CSCE 421: Machine Learning

Lecture 7: Logistic Regression and Gradient Descent

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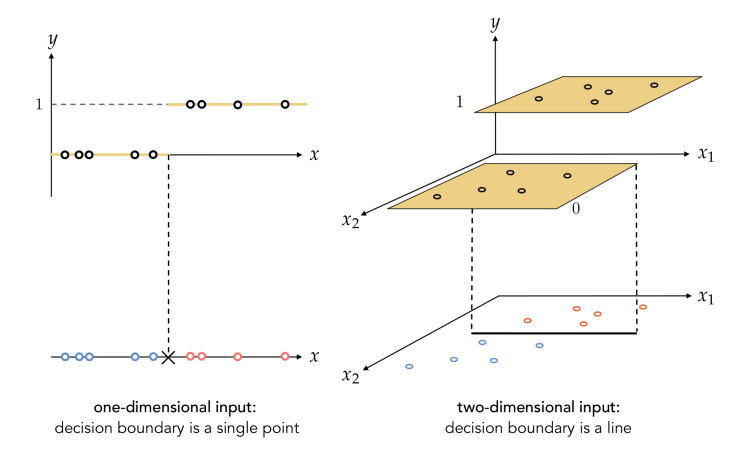
Zhale Nowroozilarki

Goals

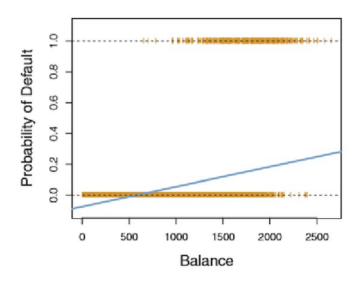
- Re-motivate Logistic Regression
- Understand why the shift from regression to classification results in needing Gradient Descent
- Learn how to use with with Cross-Validation

Decision Boundaries (Discriminative vs. Generative Models)

Classification – Step Functions

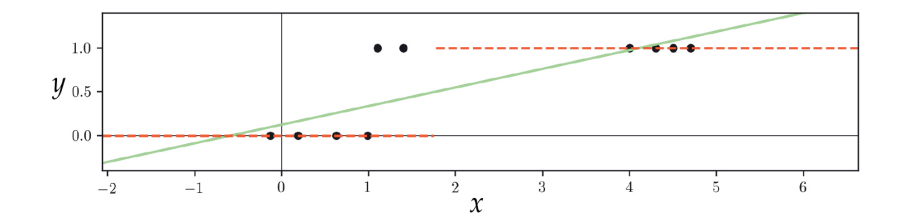


Linear Regression for Classification



If you set y > 0.5 as a decision rule – not clear it works Hard to interpret – results are not contained in [0,1]

Step function from Linear Regression Boundary



Bernouli Distribution

• We can create a probability density function that represents a single experiment asking yes/no

$$Y \sim Bernouli(\theta), Y \in \{0,1\}$$

$$p(y|\theta) = \theta^{I(y=1)} (1-\theta)^{I(y=0)} = \begin{cases} \theta, y = 1\\ 1-\theta, y = 0 \end{cases}$$

• Can think of this as a coin toss experiment and the likelihood of heads vs. tails

Logistic Regression

- Parametric classification method (not regression), which is sometimes referred to as a "generalization" of linear regression because
 - We still compute a linear combination of feature inputs, $x^T w$ (sometimes written $w^T x$)
 - However, instead of estimating the continuous output variable, we pass this into a function

$$\mu(\mathbf{w}^T \mathbf{x}) = \frac{1}{1 + e^{-(\mathbf{w}^T \mathbf{x})}}$$

— Where $0 \le \mu(\mathbf{w}^T \mathbf{x}) \le 1$, and where the Gaussian noise of linear regression is replaced by the Bernoulli Distribution so that

$$p(y|x, \mathbf{w}) = Ber(y|\mu(\mathbf{w}^T x))$$

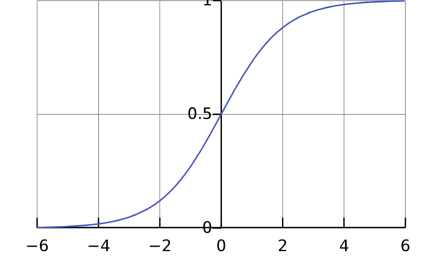
– Therefore, the output belongs to a class 1 (y = 1) with probability $\mu(w^Tx)$, and class 0 (y = 0) with probability $1 - \mu(w^Tx)$

