

CS 559 Machine Learning Linear Regression

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Plan for today



Linear Regression

Gradient Descent

Learning Rate

Features

Beyond Linear Regression Models

Bias and Variance

The Curse of Dimensionality

Maximum Likelihood Estimator

Regression Task

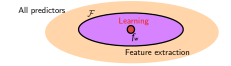


 \times \rightarrow function f \rightarrow \rightarrow

The Learning Problem



▶ **Hypothesis class**: we consider some restricted set \mathcal{F} of mappings $f: \mathcal{X} \to \mathcal{Y}$ from input data to output.



- ▶ **Estimation**: on the basis of a training set of examples of their labels, $\{(\mathbf{x}_1, y_1), ..., (\mathbf{x}_N, y_N)\}$, we find an estimate $\hat{f} \in \mathcal{F}$
- **Evaluation**: we measure how well \hat{f} generalizes to yet unseen examples, i.e., whether $\hat{f}(\mathbf{x}_{\text{new}})$ agrees with y_{new}



Loss

A loss function $Loss(\mathbf{x}, y, \mathbf{w})$ quantifies how wrong you would be if you used \mathbf{w} to make a prediction on \mathbf{x} when the correct output is y. It is the object we want to minimize.

► Loss is a function of the parameters **w** and we can try to minimize it directly.



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- Loss is a function of the parameters w and we can try to minimize it directly.
- ▶ We reduce the estimation problem to a minimization problem.
- Valid for any parameterized class of mapping from examples to predictions.
- ▶ Valid when the predictions are discrete labels or real valued as long as the loss is defined properly.

Linear Regression



Prediction

$$f_w(\mathbf{x}) = \mathbf{w}^T \mathbf{x}$$

Residual

The residual is the amount by which prediction overshoots the target.

Squared loss

$$L(\mathbf{x}, y, \mathbf{w}) = \underbrace{(f_w(\mathbf{x}) - y)^2}_{\text{residual}}$$

Example:

$$\mathbf{w} = [2, -1], \mathbf{x} = [2, 0], y = -1, L(\mathbf{x}, y, \mathbf{w}) = 25$$

Loss Minimization Framework



Loss is easy to minimize if there is only one example.

Minimize Training Loss

$$\min_{\mathbf{w} \in \mathbb{R}^d} \mathsf{TrainLoss}(\mathbf{w}) = \underbrace{\frac{1}{N} \sum_{(x,y) \in D_{\mathsf{train}}} L(\mathbf{x}, y, \mathbf{w})}_{\mathsf{empirical loss}}$$

Key: learn a **w** to make global tradeoffs (N is the number of examples in D_{train}).

Training and Testing performance: sampling



- Assume each training and test example-label pair (\mathbf{x}, y) is drawn independently at random from the same but unknown population of examples and labels.
- We can represent this population as a joint probability distribution $P(\mathbf{x}, y)$ so that each training/testing example is a sample from this distribution $(\mathbf{x}_i, y_i) \sim P$.

Empirical (training) loss =
$$\frac{1}{N} \sum_{i} \text{Loss}(y_i, f_w(\mathbf{x}))$$

Expected (testing) loss =
$$E_{(\mathbf{x},y)\sim P}\{\text{Loss}(y_i, f_w(\mathbf{x}))\}$$

► The training loss based on a few sampled examples and labels serves as a proxy for the test performance measured over the whole population.

Regression Loss Functions



Squared loss

$$L(\mathbf{x}, y, \mathbf{w})_2 = (f_w(\mathbf{x}) - y)^2$$

Absolute loss

$$L(\mathbf{x},y,\mathbf{w})_1 = |f_w(\mathbf{x}) - y|$$

Which loss to use?



