STA 314: Statistical Methods for Machine Learning I

Lecture 9 - Moving beyond linearity, k-Nearest Neighbor

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Moving Beyond Linearity

The linearity assumption in the feature space is almost always an approximation, and sometimes a poor one.

We consider the following extensions to relax the linearity assumption.

- Univariate feature (p = 1):
 - Polynomial regression
 - Step functions
 - Regression splines
- Multivariate feature (p > 1):
 - Local regression
 - Generalized additive models

Polynomial Regression

The polynomial regression

$$y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \dots + \beta_d x_i^d + \epsilon_i,$$

where ϵ_i is the error term and $x_i \in \mathbb{R}$.

- Can be fitted by the OLS approach.
- Coefficients themselves are not interpretable; we are more interested in the trend of the fitted function

$$\hat{f}(x_0) = \hat{\beta}_0 + \hat{\beta}_1 x_0 + \hat{\beta}_2 x_0^2 + \dots + \hat{\beta}_d x_0^d.$$

• The degree *d* in practice is typically no greater than 4, and can be chosen via cross-validation.

Polynomial Regression

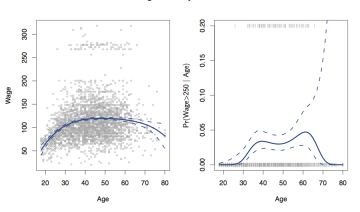
- The polynomial regression can be used for classification as well.
 - For instance, in the logistic regression,

logit (
$$\mathbb{P}(Y_i = 1 \mid X_i = x_i)$$
) = $\beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \dots + \beta_d x_i^d$.

- Can be fit by maximizing the likelihood.
- However, polynomials have notorious tail behavior very bad for extrapolation.

The Wage Data

Degree-4 Polynomial



Left: The solid blue curve is a degree-4 polynomial of wage as a function of age, fit by the OLS. The dotted curves are estimated 95 % confidence intervals. Right: We model the binary event 1{wage > 250} using logistic regression, with a

degree-4 polynomial.

Step Functions

- The polynomial regression imposes a global structure on the non-linearity of X.
- The **step function** approach avoids such a global structure by breaking the range of *X* into bins.
- For pre-specified K cut points c_1, \ldots, c_K , define

$$C_0(X) = 1\{X < c_1\},\$$
 $C_1(X) = 1\{c_1 \le X < c_2\},\$
 \vdots
 $C_K(X) = 1\{c_K \le X\}.$

 $C_0(X), \ldots, C_K(X)$ are in fact (K+1) dummy variables, and they sum up to 1.

Step Functions

• Step function approach assumes

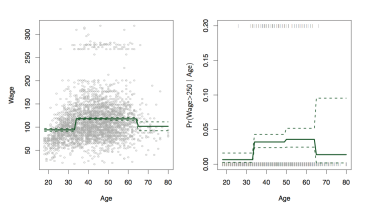
$$y_i = \beta_0 + \beta_1 C_1(x_i) + \beta_2 C_2(x_i) + ... + \beta_K C_K(x_i) + \epsilon_i$$

where ϵ_i is the error term. (Note we don't need $C_0(x_i)$ in the model.)

- Can be fitted by the OLS.
- β_j represents the average change in the response Y for $c_j \le X < c_{j+1}$ relative to $X < c_1$.

The Wage Data

Piecewise Constant



Left: The solid blue curve is a step function of wage as a function of age, fit by least squares. The dotted curves indicate an estimated 95~% confidence interval. Right: We model the binary event wage>250 using logistic regression, with the step function.

Pros and Cons of Step Function

- The step function approach is widely used in biostatistics and epidemiology among other areas, because the model is easy to fit and the regression coefficient has a natural interpretation.
- However, piecewise-constant functions can miss the trend of the true relationship between Y and X. The choice of cut points can be problematic.
- How about combining polynomial and step function?

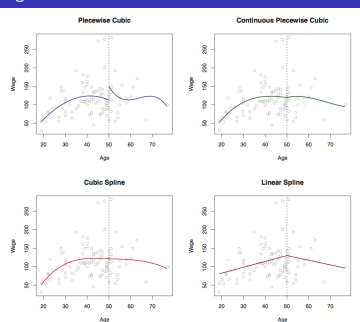
Piecewise Polynomials

• Instead of a single polynomial in *X* over its whole domain, we can use different polynomials in distinct regions:

$$y_i = \begin{cases} \beta_{01} + \beta_{11} x_i + \beta_{21} x_i^2 + \beta_{31} x_i^3 + \epsilon_i & \text{if } x_i < c; \\ \beta_{02} + \beta_{12} x_i + \beta_{22} x_i^2 + \beta_{32} x_i^3 + \epsilon_i & \text{if } x_i \ge c. \end{cases}$$

- The cut point c is called knot. Using more knots leads to a more flexible piecewise polynomial.
- In general, if we place K different knots throughout the range of X, then we will end up fitting (K+1) different cubic polynomials.

