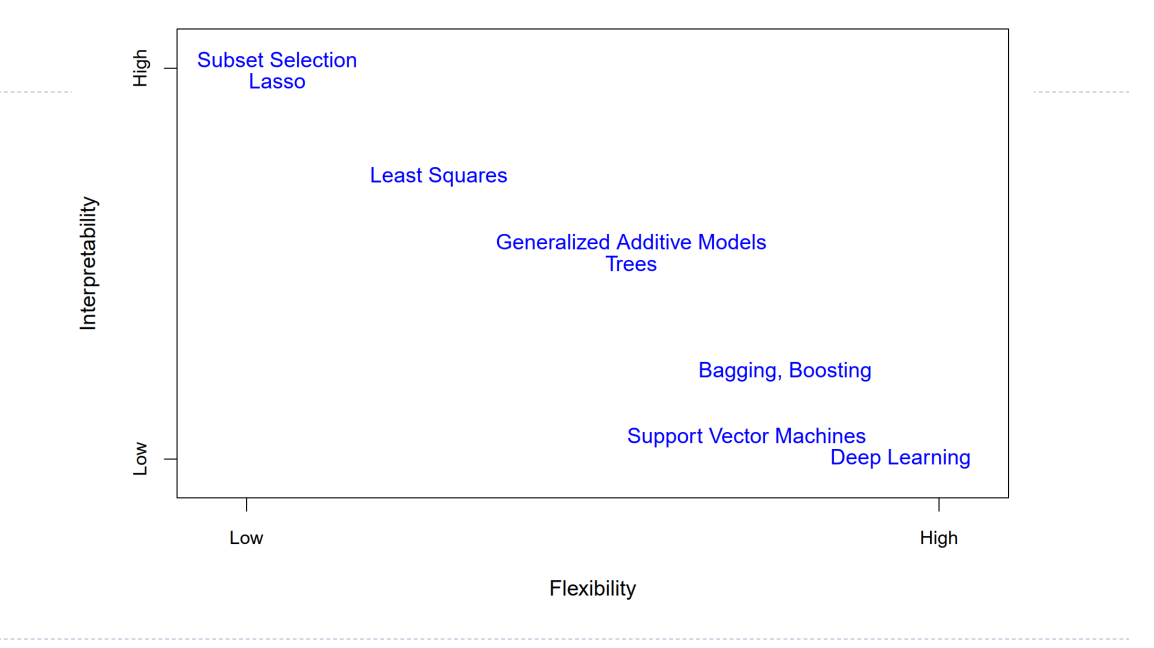


• Even more flexible *spline regression model*  $\hat{f}_s(education, seniority)$  fit to the simulated data. Here the fitted model makes no errors on the training data! Also known as <u>overfitting</u>

## Some trade-offs

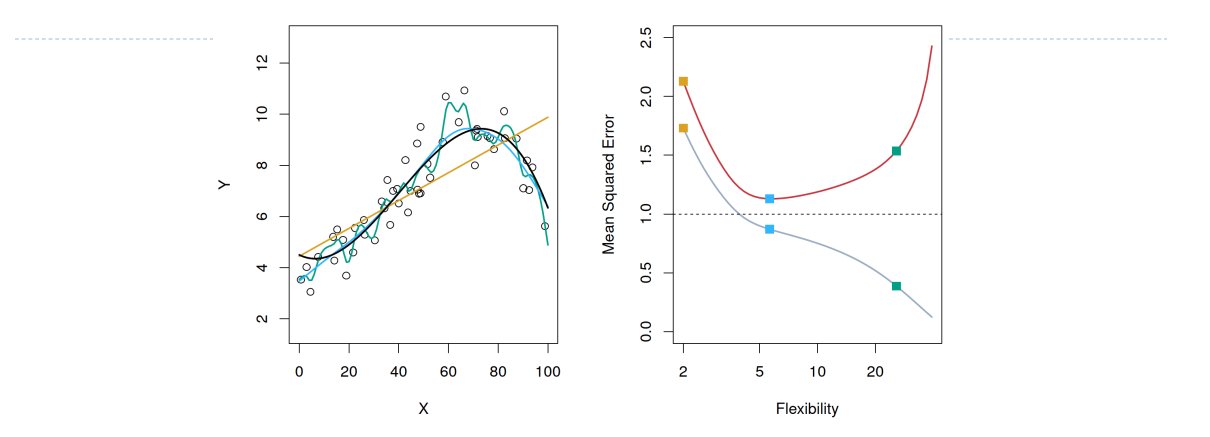
Prediction accuracy versus interpretability