Example:

$$D_{\mathsf{train}} = \{(1,0), (1,2), (1,1000)\}$$

► For least square regression (L2): w that minimizes training loss is mean y

$$\frac{\partial L_2}{\partial f_w} = -\frac{2}{N} \sum_{i=1}^{N} (y_i - f_w) = 0 \to f_w = \frac{1}{N} \sum_{i=1}^{N} y_i$$

Mean: tries to accommodate every example, popular

Which loss to use?



Example:

$$D_{\mathsf{train}} = \{(1,0), (1,2), (1,1000)\}$$

► For least absolute deviation regression (L1): w that minimizes training loss is median y

$$\frac{\partial L_1}{\partial f_w} = -\frac{1}{N} \sum_{i=1}^{N} \operatorname{sgn}(y_i - f_w)$$

(The derivative equals to 0 when there is the same number of positive and negative terms among $y_i - f_w$, which means f_w should be the median of y_i)

Median: more robust to outliers.

How to optimize?



Gradient

$$\nabla_{\mathbf{w}} \operatorname{TrainLoss}(\mathbf{w})$$

The direction that increases the loss the most

Algorithm: Gradient Descent

Initialize w

For t = 1, ..., T

$$\mathbf{w} \leftarrow \mathbf{w} - \underbrace{\eta}_{\text{step size}} \underbrace{\nabla_{\mathbf{w}} \text{TrainLoss}(\mathbf{w})}_{\text{gradient}}$$

Least squares regression



Objective function

$$\mathsf{TrainLoss}(\mathbf{w}) = \frac{1}{N} \sum_{(\mathbf{x}, y) \in D_{\mathsf{train}}} (\mathbf{w}^\mathsf{T} \mathbf{x} - y)^2$$

Gradient with respect to **w** (use chain rule)

$$\nabla_{w} \mathsf{TrainLoss}(\mathbf{w}) = \frac{1}{N} \sum_{(\mathbf{x}, y) \in D_{\mathsf{train}}} 2(\underbrace{\mathbf{w}^{\mathsf{T}} \mathbf{x}}_{\mathsf{prediction}} - \underbrace{y}_{\mathsf{target}}) \mathbf{x}$$

Gradient descent is slow



Gradient

$$\nabla_{w} \mathsf{TrainLoss}(\mathbf{w}) = \frac{1}{N} \sum_{(x,y) \in D_{\mathsf{train}}} 2(\underbrace{\mathbf{w}^{\mathsf{T}} \mathbf{x}}_{\mathsf{prediction}} - \underbrace{y}_{\mathsf{target}}) \mathbf{x}$$

Gradient descent update

$$\mathbf{w} \leftarrow \mathbf{w} - \underbrace{\eta}_{\text{step size}} \underbrace{\nabla_{w} \text{TrainLoss}(\mathbf{w})}_{\text{gradient}}$$

Each iteration requires going over all training examples – expensive when have lots of data

Stochastic Gradient Descent



Gradient Descent (GD)

$$\mathbf{w} \leftarrow \mathbf{w} - \underbrace{\eta}_{\text{step size}} \underbrace{\nabla_{w} \text{TrainLoss}(\mathbf{w})}_{\text{gradient}}$$

Stochastic Gradient Descent (SGD)

For each $(\mathbf{x}, y) \in D_{train}$

$$\mathbf{w} \leftarrow \mathbf{w} - \eta \nabla_{w} \mathsf{Loss}(\mathbf{x}, y, \mathbf{w})$$

Key: It's not about quality, it's about quantity.

Gradient Descent



- 1: procedure Batch Gradient Descent
- 2: **for** i in range(epochs) **do**
- 3: $g^{(i)}(\mathbf{w}) = \text{evaluate_gradient}(\text{TrainLoss, data, } \mathbf{w})$
- 4: $\mathbf{w} = \mathbf{w} \text{learning_rate} * g^{(i)}(\mathbf{w})$

- Can be very slow.
- Intractable for datasets that don't fit in memory.
- Doesn't allow us to update our model online, i.e. with new examples on-the-fly.
- Guaranteed to converge to the global minimum for convex error surfaces and to a local minimum for non-convex surfaces.