The Wage Data



Regression splines

- Better to add constraints to polynomials at the knots for:
 - continuity: equal function values
 - smoothness: equal first and second order derivatives
 - higher order derivatives
- The constrained polynomials are called **splines**. A degree-d spline contains piecewise degree-d polynomials, with continuity in derivatives up to degree (d-1) at each knot.
- How can we construct the degree-d spline?

Linear Splines

• A linear spline has piecewise linear functions continuous at each knot. That is, with knots at $\xi_1 < \xi_2 < \cdots < \xi_K$,

$$y_i = \beta_0 + \beta_1 x_i + \beta_2 (x_i - \xi_1)_+ \cdots + \beta_{K+1} (x_i - \xi_K)_+ + \epsilon_i$$

where

$$(x_i - \xi_k)_+ = \begin{cases} x_i - \xi_k & \text{if } x_i > \xi_k \\ 0 & \text{otherwise} \end{cases}.$$

A basis representation:

$$y_i = \beta_0 + \beta_1 b_1(x_i) + \beta_2 b_2(x_i) + \dots + \beta_{K+1} b_{K+1}(x_i) + \epsilon_i,$$

where b_k are basis functions

$$b_1(x_i) = x_i, \quad b_{k+1}(x_i) = (x_i - \xi_k)_+, \quad k = 1, \dots, K,$$

• Interpretation of β_1 : the averaged increase of Y associated with one unit of X for $X < \xi_1$.

Cubic Splines

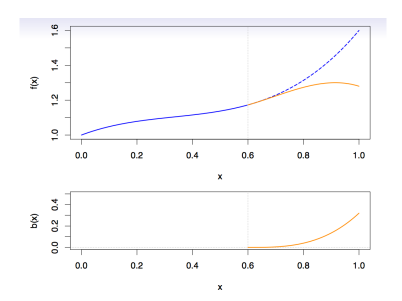
• A cubic spline has piecewise cubic polynomials with continuous derivatives up to order 2 at each knot. That is, with K knots at $\xi_1 < \xi_2 < \cdots < \xi_K$,

$$y_i = \beta_0 + \beta_1 b_1(x_i) + \beta_2 b_2(x_i) + \dots + \beta_{K+3} b_{K+3}(x_i) + \epsilon_i,$$

where b_k are basis functions

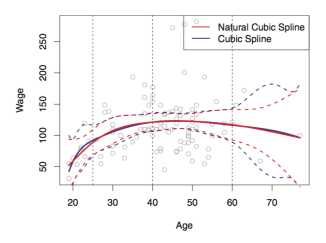
$$b_1(x_i) = x_i,$$
 $b_2(x_i) = x_i^2,$ $b_3(x_i) = x_i^3,$
 $b_{k+3}(x_i) = (x_i - \xi_k)_+^3,$ $k = 1, ..., K.$

Cubic Splines



Natural Splines

A natural spline is a regression spline with additional boundary constraints: the function is required to be linear at the boundary.



More on splines

- Choosing the number and locations of the knots
 - ▶ Typically, we place *K* knots at the corresponding quantiles of the data or place on the range of *X* with equal space. Oftentimes, the placement of knots is not very crucial.
 - ▶ We use cross-validation to choose K.
- Polynomial regressions and step functions are special cases of splines.
- Another variant: smoothing spline (ISLR 7.5).

Local Regression

Local regression predicts at a target point x_0 using only the nearby training observations.

Algorithm 7.1 Local Regression At $X = x_0$

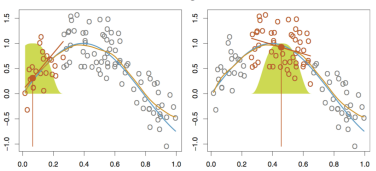
- 1. Gather the fraction s = k/n of training points whose x_i are closest to x_0 .
- 2. Assign a weight $K_{i0} = K(x_i, x_0)$ to each point in this neighborhood, so that the point furthest from x_0 has weight zero, and the closest has the highest weight. All but these k nearest neighbors get weight zero.
- 3. Fit a weighted least squares regression of the y_i on the x_i using the aforementioned weights, by finding $\hat{\beta}_0$ and $\hat{\beta}_1$ that minimize

$$\sum_{i=1}^{n} K_{i0} (y_i - \beta_0 - \beta_1 x_i)^2.$$
 (7.14)

4. The fitted value at x_0 is given by $\hat{f}(x_0) = \hat{\beta}_0 + \hat{\beta}_1 x_0$.

Simulated Example

Local Regression



The blue curve is true f(x), and the light orange curve is the local regression $\hat{f}(x)$. The orange points are local to the target point x_0 , represented by the orange vertical line. The yellow bell-shape indicates weights assigned to each point. The fit $\hat{f}(x_0)$ at x_0 is obtained by fitting a weighted linear regression (orange line segment), and using the fitted value at x_0 (orange solid dot) as the estimate $\hat{f}(x_0)$.