- Linear models are easy to interpret; thin-plate splines are not
- Good fit versus over-fit or under-fit
  - How do we know when the fit is just right?
- Parsimony versus black-box
  - We often prefer a simpler model involving fewer variables over a black-box predictor involving them all

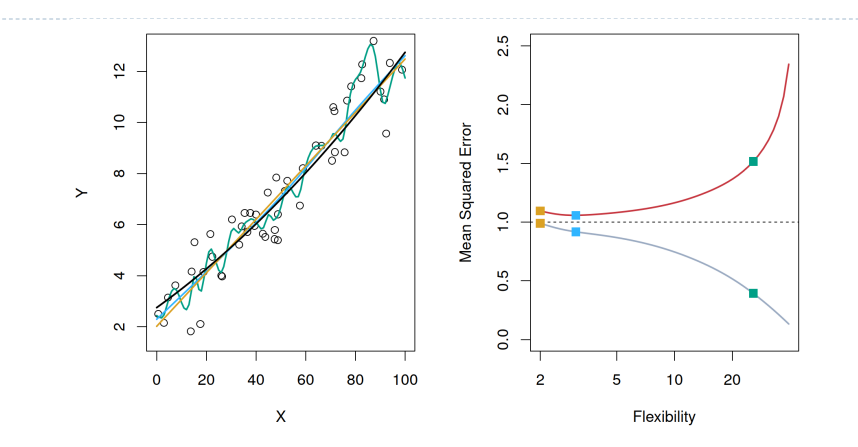


### Assessing Model Accuracy

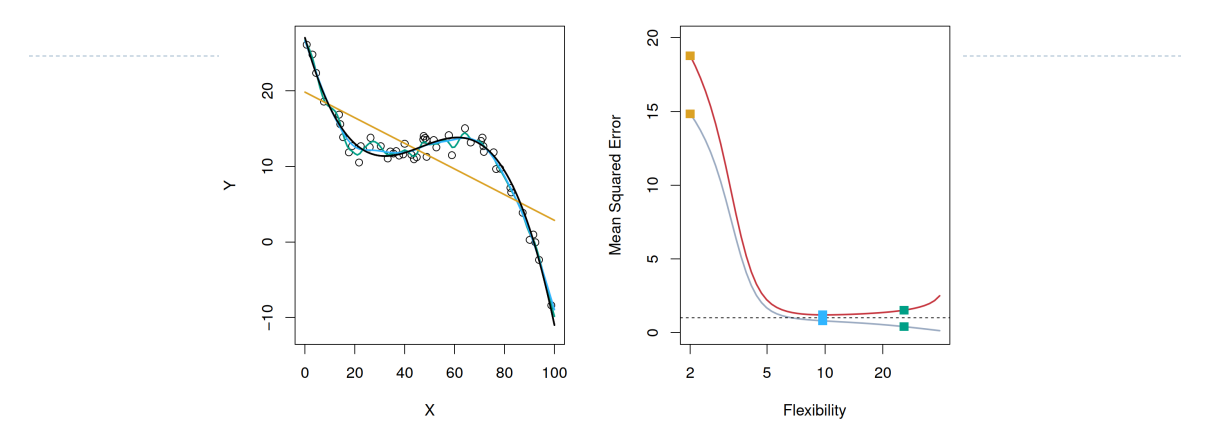
- Suppose we fit a model f(x) to some training data Tr = {x<sub>i</sub>, y<sub>i</sub>}, i = 1 ... n, and we wish to see how well it performs
  - We could compute the average squared prediction error over Tr:  $MSE_{Tr} = Ave_{i \in Tr}[y_i - \hat{f}(x_i)]^2$
- This may be biased toward more overfit models
  - Instead, we should, if possible, compute it using fresh test data  $Te = \{x_i, y_i\}, i = 1 \dots m$ ,  $MSE_{Te} = Ave_{i \in Te} [y_i - \hat{f}(x_i)]^2$