Stochastic Gradient Descent



```
    procedure STOCHASTIC GRADIENT DESCENT
    for i in range(epochs) do
    np.random.shuffle(data)
    for example ∈ data do
    g(i)(w) = evaluate_gradient(loss, example, w)
    w = w - learning_rate * g(i)(w)
```

- Allow for online update with new examples.
- With a high variance that cause the objective function to fluctuate heavily.

Mini-batch Gradient Descent



```
    procedure MINI-BATCH GRADIENT DESCENT
    for i in range(epochs) do
    np.random.shuffle(data)
    for batch ∈ get_batches(data, batch_size=50) do
    g<sup>(i)</sup>(w) = evaluate_gradient(loss, batch, w)
    w = w - learning_rate * g<sup>(i)</sup>(w)
```

- Reduces the variance of the parameter updates, which can lead to more stable convergence;
- Can make use of highly optimized matrix optimizations common to state-of-the-art deep learning libraries that make computing the gradient w.r.t. a mini-batch very efficient.

Learning Rate



Gradient Update:

$$\mathbf{w} \leftarrow \mathbf{w} - \eta \nabla_{w} \mathsf{Loss}(x, y, \mathbf{w})$$

Question: what should the step size (learning rate) be?

Strategies:

For each $(x, y) \in D_{train}$

- ▶ Constant: $\eta = 0.1$
- lackbox Descreasing: $\eta = \frac{1}{\sqrt{\# \text{ updates made so far}}}$

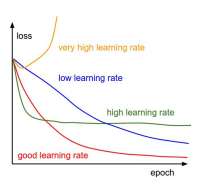
Learning Rate



Set the learning rate η carefully:

$$\mathbf{w}^{(i)} \leftarrow \mathbf{w}^{(i-1)} - \eta \nabla L(\mathbf{w}^{(i-1)})$$

If there are more than 3 parameters, it is hard to visualize the loss w.r.t the parameters. But you can visualize the loss w.r.t the # of parameter updates (epoch).



Adaptive Learning Rates



- ► Popular & Simple Idea: Reduce the learning rate by some factor every few epochs.
 - ► At the beginning, we are far from the destination, so we use larger learning rate
 - After several epochs, we are close to the destination, so we reduce the learning rate
 - e.g. $\frac{1}{t}$ decay: $\eta^{(t)} = \frac{\eta}{\sqrt{t+1}}$
- Learning rate cannot be one-size-fits-all
 - ► Giving different parameters different learning rates

[An overview of gradient descent optimization algorithms]

Adagrad



$$\eta^{(t)} = \frac{\eta}{\sqrt{t+1}}$$

$$g_i^{(t)} = \frac{\partial L(x, y, \mathbf{w}_i^{(t)})}{\partial w_i^{(t)}}$$

Vanilla Gradient descent

$$\mathbf{w}^{(t+1)} \leftarrow \mathbf{w}^{(t)} - \eta^{(t)} g^{(t)}$$

➤ Adagrad ¹: Divide the learning rate of each parameter by the root mean square of its previous derivatives

$$\mathbf{w}^{(t+1)} \leftarrow \mathbf{w}^{(t)} - rac{\eta^{(t)}}{\sigma^{(t)}} g^{(t)}$$

where $\boldsymbol{\sigma}^{(t)}$ is the root mean square of the previous derivatives of parameter \boldsymbol{w}