Why use a Sigmoid Function?

$$\sigma(\boldsymbol{\eta}) = \frac{1}{1 + e^{-(\boldsymbol{\eta})}} = \frac{e^{\boldsymbol{\eta}}}{1 + e^{\boldsymbol{\eta}}}$$



- Bounded between 0 and 1 <- thus interpretable as a probability
- Monotonically increasing <- thus can be used for classification rules



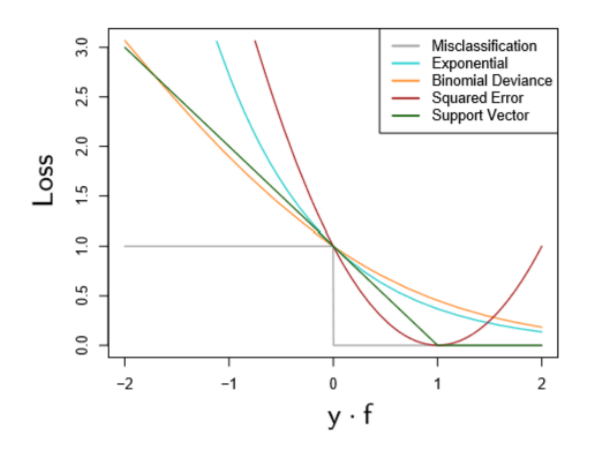
$$\sigma(\pmb{\eta}) > 0.5$$
 , positive class (y = 1)

$$\sigma(\eta) \le 0.5$$
 , negative class (y = 0)

Nice computational properties for optimizing criterion function

Decision Boundaries (Margin Classification)

Types of Loss



Logistic Regression: Representation

- Setup classification problem for two classes
 - Input: $x \in \mathbb{R}^p$
 - Output: y ∈ {0, 1}
 - Training Data: $D = \{(x_1, y_1), ..., (x_n, y_n)\}$
 - Model parameters: w (weights)
 - Model:

$$p(y | \mathbf{x}, \mathbf{w}) = \sigma(\mathbf{w}^T \mathbf{x}), \sigma(\mathbf{\eta}) = \frac{1}{1 + e^{-(\mathbf{\eta})}} = \frac{e^{\mathbf{\eta}}}{1 + e^{\mathbf{\eta}}}$$

$$y = \begin{cases} 1, p(y \mid x, w) > 0.5 \\ 0, otherwise \end{cases}$$

Maximum Likelihood Estimation

$$\hat{\theta} = \operatorname*{argmax}_{\theta} L(\theta; x)$$

- Likelihood is $\prod_{p:y_p=1} p(x_p) \prod_{p':y_{p'}=0} (1 p(x_{p'}))$
- With one dimension this means optimizing intercept w_0 and slope w_1
- So maximizing the parameters means finding the optimal w that maximize the product

Multiple Logistic Regression

$$\log\left(\frac{p(x)}{1-p(x)}\right) = w_0 + w_1 x_1 + \dots + w_p x_p$$

- $p(y|x,w) = Ber(y|\sigma(w^Tx))$, with a linear decision boundary at p(x) > 0.5
- We can then calculate the negative of the log of the likelihood (negative log likelihood)

$$NLL(w) = -\sum_{i=1}^{n} \left(y_i \log \sigma(w^T x_i) + (1 - y_i) \log \left(1 - \sigma(w^T x_i) \right) \right)$$

Multiple Logistic Regression

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• Now suppose we recast $\tilde{y} = \{-1, 1\}$, then

$$NLL(w) = -\sum_{i=1}^{n} \left(\log \left(1 + e^{-\widetilde{y_i} w^T x_i} \right) \right)$$

Which has no closed form solution



Logistic Regression: Evaluation

Data likelihood (1 training sample)

$$p(y|x) = \begin{cases} \sigma(w^T x) & y = 1 \\ 1 - \sigma(w^T x) & otherwise \end{cases} = \sigma(w^T x)^y (1 - \sigma(w^T x))^{1-y}$$

Data likelihood (all training samples)

$$L(D, w) = \prod_{i=1}^{n} p(y_i | x_i, w) = \prod_{i=1}^{n} \sigma(w^T x_i)^{y_i} (1 - \sigma(w^T x_i))^{1 - y_i}$$