• The black curve is truth. Red curve on the right is  $MSE_{Te}$ , grey curve is  $MSE_{Tr}$ . Orange, blue and green curves/squares correspond to fits of different flexibility



• Here, the truth is smoother, so the smoother fit and linear model do really well

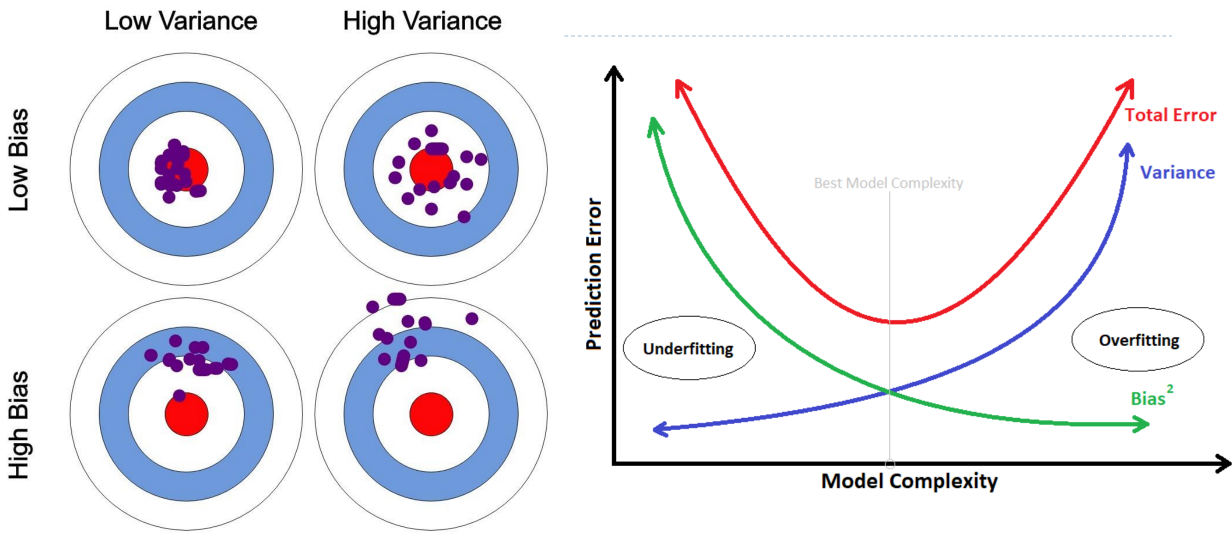


Here, the truth is wiggly and the noise is low, so the more flexible fits do the best

• Proof of testing error is usually larger than the training error

#### **Bias-Variance Trade-off**

- Suppose we have fit a model  $\hat{f}(x)$  to some training data Tr, and let  $(x_0, y_0)$  be a test observation drawn from the population. If the true model is  $Y = f(X) + \epsilon$  (with f(x) = E(Y|X = x)), then  $E\left[(y_0 - \hat{f}(x_0))^2\right] = Bias_{Tr}[\hat{f}(x_0, Tr)]^2 + Var_{Tr}[\hat{f}(x_0, Tr)] + Var(\epsilon)$
- The expectation averages over the variability of  $y_0$  as well as the variability in Tr. Note that  $Bias_{Tr}[\hat{f}(x_0, Tr)] = E[\hat{f}(x_0, Tr)] f(x_0)$ 
  - Typically, as the *flexibility* of  $\hat{f}$  increases, its variance increases, and its bias decreases. So choosing the flexibility based on average test error amounts to a *bias-variance trade-off*
  - Proof of the decomposition