Adagrad



$$\begin{split} w^{(1)} \leftarrow w^{(0)} - \frac{\eta^{(0)}}{\sigma^{(0)}} g^{(0)} & \sigma^{(0)} = \sqrt{(g^{(0)})^2 + \epsilon} \\ w^{(2)} \leftarrow w^{(1)} - \frac{\eta^{(1)}}{\sigma^{(1)}} g^{(1)} & \sigma^{(1)} = \sqrt{\frac{1}{2} [(g^{(0)})^2 + (g^{(1)})^2] + \epsilon} \\ w^{(3)} \leftarrow w^{(2)} - \frac{\eta^{(2)}}{\sigma^{(2)}} g^{(2)} & \sigma^{(2)} = \sqrt{\frac{1}{3} [(g^{(0)})^2 + (g^{(1)})^2 + (g^{(2)})^2] + \epsilon} \\ \dots & \dots \\ w^{(t+1)} \leftarrow w^{(t)} - \frac{\eta^{(t)}}{\sigma^{(t)}} g^{(t)} & \sigma^{(t)} = \sqrt{\frac{1}{t+1} \sum_{i=1}^{t} (g^{(i)})^2 + \epsilon} \end{split}$$

 ϵ is a smoothing term that avoids division by zero (usually on the order of 1e-8)

Summary so far



- Linear predictors:
 f_w(x) based on score w^Tx
- ▶ Loss minimization: learning as optimization:

$$\min_{w} \mathsf{TrainLoss}(w)$$

Stochastic gradient descent: optimization algorithm:

$$\mathbf{w} \leftarrow \mathbf{w} - \eta \nabla_{w} \mathsf{Loss}(\mathbf{x}, y, \mathbf{w})$$

Two components



Score:

$$\mathbf{w}^T\mathbf{x} = \sum_{i=1}^d w_i x_i$$

- Previous: learning w via optimization
- Next: feature extraction and vector representation based on prior domain knowledge

Example: Data



Example: specifies that y is the ground-truth output for \mathbf{x}

$$(\mathbf{x}, y)$$

Training data: a set of examples

$$\begin{split} D_{\text{train}} &= [\text{(``...located in Hoboken, NJ...''}, 1.4M),\\ &\quad \text{(``...Fairfax, VA...''}, 0.7M),\\ &\quad \text{...]} \end{split}$$

Feature Extraction



- Example task: predict house price y, given a house
- Question: what properties of x might be relevant for predicting y?
- Feature extractor: Given input x, output a set of (feature name, feature value) pairs.

Faverage neighborhood house price: \$1.1M average school score: 5.6 distance to train stations: 3.4(m) parking garage: 1 size: 1200(sqrt)

Feature Vector



Mathematically, feature vector doesn't need feature names:

Taverage neighborhood house price:	\$1.1 <i>M</i>]		[1.1]
average school score:	5.6		5.6
distance to train stations:	3.4(m)	\rightarrow	3.4
parking garage:	1		1
size:	1200(sqrt)		[1200]

Feature vector

For in input X, its feature vector is:

$$\mathbf{x} = x_1, ..., x_d$$

x as a point in a high-dimensional space.

Linear Predictors



Weight vector $\mathbf{w} \in \mathbb{R}^d$

Feature vector $\mathbf{x} \in \mathbb{R}^d$

Score

weighted combination of features

$$\mathbf{w}^T\mathbf{x} = \sum_{i=1}^d w_i x_i$$

Organization of Features



Task: predict the price of a house

A house
$$\xrightarrow{\text{feature extraction}}$$

$$\begin{bmatrix} \text{avg_n_price} & \$1.1 \\ \text{avg_school} & 5.6 \\ \text{dist_train:} & 3.4 \\ \text{parking:} & 1 \\ \text{size:} & 1200 \end{bmatrix}$$

Which features to construct? need prior knowledge

Feature Templates



A feature template is a group of features all computed in a similar way. Input:

a house located in.....

Some feature templates:

- Average school score is __
- Distance to train stations less than __
- Located in __

Sparsity in feature vectors



Feature template: located in __

A house located in ... $\xrightarrow{\text{feature extraction}}$

[located_in_nyc	17
located_in_DC	0
located_in_seattle	0
located_in_pittsburg	0
located_in_sf	0

Inefficient to represent all the zeros...