Local Regression

- The size of the neighborhood (fraction s of training data) is a tuning parameter, which can be chosen by cross-validation.
- When we have two dimensional predictors x_1 and X_2 , we can simply use 2-dimensional neighborhoods, and fit bivariate linear regression models using the observations that are near each target point in 2-dimensional space.
- However, local regression can perform poorly if p is much larger than about 3 or 4 (the curse of dimensionality).
- *k*-Nearest Neighbour is one of the most common local regression approaches. (Later)

Generalized Additive Models

 Generalized additive models (GAMs) provide a general framework for extending a standard linear model by allowing non-linear functions of each of the variables, while maintaining additivity,

$$y_i = \beta_0 + f_1(x_{i1}) + f_2(x_{i2}) + \dots + f_p(x_{ip}) + \epsilon_i.$$

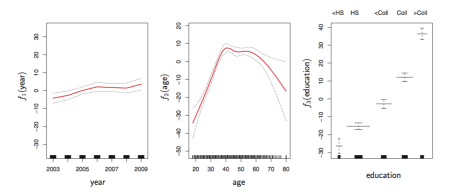
- Each f_k can be linear, polynomials, step function, splines and local regression.
- Can be applied to classification problems.
 - Logistic regression:

logit (
$$\mathbb{P}(Y_i = 1 \mid X_i = x_i)$$
) = $\beta_0 + f_1(x_{i1}) + f_2(x_{i2}) + \cdots + f_p(x_{ip})$.

Wage Data

Consider the wage data

wage =
$$\beta_0 + f_1(\text{year}) + f_2(\text{age}) + f_3(\text{education}) + \epsilon$$
.



The first two functions are natural splines in year and age. The third function is a step function, fit to the qualitative variable education.

Pros and Cons of GAMs

- GAMs allow us to fit a non-linear f_j to each X_j , so that we can automatically model non-linear relationships that standard linear regression won't be able to capture.
- The non-linear fits can potentially make more accurate predictions for the response *Y*.
- Because the model is additive, we can still examine the effect of each X_i on Y individually while holding all of the other variables fixed.
- It avoids the curse of dimensionality.
- However, GAMs fail to incorporate the interaction of variables.

A Classical Local Approach: Nearest Neighbors

- Suppose we're given a new feature vector $\mathbf{x} \in \mathbb{R}^p$ we consider classification.
- The idea: find the nearest feature vector to x in the training set and use its label.
- Can formalize "nearest" in terms of the Euclidean distance

$$\|\mathbf{x}_i - \mathbf{x}_{i'}\|_2 = \sqrt{\sum_{j=1}^p (x_{ij} - x_{i'j})^2}$$

Algorithm:

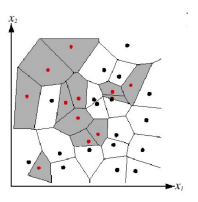
1. Find example (\mathbf{x}_*, y_*) (from the stored training set) closest to \mathbf{x} . That is:

$$\mathbf{x}_* = \underset{\mathbf{x}_i \in \{\mathbf{x}_1, \dots, \mathbf{x}_n\}}{\operatorname{argmin}} \operatorname{distance}(\mathbf{x}_i, \mathbf{x})$$

2. Output y_* as the label

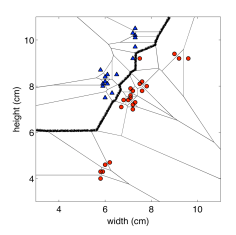
Nearest Neighbors: Decision Boundaries

We can visualize the behavior in the classification setting using a Voronoi diagram.

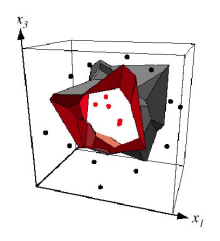


Nearest Neighbors: Decision Boundaries

Decision boundary: the boundary between regions of the feature space assigned to different categories.

