# Statistical Learning and Prediction

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### **Statistical Learning**

Vast set of tools for understanding data.

- − Supervised:  $Y \leftarrow f(X_1, ..., X_p)$ ; predict Y on the basis of X
- Unsupervised:  $X_1, \ldots, X_p$ ; finding structure in X (underlying dimensions/groups)

### **Statistical Learning**

• The linear regression model

$$Y = f(X_1, \dots, X_p) + \epsilon$$
$$= \beta_0 + \beta_1 X_1 + \dots + \beta_p X_p + \epsilon$$

can be used for explanation and/or prediction.

- Explanation: understanding how the X's are related to Y; possibly causally.
- <u>Prediction</u>: if we have new observations with known values of *X*'s, what is the expected (predicted) value of *Y* and how accurate are these predictions?

### **Explanatory Regression**

Suppose we have data and obtained estimates:

$$\hat{y}_i = 2 + 0.5x_{i1} + 1.5x_{i2}$$

- Estimated coefficients indicate magnitude of the effects, standard errors indicate variability of estimates.
- Use statistical tests for deciding whether the explanatory variables really affect the response, given the other variables.
- Adequate estimation of coefficients  $(\hat{\beta})$  is assumed crucial: Accurate estimates = unbiased estimates! That is:

$$\mathbb{E}[\hat{\beta}] = \beta$$

### **Explanatory Regression**

• Unbiased estimates can be obtained with e.g., OLS:

$$\hat{\beta} = \arg\min_{\beta} \left( \sum_{i=1}^{N} (y_i - x_i^{\top} \beta)^2 \right)$$

• "From a Bayesian perspective, the principle of unbiasedness is reasonable in the limit of large samples, but otherwise it is potentially misleading" (Gelman et al., 1995)

### **Predictive Regression**

• Suppose we have data and have obtained estimates:

$$\hat{y}_i = 2 + 0.5x_{i1} + 1.5Xx_{i2}$$

- Suppose we have a new observation  $x_0 = \begin{bmatrix} 2 & 3 \end{bmatrix}$
- With these values we can predict Y, i.e.,  $2 + 0.5 \times 2 + 1.5 \times 3 = 7.5$
- We do not care to recover parameters that generated the data, but want to obtain a model that yields as accurate as possible  $\hat{Y}$ .
- E.g., minimize

$$\mathbb{E}[(\hat{Y} - Y)^2]$$

#### Exercise 1: Bias can be beneficial

• Generate some training observations and combine them into a training dataset. Use set.seed() to allow for later replication:

```
x <- runif(50, min = -3, max = 3)
epsilon <- rnorm(50)
y <- 0.1*x + epsilon
train_dat <- data.frame(x, y)</pre>
```

- Generate 1,000 new observations from the same distributions. Combine the observations into a test dataset.
- Generate a range of shrinkage factor values:

```
s \leftarrow seq(0, 1, by = .1)
```

• Estimate an OLS regression on the training observations, using 1m. Omit the intercept, by adding 0 or −1 in the right-hand side

of the model formula. Extract the estimated  $\beta$  coefficient using coef.

• Compute predictions for the test observations, but do not use function predict. Instead, compute predictions yourself and apply shrinkage to the OLS coefficient you just estimated:  $\hat{Y} = X \cdot s \cdot \hat{\beta}$ .

Thus, generate predictions for the test observations 11 times: once for each value of the shrinkage factor *s*.

- For each value of *s*, compute MSE (mean squared prediction error) on the test observations.
- Plot the test MSE values as a function of shrinkage factor *s*. Is shrinkage beneficial for prediction? What is the optimal value for shrinkage factor *s*?

- Repeat the above experiment 100 times, and plot the average MSE (over the 100 replications) as a function of shrinkage *s*.
- What do you expect to happen to the *optimal* value of the shrinkage if sample size doubles (i.e.,  $N_{train} = 100$  instead of 50)? And if effect size doubles (i.e., the effect of x is .20 instead of .10)?

### **Predictive Regression**

- Traditional statistical textbooks focus on obtaining *unbiased* estimates (e.g., OLS, ML).
- (Modern) statistical learning accepts biased parameter estimates as long as *variance* decreases more than *squared bias* increases.

### Predictive perspective: Bias-variance trade-off

- We have a probability distribution  $P^*$ , from which we draw a sample  $\mathcal{T}$  of size N
- Let  $f_B$  be Bayes optimal f (unknown, depends on  $P^*$ )
- Let  $\bar{f}(X) = \mathbb{E}_{\mathcal{T}}[\hat{f}(X)]$
- Aim is to minimize *expected prediction error*:

$$\mathbb{E}_{\mathcal{T}}[\text{EPE}(\hat{f})] = \mathbb{E}_{X}[Var(Y|X)] +$$

$$\mathbb{E}_{X}[(f_{B}(X) - \bar{f}(X))^{2}] +$$

$$\mathbb{E}_{\mathcal{T}}\mathbb{E}_{X,Y}[(\hat{f}(X) - \bar{f}(X))^{2}]$$

• EPE = irreducible error (or  $\sigma^2$ ) + bias<sup>2</sup> + variance

### **Population: Non-linear Regression**

Often we fit a linear model, assuming that the conditional means in the population lie on a straight line.

This assumption is most likely false! Why does it often work so well?