Feature vector representation



Array representation: (good for dense features):

Map representation: (good for sparse features):

 $\left\{ \text{``avg_n_price''} : 1.1, \text{``avg_school''} : 5.6 \right\}$

Example: Vector Representation



Linear functions:

$$\mathbf{x} = [x]$$
 $\mathcal{F}_1 = \{x \to w_1 x + w_2 x^2 : w_1 \in \mathbb{R}, w_2 = 0\}$

Quadratic functions:

$$\mathbf{x} = [x, x^2]$$

$$\mathcal{F}_2 = \{x \to w_1 x + w_2 x^2 : w_1 \in \mathbb{R}, w_2 \in \mathbb{R}\}$$

Quadratic functions are supersets of linear functions

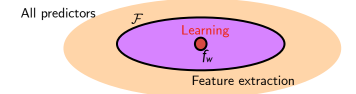
Expressivity



Hypothesis class

A hypothesis class is the set of possible predictors with a fixed \boldsymbol{x} and a varying \boldsymbol{w}

$$\mathcal{F} = \{f_w : \mathbf{w} \in \mathbb{R}^d\}$$



Features in linear predictors



Example: medical diagnosis

- Input features: height, weight, body temperature, blood pressure, etc.
- Output: real value.

Three issues: (non-linear in original measurements)

- Non-monotonicity
- Saturation
- ▶ Interactions between features

Non-monotonicity (1)



► Features, Try(1):

$$\mathbf{x} = [1, temperature(x)]$$

Output:

health
$$y \in \mathbb{R}$$

▶ Problem: favor extremes; true relationship is non-monotonic

Non-monotonicity (2)



► Features, Try(2): transform inputs

$$\mathbf{x} = [1, (temperature(x) - 37)^2]$$

Output:

health
$$y \in \mathbb{R}$$

Disadvantage: requires manually-specified domain knowledge

Non-monotonicity (3)



► Features, Try(3): transform inputs

$$\mathbf{x} = [1, temperature(x), temperature(x)^2]$$

Output:

health
$$y \in \mathbb{R}$$

 General rule: features should be simple building blocks to be pieced together

Saturation (1)



Example: product recommendation

Input: product information x

Output: relevance y

The number of people who bought x

Identity:
$$\mathbf{x} = N(x)$$

Problem: is 1000 people really 10 times more relevant than 100 people? Not quite...

Saturation (2)



▶ The number of people who bought x

Identity:
$$\mathbf{x} = N(x)$$

Log

$$\mathbf{x} = \log N(x)$$

Discretization

$$\mathbf{x} = [\mathbb{I}(1 < N(x) \le 10), \mathbb{I}(10 < N(x) \le 100), ...]$$

Interaction between features (1)



Example: predicting health

- Input: patient information x
- Output: health y

$$\mathbf{x} = [\mathsf{height}(x), \mathsf{weight}(x)]$$

Problem: cannot capture relationship between height and weight.

Interaction between features (2)



Try (2):

$$\mathbf{x} = (52 + 1.9(\text{height}(x) - 60) - \text{weight}(x))^2$$

Solution: define features that combine inputs.

Disadvantage: requires manually-specified domain knowledge.

Interaction between features (3)



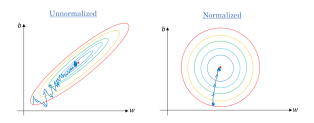
Try (3):

 $\mathbf{x} = [1, \mathsf{height}(x), \mathsf{weight}(x), \mathsf{height}(x)^2, \mathsf{weight}(x)^2, \mathsf{height}(x) \mathsf{weight}(x)]$

Solution: add features involving multiple measurements.

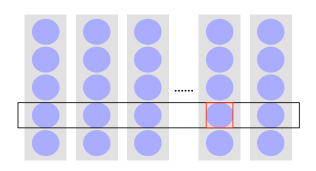
Feature Scaling





Feature Scaling





Assuming we have n (r = 1, ..., n) examples. For each dimension i, the mean is m_i , and the standard deviation is σ_i . Scaling features:

$$x_i^r \leftarrow \frac{x_i^r - m_i}{\sigma_i}$$

Linear in what?