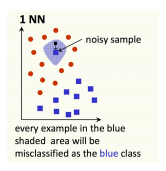


Nearest Neighbors: Decision Boundaries



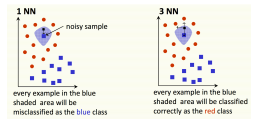
Example: 2D decision boundary

Nearest Neighbors



- Nearest neighbors sensitive to noise or mis-labeled data ("class noise").
- Solution? Smooth by having k nearest neighbors vote

k-Nearest Neighbors



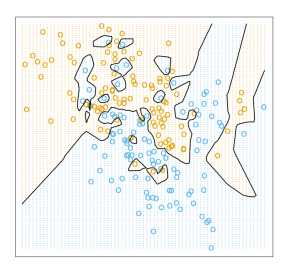
Algorithm (k-NN):

- 1. Find k data points $(\mathbf{x}_{(1)}, y_{(1)}), \dots, (\mathbf{x}_{(k)}, y_{(k)})$ closest to the test instance \mathbf{x}
- 2. Classification output is majority class

$$y = \arg \max_{y \in C} \sum_{i=1}^{k} \mathbb{I}(y = y_{(i)})$$

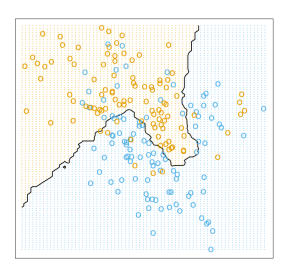
k-NN

k=1



k-NN

k=15



k-NN

Tradeoffs in choosing k

- Small k
 - Good at capturing fine-grained patterns
 - Overfitting: may be sensitive to random idiosyncrasies in the training data (i.e.).
- Large k
 - Makes stable predictions by averaging over lots of examples
 - ▶ Underfitting: may fail to capture important regularities.
- Balancing k
 - Optimal choice of k depends on number of data points n.
 - ▶ Nice theoretical properties if

$$k \to \infty$$
, and $\frac{k}{n} \to 0$ (ESL 2.4).

▶ Rule of thumb: choose $k < \sqrt{n}$ via cross-validation!

Pitfalls: The Curse of Dimensionality

- k-NN suffers the curse of dimensionality!
 - ▶ In high dimensions, "most" points are approximately the same distance because they are far away from each other.
- Saving grace: some datasets (e.g. images) may have low intrinsic dimension, i.e. lie on or near a low-dimensional manifold.



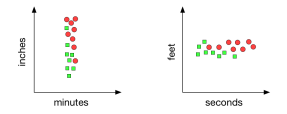


Image credit: https://scikit-learn.org/stable/modules/generated/sklearn.datasets.make_swiss_roll.html

The neighborhood structure depends on the intrinsic dimension.

Pitfalls: Normalization

- Nearest neighbors can be sensitive to the ranges of different features.
- Often, the units are arbitrary:



• Simple fix: normalize each dimension to be zero mean and unit variance. I.e., compute the mean μ_j and standard deviation σ_j , and take

$$\tilde{x}_j^{(i)} = \frac{x_j^{(i)} - \mu_j}{\sigma_j}$$

Caution: depending on the problem, the scale might be important!

Pitfalls: Computational Cost

- Computational cost for training: 0
- Computational cost for classifying test data, per data point (un-modified algorithm)
 - ▶ Calculuate *p*-dimensional Euclidean distances with *n* data points: $\mathcal{O}(np)$
 - ▶ Sort the distances: $\mathcal{O}(n \log n)$
- This must be done for each test data point, which is very expensive by the standards of a learning algorithm!
- Need to store the entire dataset in memory!
- Tons of work has gone into algorithms and data structures for efficient nearest neighbors with high dimensions and/or large datasets.

Example: Digit Classification

Decent performance when lots of data

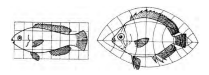
0123456789

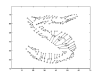
- Yann LeCunn MNIST Digit Recognition
 - Handwritten digits
 - 28x28 pixel images: d = 784
 - 60,000 training samples
 - 10,000 test samples
- Nearest neighbour is competitive

Test Error Rate (%)	
12.0	
5.0	
2.4	
1.1	
0.67	
3.6	
1.1	
4.7	
1.6	
8.0	
0.7	

Example: Digit Classification

- Changing the similarity measure can really improve *k*-NN.
- Example: shape contexts for object recognition. In order to achieve invariance to image transformations, they tried to warp one image to match the other image.
 - ▶ Distance measure: average distance between corresponding points on warped images
- Achieved 0.63% error on MNIST, compared with 3% for Euclidean KNN.
- Competitive with the state of the art at the time, but required careful engineering.







[Belongie, Malik, and Puzicha, 2002. Shape matching and object recognition using shape contexts.]