Log Likelihood

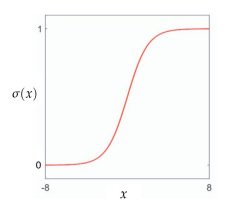
$$l(D, w) = \sum_{i=1}^{n} (y_i \log \sigma(w^T x_i) + (1 - y_i) \log(1 - \sigma(w^T x_i)))$$

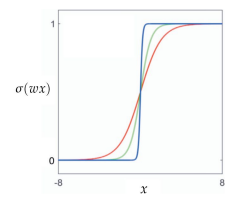
Negative Log Likelihood (the cross-entropy error)

$$nll(w) = -\sum_{i=1}^{n} \left(y_i \log \sigma(w^T x_i) + (1 - y_i) \log \left(1 - \sigma(w^T x_i) \right) \right)$$



Training Logistic Regression





Where training in linear regression shifted line (slope/intercept) training in logistic regression alters the "step"-like nature of the sigmoid

Training Logistic Regression with Gradient Descent

- Gradient Descent Methods
- How to optimize the coefficients

Logistic Regression: Optimization

The cross-entropy error

$$\mathcal{E}(w) = -\sum_{i=1}^{n} \left(y_i \log \sigma(w^T x_i) + (1 - y_i) \log \left(1 - \sigma(w^T x_i) \right) \right)$$

- Must find weights w of logistic regression that minimize the error
- Because there is no closed form solution, unlike linear regression, we need gradient descent

$$\nabla \mathcal{E}(w) = \sum_{i=1}^{n} (\sigma(w^{T} x_{i}) - y_{i}) x_{i}$$

Is this convex?

Cross Entropy Error: Convex

$$\nabla \mathcal{E}(w) = \frac{\partial \mathcal{E}(w)}{\partial w} = \sum_{i=1}^{n} (\sigma(w^{T} x_{i}) - y_{i}) x_{i}$$

$$H = \frac{\partial^2 \mathcal{E}(w)}{\partial w w^T} = \sum_{i=1}^n \sigma(w^T x_i) (1 - \sigma(w^T x_i)) (x_i x_i^T)$$

For all **v**, substituting $\mu = \sigma(w^T x_i) (1 - \sigma(w^T x_i)) \ge 0$

$$\boldsymbol{v}^T H \boldsymbol{v} = \mu \boldsymbol{v}^T \left(\sum_{i=1}^n x_i x_i^T \right) \boldsymbol{v} = \mu \sum_{i=1}^n (x_i^T \boldsymbol{v})^T (x_i^T \boldsymbol{v}) = \mu \sum_{i=1}^n ||x_i^T \boldsymbol{v}||_2^2 \ge 0$$

Gradient Descent in Logistic Regression

• The cross-entropy error

$$\mathcal{E}(w) = -\sum_{i=1}^{n} \left(y_i \log \sigma(w^T x_i) + (1 - y_i) \log \left(1 - \sigma(w^T x_i) \right) \right)$$

• Gradient Descent Optimization Expression

$$\mathbf{w} \leftarrow \mathbf{w} - \alpha(k) \nabla \mathcal{E}(\mathbf{w})$$
$$\nabla \mathcal{E}(\mathbf{w}) = \sum_{i=1}^{n} (\sigma(\mathbf{w}^{T} x_{i}) - y_{i}) x_{i}$$

Batch Gradient Descent

- 1. Initialize $w, \epsilon, \alpha, k = 0$
- 2. While $\|\nabla \mathcal{E}(w)\|_2 > \epsilon$

$$2a. k = k + 1$$

2b.
$$w = w - \alpha(k)(\sum_{i=1}^{n} (\sigma(w^{T}x_{i}) - y_{i}) x_{i})$$

Stochastic Gradient Descent

Update weights one sample at a time

- 1. Initialize $w, \epsilon, \alpha, k = 0$
- 2. Loop until convergence
 - 2a. k = k + 1
 - 2b. Randomly choose a single sample (x_i, y_i)
 - 2c. Compute its contribution to the gradient $g_i = (w^T x_i y_i)x_i$
 - 2d. Update weights $w = w \alpha(k)g_i$

Mini Batch Gradient Descent

Update weights with a subset of samples at a time

- 1. Initialize $w, \epsilon, \alpha, k = 0$
- 2. Loop until convergence
 - 2a. k = k + 1
 - 2b. Randomly choose a set of samples (x_i, y_i)
 - 2c. Compute their contribution to the gradient $g_i = (w^T x_i y_i)x_i$
 - 2d. Update weights $w = w \alpha(k) \sum_{i=1}^{M} g_i$