Prediction driven by score:

$\mathbf{w}^T \mathbf{x}$

- ▶ Linear in w? Yes
- ▶ Linear in x? Yes
- Linear in each input feature? No!

Non-linearity

- Predictors can be expressive non-linear functions and decision boundaries of x
- ► Score is linear function of w, which permits efficient learning

Summary so far



- ► Goal: define features so that the hypothesis class contains good predictors
- Pay attention to non-linearity in x: non-monotonicity, saturation, interaction between features
- Suggested approach: define features x to be building blocks (e.g., monomials)
- Linear prediction: actually very powerful!



Objective:

$$\min L = \min \sum_{n=1}^{N} \{y_n - \mathbf{w}^T \mathbf{x}_n\}^2$$

▶ Transform into matrix operation $(\mathbf{y}_{N\times 1}, X_{N\times d}, \mathbf{w}_{d\times 1})$:

$$\min(\mathbf{y} - X\mathbf{w})^T(\mathbf{y} - X\mathbf{w})$$



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

where X is an $N \times d$ matrix. The solution if a linear function of the outputs \mathbf{y}



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▶ The resulting prediction errors $\epsilon_i = y_i - f(\mathbf{x}_i)$ are uncorrelated with any linear function of the inputs \mathbf{x} .



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- ▶ The resulting prediction errors $\epsilon_i = y_i f(\mathbf{x}_i)$ are uncorrelated with any linear function of the inputs \mathbf{x} .
- Easily extend to non-linear functions of the inputs. =

Beyond Linear Regression



► Example extension: m^{th} order polynomial regression on one-dimensional input where $f : \mathcal{R} \to \mathcal{R}$ is given by:

$$f(\mathbf{w}) = w_0 + w_1 x + ... + w_{m-1} x^{m-1} + w_m x^m$$

Beyond Linear Regression



▶ Example extension: m^{th} order polynomial regression on one-dimensional input where $f : \mathcal{R} \to \mathcal{R}$ is given by:

$$f(\mathbf{w}) = w_0 + w_1 x + ... + w_{m-1} x^{m-1} + w_m x^m$$

Solution as before:

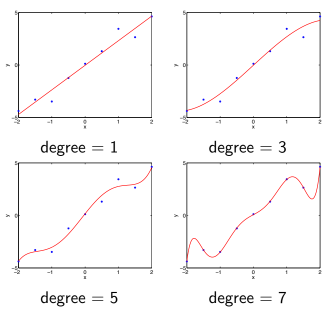
$$\mathbf{w} = (X^T X)^{-1} X^T \mathbf{y}$$

where:

$$\mathbf{w} = \begin{pmatrix} w_0 \\ w_1 \\ \dots \\ w_m \end{pmatrix}, \ \mathbf{X} = \begin{pmatrix} 1 & x_1 & x_1^2 & \dots & x_1^m \\ 1 & x_2 & x_2^2 & \dots & x_2^m \\ \dots & \dots & \dots & \dots \\ 1 & x_N & x_N^2 & \dots & x_N^m \end{pmatrix}$$

Polynomial Regression







- Underfitting:
 - ▶ Training an ML algorithm on training data \rightarrow 0.24 accuracy
 - ▶ Applying the ML algorithm on testing data → 0.25 accuracy
- Overfitting:
 - ▶ Training an ML algorithm on training data \rightarrow 0.95 accuracy
 - Applying the ML algorithm on testing data → 0.27 accuracy (huge generalization error!)