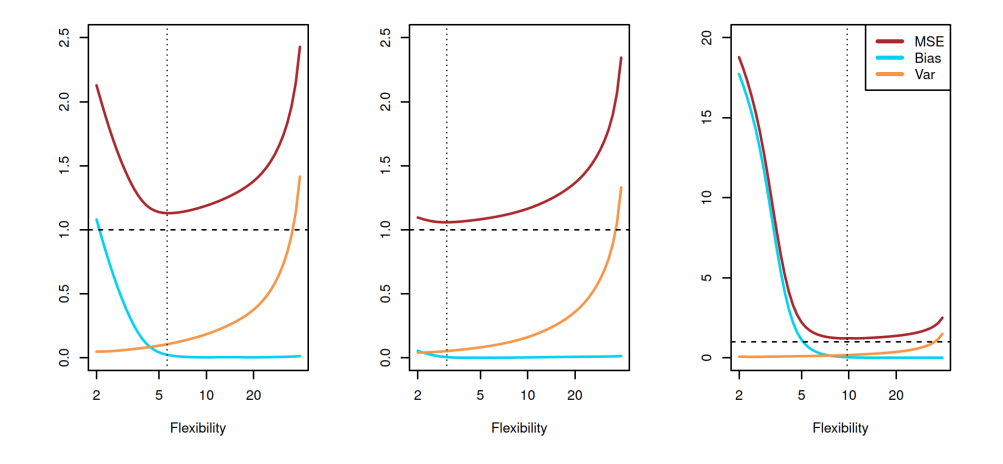


https://nvsyashwanth.github.io/machinelearningmaster/bias-variance/

https://jason-chen-1992.weebly.com/home/-bias-variance-tradeoff

29

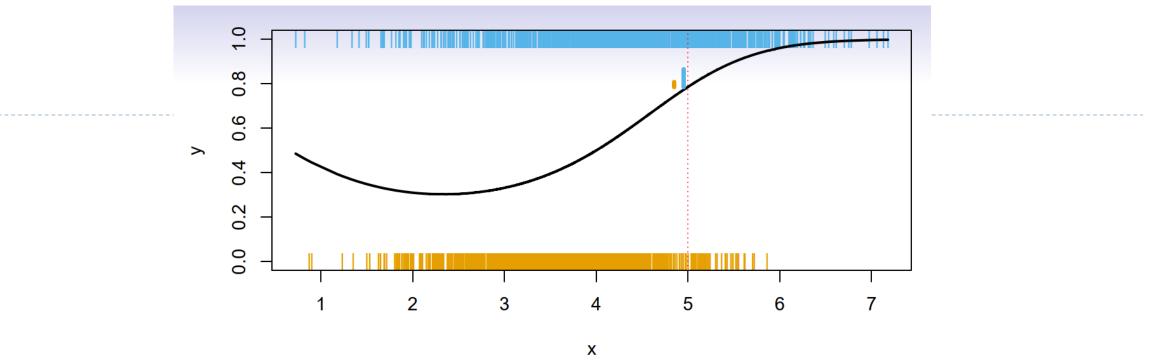
## Bias-variance trade-off for the three examples



30

## **Classification Problems**

- Here, the response variable Y is qualitative e.g. email is one of C = (spam, ham) (ham = good email), digit class is one of C = {0, 1, ..., 9}. Our goals are to:
  - Build a classifier C(X) that assigns a class label from C to a future unlabeled observation X
  - What is an optimal classifier?
  - Understand how flexibility affects the classification