This is a good compromise between batch gradient descent (complete gradient) and stochastic gradient descent (gradient of one sample at a time)

Common mini-batch is a set M between 50 and 250 samples

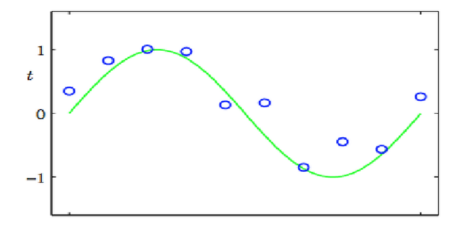
This is typically used for training neural networks

Goals

- Re-motivate Logistic Regression
- Understand why the shift from regression to classification results in needing Gradient Descent
- Non-linear transformations!
- Learn how to use with with Cross-Validation

What if data does not fit a line?

Example: Samples of a sine function



We can use a non-linear basis function

$$\phi(x) \colon x \in \mathbb{R}^p \to z \in \mathbb{R}^M$$

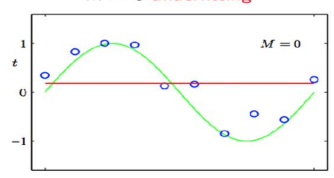
So that
$$y = w^T z = w^T \phi(x)$$

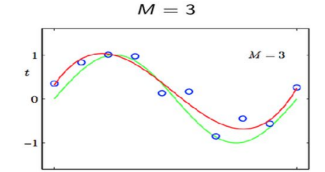
More non-linear basis functions

Example: Samples from a sine function

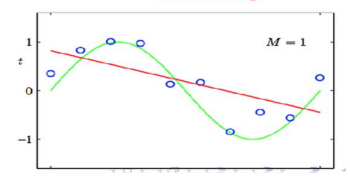
Polynomial basis function $\phi(\mathbf{x}) = [1 \times \dots \times^M]^T$

M = 0 underfitting

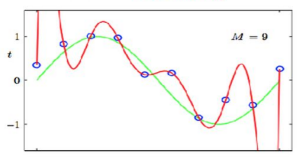




M = 1 underfitting







But higher-order non-linearity tends to overfit

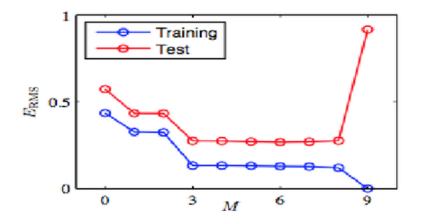
Weights of high order polynomials are very large

$$y_i = \mathbf{w}^T \mathbf{z_i} = \mathbf{w}^T \phi(\mathbf{x_i}), \ \mathbf{z_i} = \phi(\mathbf{x_i}) \in \mathbb{R}^M$$

	M=0	M=1	M=3	M=9
w_0	0.19	0.82	0.31	0.35
w_1		-1.27	7.99	232.37
w_2			-25.43	-5321.83
w_3			17.37	48568.31
w_4				-231639.30
w_5				640042.26
w_6				-1061800.52
w_7				1042400.18
w_8				-557682.99
w_9				125201.43

Overfitting

- The risk of using highly flexible (complicated) models without enough data is that the model overfits
- Leads to poor generalization
- How do you detect overfitting?
 - Can plot model complexity vs. the objective function
 - As complexity improves, will see training improves while testing improves then deteriorates



- How do you prevent this?
 - Feature selection, more data, or regularization



Linear Regression: Summarization

Representation

$$y = w^T x$$

Evaluation

$$\min_{w} RSS(w) = \sum_{i=1}^{N} (y_i - w^T x_i)^2$$

Analytic Optimization

$$\boldsymbol{w}^* = (\boldsymbol{X}^T \boldsymbol{X})^{-1} \boldsymbol{X}^T \boldsymbol{y}$$

Approximate Optimization

via gradient descent

• In linear regression the maximum likelihood estimation = the ordinary least squares solution

Logistic Regression: Summarization

• Generalization of linear regression

$$\sigma(w^Tx)$$

is interpretable as a probability

• Evaluation through cross-entropy error

$$\mathcal{E}(w) = -\sum_{i=1}^{n} \left(y_i \log \sigma(w^T x_i) + (1 - y_i) \log \left(1 - \sigma(w^T x_i) \right) \right)$$

Optimization through gradient descent

Practical Use of Logistic Regression

- Understanding re-sampling
- Measures of Accuracy when repeating modeling
- Python/Discussion!