# Excercise 2: Under- and overfitting with polynomial regression

• Generate a training and test set, each with n = 50, with a single predictor which has a non-linear effect on the response:

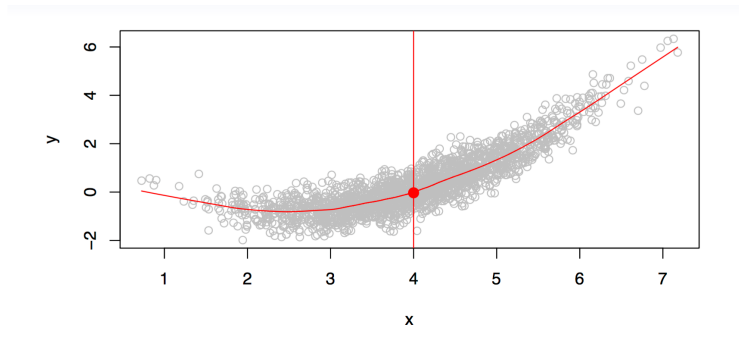
```
set.seed(42)
n <- 50
x <- runif(n, min = -5, max = 5)
y <- x + 8*sin(x/2) + rnorm(n)
train <- data.frame(x, y)
x <- runif(n, min = -5, max = 5)
y <- x + 8*sin(x/2) + rnorm(n)
test <- data.frame(x, y)</pre>
```

• Fit polynomial regression models to the training data of degree 1 trough 15, make predictions on the test set and compute the

prediction error for each degree. (You can use predict for this exercise.)

- Hint: Use functions lm, poly, and a for loop. E.g.:  $lm(y \sim poly(x, degree = 3))$
- Plot the test MSE as a function of the polynomial degree.
- If time permits, create a plot which shows the training observations, and fitted curves for the degree 1, 2, 3 and 15 polynomials.

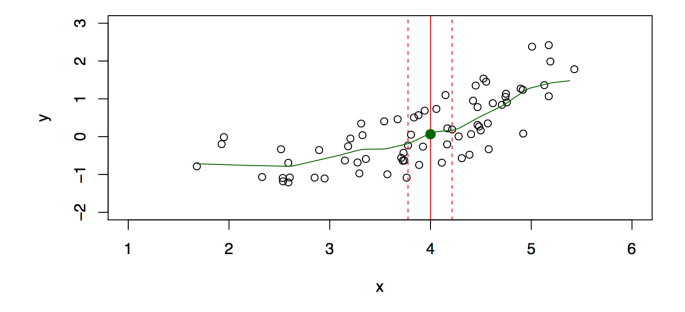
### **Population: Non-linear Regression**



The regression line in the <u>population</u> combines the conditional means at each point x

## Sample data

Using sample data, we want to obtain an estimate  $\hat{f}(X)$  of f(X).



- Due to sparsity, cannot estimate a conditional mean at all points X = x.
- $\bullet$  Thus, take a small neighbourhood around X = x and take

neighbourhood mean as predicted value, i.e. nearest neighbour averaging.

- How is the size of the neighbourhood defined in kNN?
- What happens to the *bias* if *k* increases?
- What happens to the *variance* ik *k* increases?
- Does the optimal value for *k* depend on sample size?
- Does the optimal value for *k* depend on the amount of irreducible error?

### Multiple Predictor Variables

With multiple predictors the observations are further spread out through the space:

- Nearest neighbours might not be near at every point
- Flexible models become very wild
- This is known as the *curse of dimensionality*
- More structure in *f* is needed
- How can we impose structure?

### **Exercise 3: Curse of dimensionality**

• Generate a dataset with 100 observations on 10,000 predictors. Predictors may be independent:

```
p <- 10000
N <- 100
set.seed(42)
X <- matrix(rnorm(p*N), ncol = p, nrow = N)</pre>
```

- Compute Euclidian distances between all points in the dataset. First, only use the first column of X to compute the distances (i.e., p = 1), then use the first two columns (i.e., p = 2), and so on for  $p \in \{1, 2, 10, 100, 1000, 10000\}$ .
- Create a histogram for each value of *p*.

- Hint: Use function dist to compute distances, use function hist to create a histogram. Specify argument xlim for each histogram, to make sure 0 is included on the *x*-axis.
- Are the nearest neighbours near in 1-dimensional space? In 2-dimensional space? In 10-, 100-, 1000-dimensional space?

### Evaluating predictive accuracy: Numeric outcome

• Mean squared error (MSE):

MSE = 
$$\frac{1}{N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

• Mean absolute error (MAE):

MAE = 
$$\frac{1}{N} \sum_{i=1}^{N} |y_i - \hat{y}_i|$$

### Classification

Response variable Y may be a categorical variable with categories C = 1, ..., k, ..., K.

Again, we want to predict response *Y* based on predictors *X*:

- Can directly construct a classifier  $\hat{f}(X) = C(X)$  that assigns a predicted category from C based on X.
- Preferable, though: Construct a function  $\hat{f}(X)$  that provides conditional probabilities:  $\hat{p}_k(X) = Pr(Y = k | X = x)$ .

Then Bayes classifier assigns C(X) = k if  $\hat{p}_k(x) = \max\{\hat{p}_1(x), \dots, \hat{p}_K(x)\}$ 

### **Evaluating predictive accuracy: Classification**

• Misclassification rate:

$$\frac{1}{N} \sum_{i=1}^{N} I(y_i \neq \hat{y}_i)$$

• Brier score:

$$\frac{1}{N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

• Cross-entropy (a.k.a. deviance):

$$-\frac{1}{N} \sum_{i=1}^{N} [y_i \cdot \log(\hat{y}_i) + (1 - y_i) \cdot \log(1 - \hat{y}_i)]$$

### Exercise 4: Flexibility and predictive performance

For each of parts (a) through (d), indicate whether we would generally expect the performance of a flexible statistical learning method to be better or worse than an inflexible method:

- a) Sample size *N* is extremely large, and the number of predictors *p* is small.
- b) The number of predictors *p* is extremely large, and the number of observations *N* is small.
- c) The relationship between the predictors and response is highly non-linear.
- d) The variance of the error terms,  $\sigma^2 = Var(\epsilon)$ , is extremely high.