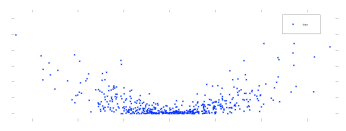


Figure: Regression data



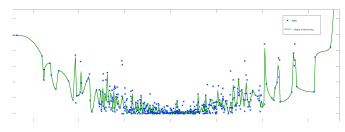


Figure: Overfitting



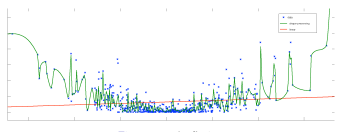


Figure: underfitting



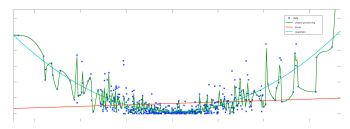


Figure: Overfitting, underfitting, and genelarization



- ► Underfitting:
 - ► Low dimensional
 - ▶ Heavily regularized
 - ► Bad modeling assumption



- Underfitting:
 - Low dimensional
 - Heavily regularized
 - ► Bad modeling assumption
- Overfitting:
 - High dimensional or non-parametric
 - Weakly regularized
 - Not enough modeling assumptions
 - Not enough data

Summary

- Training models too complex can cause overfitting
- ► Training models too simple (or wrong) can cause underfitting

Bias and Variance



Assuming a training set $(x_1,...,x_n)$ and their real associated y values. There is a function with noise $y=f(x)+\epsilon$ where the noise ϵ has mean 0 and variance σ^2 . We want to find a function $\hat{f}(x)$ that approximates the true function f(x). Expected squared prediction error at a point x is:

$$E[(y - \hat{f}(x))^{2}] = (E[\hat{f}(x) - f(x)])^{2} + (E[\hat{f}(x)^{2}] - E^{2}[\hat{f}(x)]) + \sigma^{2}$$

$$= Bias^{2} + Variance + \sigma^{2}$$

where

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

- Both contribute to errors
- ▶ Bias: error from incorrect modeling assumption
- Variance: error from random noise



Bias and Variance



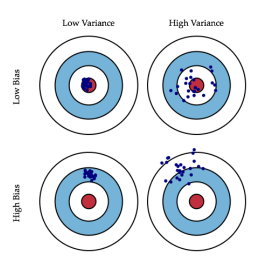


Figure: Graphical illustration of bias vs. variance

What to do with large bias?



Diagnosis:

- If your model cannot even fit the training examples, then you have large bias (underfitting)
- ▶ If you can fit the training data, but large error on testing data, then you probably have large variance (overfitting)

For bias: redesign your model

- Add more features as input
- A more complex model

What to do with large variance?



- More data (very effective, but not always practical)
- Regularization

Regularization

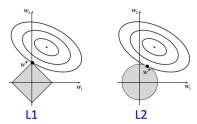


► L-2 regularization (Ridge regression)

$$L_2(X, y, \mathbf{w}) = \sum_{n=1}^{N} (f_w(x_n, y_n, \mathbf{w}) - y_n)^2 + \underbrace{\mathbf{w}^T \mathbf{w}}_{\text{L2 regularization}}$$

► L-1 regularization (LASSO regression)

$$L_1(X, y, \mathbf{w}) = \sum_{n=1}^{N} (f_w(x_n, y_n, \mathbf{w}) - y_n)^2 + \underbrace{|\mathbf{w}|}_{\text{L1 regularization}}$$



Model Selection



- Usually a trade-off between bias and variance
- Select a model that balances two kinds of error to minimize total error
- Cross-validation: it allows us to estimate the generalization error based on training examples alone.

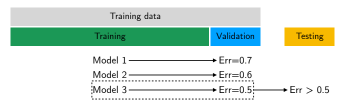


Figure: Training vs. Validation vs. Testing

N-fold Cross Validation



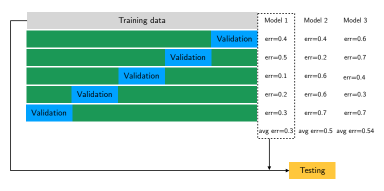


Figure: N-fold Cross Validation

- ► In general: we perform N runs, Each run, it allows to use (N-1)/N of the available data for training.
- ► If the number of data is very limited, we can set k=N (total number of data points). This gives the leave-one-out cross-validation technique.