Resampling Methods. Why?

- Resampling is a method by which we re-draw subsets of training data to see if model fit changes
- This allows us to see how stable our model is by how much the coefficient weights change
- We want to reduce model error rates
- How does one guarantee performance on a test set when only evaluating a training set?

Resampling Methods

Cross-validation

- Helps estimate measures of performance
- Allows for a choice in levels of model flexibility
- Used for hyperparameter tuning
- (You tend to see this more in computer science work)

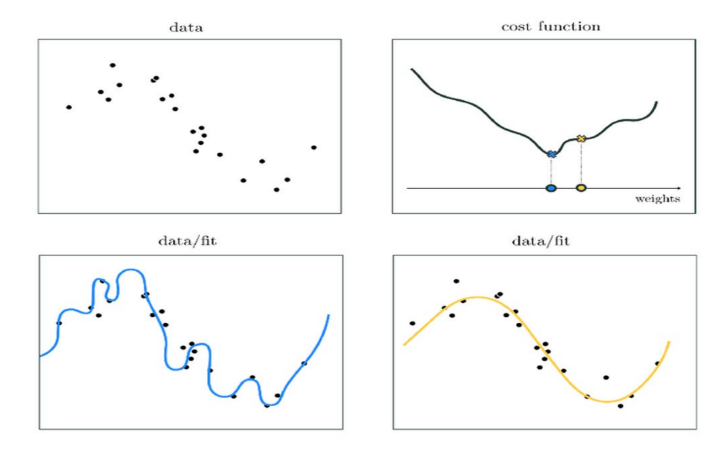
Bootstrapping

- Also helps give estimates of model performance
- Measures of accuracy surrounding parameter estimates
- Repeated trials help simulate different event rates
- (you tend to see this more in biostatistics work)



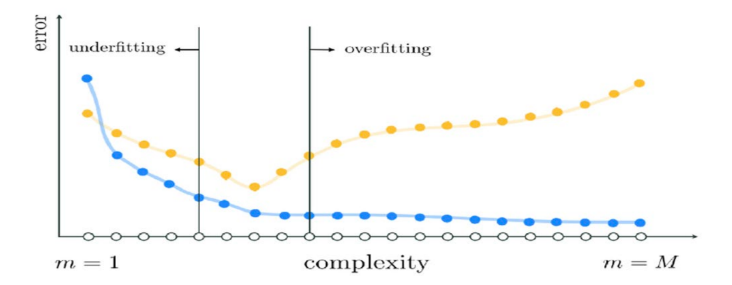
Overfitting

 Consider fitting a polynomial model to data, how does training error change with increasing degree?



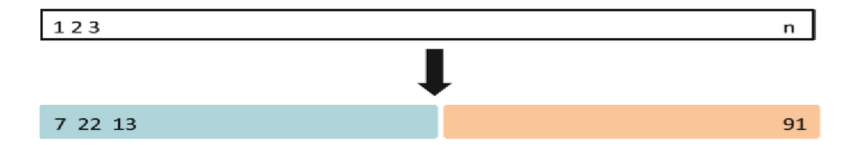
Overfitting

- As a model gets more complex, we become prone to reducing training error at the expense of testing error
- We also see sensitivity increase to specific inclusion of variables



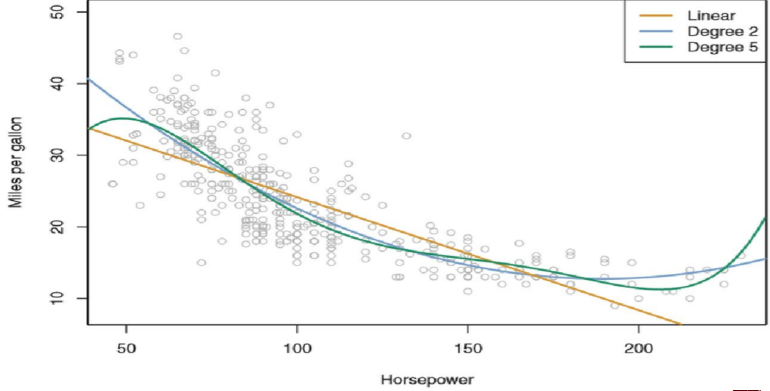
Cross-Validation: Test Set Approach

- We can randomly split the training data (after having removed our "hold out" data) to train and test (or validate).
- These splits can be 50/50, 80/20, 90/10