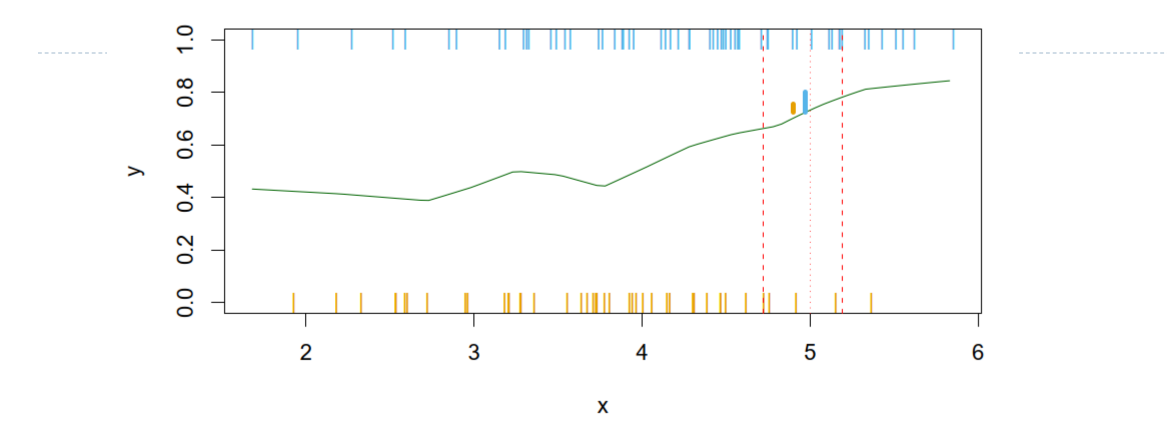


The orange/blue marks indicate the response *Y*, either 0 or 1

Is there an ideal C(X)? Suppose the K elements in C are numbered 1, 2, ..., K.

$$p_k(x) = \Pr(Y = k | X = x), k = 1, 2, \dots, K.$$

• These are the *conditional class probabilities* at x; e.g., see the little barplot at x = 5. Then the <u>Bayes optimal classifier</u> at x is C(x) = j if  $p_j(x) = \max\{p_1(x), p_2(x), \dots, p_k(x)\}$ 



Nearest-neighbor averaging can be used as before. It also breaks down as the dimension grows. However, the impact on  $\hat{C}(x)$  is less than on  $\hat{p}_k(x), k = 1, \dots, K$ 

## Classification: some details

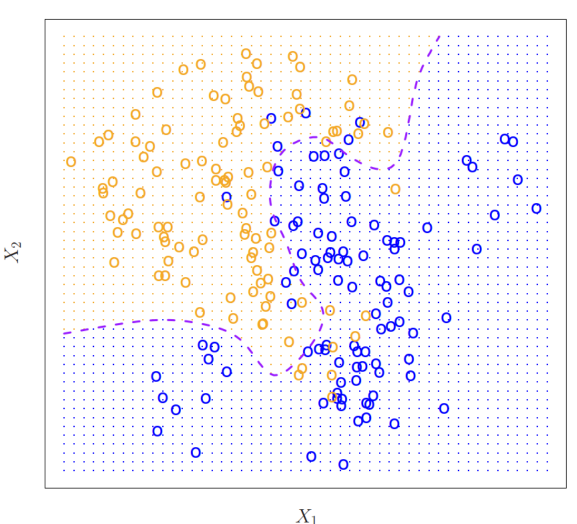
Typically, we measure the performance of Ĉ(x) using the misclassification error rate:

$$Err_{Te} = Ave_{i \in Te}I[y_i \neq \hat{C}(x_i)]$$

- The <u>Bayes classifier</u> (using the true  $\hat{p}_k(x)$ ) has the smallest error (in the population)
- Support-vector machines build structured models for C(x)
- We will also build structured models for representing the  $p_k(x)$ . e.g., *Logistic regression*, generalized additive models

 The Bayes classifier produces the lowest possible test error rate, called the <u>Bayes error rate</u>