The Curse of Dimensionality - when d is large



- As dimensionality grows: fewer observations per region.
- When a measure such as a Euclidean distance is defined using many coordinates, there is little difference in the distances between different pairs of samples.
- ▶ The proportion of an inscribed hypersphere with radius *r* and dimension *d*, to that of a hypercube with edges of length 2*r*:

$$\frac{V_{\text{hypersphere}}}{V_{\text{hypercube}}} = \frac{\frac{2r^d\pi^{d/2}}{d\Gamma(d/2)}}{(2r)^d} = \frac{\pi^{d/2}}{d2^{d-1}\Gamma(d/2)} \to 0 \text{ as } d \to \infty$$

The hypersphere becomes an insignificant volume relative to that of the hypercube

The Curse of Dimensionality - another angle



Consider a hypersphere of radius r=1 in a space of d dimensions, and ask what is the fraction of the volume of the hypersphere that lies between radius $r=1-\epsilon$ and r=1. We can evaluate this fraction by noting that the volume of a hypersphere of radius r in d dimensions must scale as r^d , and so we write:

$$V_d(r) = K_d r^d$$

where k_d depends on d. Thus the fraction is:

$$\frac{V_d(1) - V_d(1 - \epsilon)}{V_d(1)} = 1 - (1 - \epsilon)^d$$

Most of the volume of a hypersphere is concentrated in a thin shell near the surface!

The Curse of Dimensionality



- Real data will often be confined to a region of the space having lower effective dimensionality, and in particular the directions over which important variations in the target variables occur may be so confined.
- Real data will typically exhibit some smoothness properties (at least locally) so that for the most part small changes in the input variables will produce small changes in the target variables, and so we can exploit local interpolation-like techniques to allow us to make predictions of the target variables for new values of the input variables.



Warning of Math

Maximum Likelihood



We assume there is a true function f(x) and the true value is given by $y = f(x) + \epsilon$ where ϵ is a Gaussian distribution with mean 0 and variance σ^2 . Thus we can write:

$$p(y|x, \mathbf{w}, \beta) = \mathcal{N}(y|f(x), \beta^{-1})$$

where $\beta^{-1}=\sigma^2$. Assuming the data points are drawn independently from the distribution, we obtain the likelihood function:

$$p(\mathbf{y}|\mathbf{X},\mathbf{w},\beta) = \prod_{n=1}^{N} \mathcal{N}(y_n|\mathbf{w}^T\mathbf{x}_n,\beta^{-1})$$

Maximum Likelihood



The likelihood function:

$$p(\mathbf{y}|\mathbf{X},\mathbf{w},\beta) = \prod_{n=1}^{N} \mathcal{N}(y_n|\mathbf{w}^T\mathbf{x}_n,\beta^{-1})$$

The log-likelihood function:

$$\log p(\mathbf{y}|\mathbf{X}, \mathbf{w}, \beta) = \sum_{n=1}^{N} \mathcal{N}(y_n | \mathbf{w}^T \mathbf{x}_n, \beta^{-1})$$

$$= \underbrace{\frac{N}{2} \log \beta - \frac{N}{2} \log(2\pi)}_{\text{not related to } \mathbf{w}} - \beta E_D(\mathbf{w})$$
(2)

where:

$$E_D(\mathbf{w}) = \frac{1}{2} \sum_{n=1}^{N} \{y_n - \mathbf{w}^T \mathbf{x}_n\}^2$$

is the sum-of-squares loss function.

Maximum Likelihood



- ► Thus, maximum likelihood is equivalent to minimizing the sum-of-squares loss function.
- Solving for w:

$$\mathbf{w}_{ML} = (X^T X)^{-1} X^T \mathbf{y}$$

where X is an $N \times d$ matrix. The solution if a linear function of the outputs \mathbf{y}



End of Warning



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