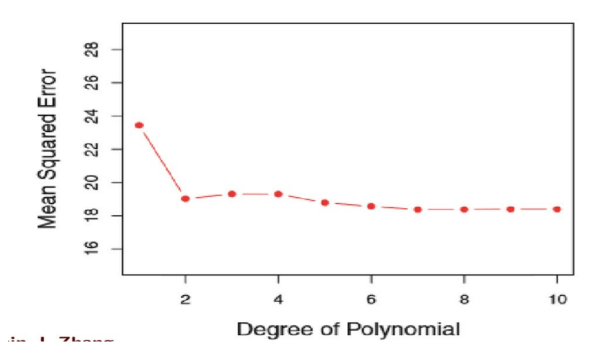
Let's compare polynomial fit for horsepower and mpg

- Model for estimating relationship between engine horsepower and the miles per gallon achieved
- Best fit is $mpg = w_0 + w_1 horsepower + w_2 horsepower^2 + \epsilon$
- So how do we determine the best degree of polynomial?



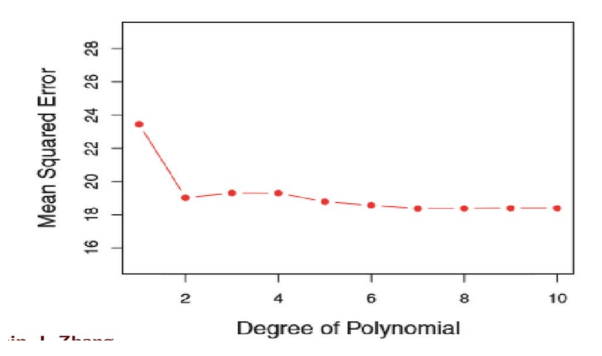
Cross-validation: Test set approach

- Best fit is $mpg = w_0 + w_1 horsepower + w_2 horsepower^2 + \epsilon$
- Split the data 50/50
- See that the cubic term actually increases error



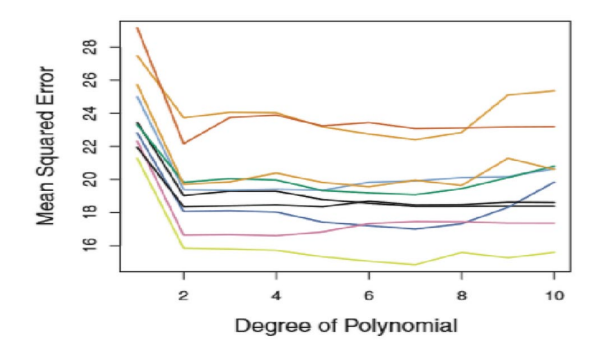
Cross-validation: Test set approach

- Best fit is $mpg = w_0 + w_1 horsepower + w_2 horsepower^2 + \epsilon$
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- See that the cubic term actually increases error
- What if we repeat this experiment multiple times?



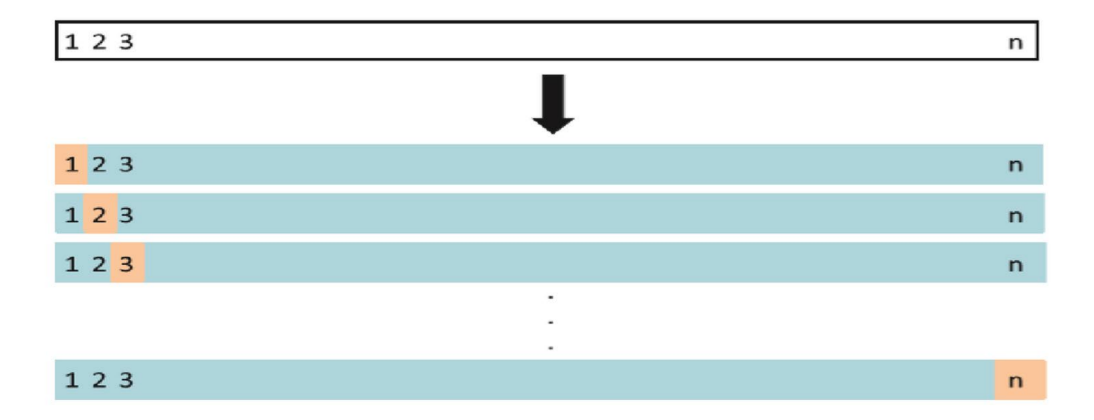
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Leave-one-out Cross-validation