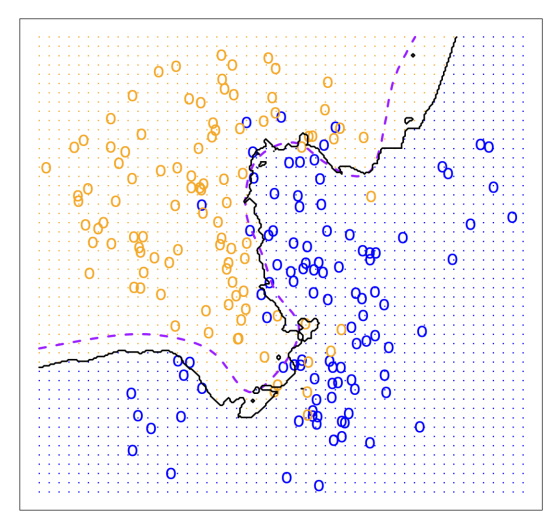
 $1 - max_j \Pr(Y = j | X = x_0)$ 



 K-nearest neighbors (KNN) classifier

•  $\Pr(Y = j | X = x_0) = \frac{1}{K} \sum_{i \in Tr} I(y_i = j)$ 

KNN: K=10

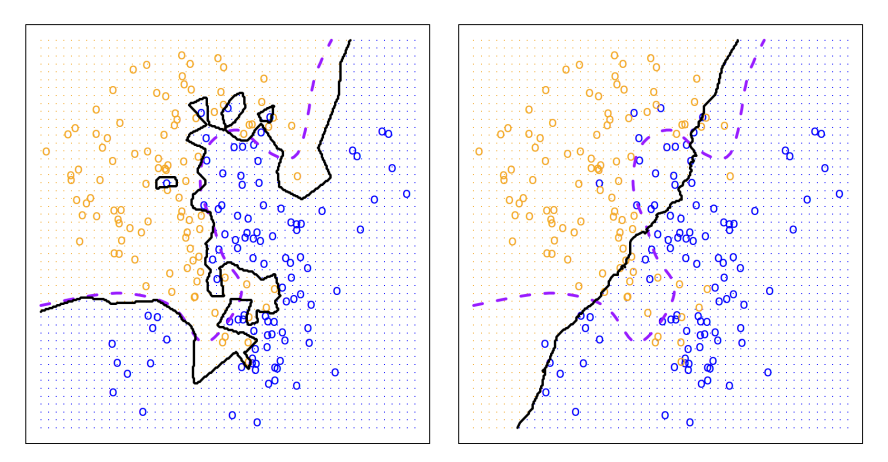


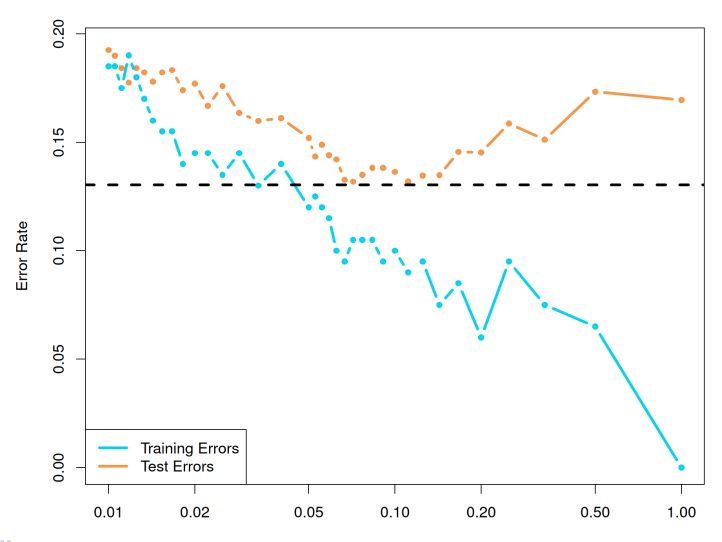
 $X_1$ 

 $X_2$ 

KNN: K=1

KNN: K=100





38

1/K

# Appendix

#### The Bias-variance tradeoff

• 
$$f = f(x), \hat{f} = \hat{f}(x, Tr), Var(X) = E(X^2) - E[X]^2$$

y = f + ε → E(y) = E(f) = f (f is deterministic, independent of Tr and f̂ is independent of ε)

•  $MSE = E_x[Bias_{Tr}[\hat{f}(x,Tr)]^2 + Var_D[\hat{f}(x,Tr)]] + \sigma^2$  (Taking expectation over x)