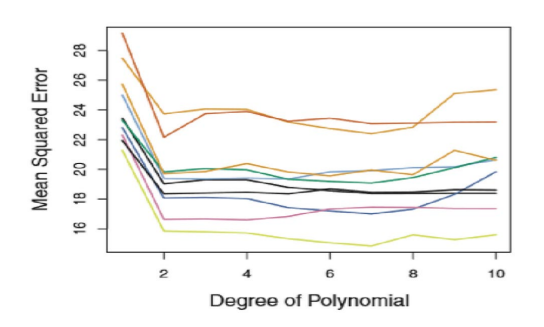


Advantages of LosoCV

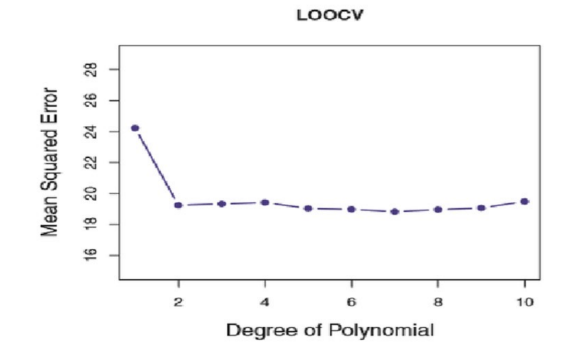
- In each iteration, train with as much data as possible
- Consider error as the average across all iterations
- This presents far less bias
- It won't overestimate/underestimate test error as much as a single train/test split can

Comparison with MPG and Horsepower

Train-test split

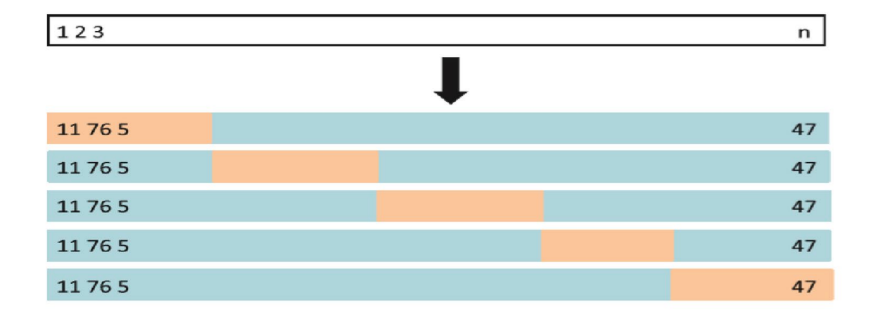


Loso cv



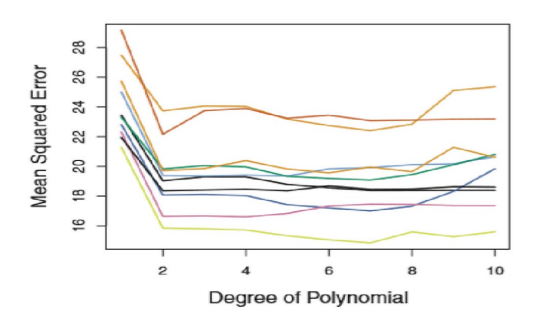
K-fold Cross-Validation

- LOSO CV can be computationally expensive. A single train/test split not robust enough.
- Iterate subsets instead to find a balance

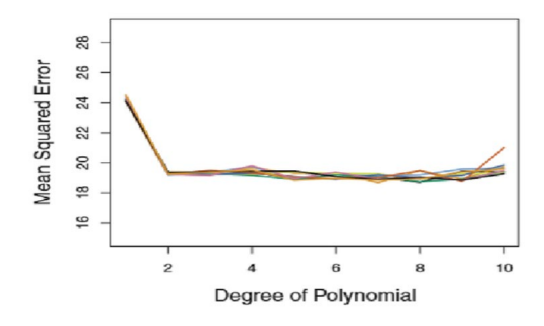


Comparison with MPG and Horsepower

Train-test split



K=10 fold cross-validation





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