3. Linear Neural Networks

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Summary for Dive Into Deep Learning, https://d21.ai/chapter_preface/index.html

- 3.1 Linear Regression
- 3.2. Linear Regression Implementation from Scratch
- 3.3 Concise Implementation of Linear Regression search Search Quick search

Linear Neural Networks | 3.1 Linear Regression

A method for modeling the relationship between one or more independent variables and a dependent variable

Assumptions on Linear Regression

- All data are independently and identically distributed
 - All data are randomly sampled from the same distribution independently
 - Data not following IID: time-series data —



- Inductive bias (a bias required for generalization) for the model
- Well-behaved **noise** (following a Gaussian distribution)
 - Not expect to find a real-world dataset exactly equals the linear relationship due to factors such as measurement error
 - Thus, incorporating a noise term to account for such errors.

Model

price =
$$w_{\text{area}} \cdot \text{area} + w_{\text{age}} \cdot \text{age} + b$$
.

dependent variable y independent variables \mathbf{x} (output) (input) $\hat{y} = w_1 x_1 + \ldots + w_d x_d + b$.

$$\hat{y} = w_1 x_1 + \ldots + w_d x_d + b$$
.

$$\hat{y} = \mathbf{w}^\mathsf{T} \mathbf{x} + b$$
.

- w: weight
- b: bias (also called as offset or intercept)
- Strictly speaking, linear regression is an affine transformation
 - a kind of space transformation conserving linearity and parallelism including rotation, reflection, scaling, and others
- Goal: Choosing weights w and bias b which optimally fit the given set of pairs between input and output data



 $\begin{bmatrix} x' \\ y' \\ 1 \end{bmatrix} = \begin{bmatrix} a & b & e \\ c & d & f \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} x \\ y \\ 1 \end{bmatrix}$

Affine transformation

Linear Neural Networks | 3.1 Linear Regression

Elements to find optimal parameter w, b

- Loss function: a quality measure for some given model
- Stochastic Gradient Descent: a procedure for updating the model to improve its quality

Loss function

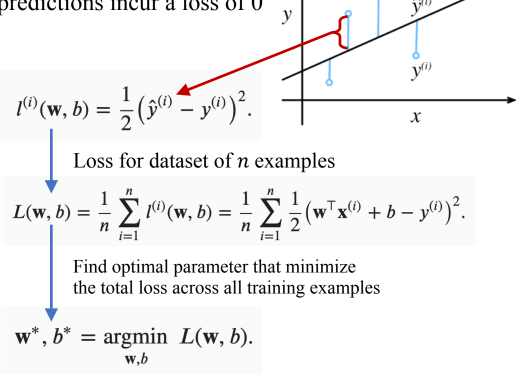
- Quantifying the distance between the real and predicted value of the target
 - Non-negative number where smaller values are better and perfect predictions incur a loss of 0
- Squared error
 - prediction for an data sample i, $\hat{y}^{(i)}$
 - sample target (true label) i, $y^{(i)}$

Normal Equation

Analytic solution for the simple liner regression problem

$$\mathbf{y} = X\mathbf{\Theta} + \mathbf{e} \xrightarrow{\text{error}} \sum_{j=1}^{n} \epsilon_{j}^{2} = \mathbf{e}^{T} \mathbf{e} = (\mathbf{y} - X\mathbf{\Theta})^{T} (\mathbf{y} - X\mathbf{\Theta})$$

$$\mathbf{e} = \mathbf{y} - X\mathbf{\Theta} \xrightarrow{\text{Minimize}} \sup_{\mathbf{y} = \mathbf{y} = \mathbf$$



Linear Neural Networks | 3.1 Linear Regression

Limitations of Normal Equation

- Hard to calculate a inverse matrix for large-scale data or features, $O(n^{2.376})$
- No inverse matrix can be existed if collinearity between features exists (singular matrix)

Gradient Descent

- Iteratively reducing the error by updating the parameters in the direction that incrementally lowers the loss function
- Taking the derivative of the loss function, which is an average of the losses computed on every single example in the dataset
- Stochastic Gradient Descent \longrightarrow for i = 1 to $n : \{ \theta_j^{t+1} = \theta_j^t \alpha(\Theta^T x_i y_i) x_i^{(j)} \text{ for every } j \}$
 - Updating parameters for a gradient of every sampled data
 - Pros: fast calculation speed, small memory requirement
 - Cons: non-stable (largely fluctuating) learning procedure, higher possibility to fall into local minimum
- Batch Gradient Descent

 $\theta_j^{t+1} = \theta_j^t - \alpha \sum_{i=1}^n (\Theta^T \mathbf{x}_i - y_i) \, x_i^{(j)} \quad \text{for every } j$

- update parameters for the mean of calculated gradients for all data
- Pros: converged into global minimum
- Cons: low calculation speed, large memory requirement
- Mini Batch Gradient Descent
 - update parameters for the mean of calculated gradients for a subset of given data



Mini Batch Gradient Descent

- Initializing the values of the model parameters, typically at random
- Randomly sampling a minibatch \mathcal{B} consisting of a fixed number of training examples
 - $|\mathcal{B}|$ represents the number of examples in each minibatch
- Updating the parameters in the direction of the negative gradient
 - ∂_i denotes the partial derivative of parameter I
 - η denotes the learning rate

$$\mathbf{w} \leftarrow \mathbf{w} - \frac{\eta}{|\mathcal{B}|} \sum_{i \in \mathcal{B}} \partial_{\mathbf{w}} l^{(i)}(\mathbf{w}, b) = \mathbf{w} - \frac{\eta}{|\mathcal{B}|} \sum_{i \in \mathcal{B}} \mathbf{x}^{(i)} \left(\mathbf{w}^{\mathsf{T}} \mathbf{x}^{(i)} + b - y^{(i)} \right),$$
$$b \leftarrow b - \frac{\eta}{|\mathcal{B}|} \sum_{i \in \mathcal{B}} \partial_{b} l^{(i)}(\mathbf{w}, b) = b - \frac{\eta}{|\mathcal{B}|} \sum_{i \in \mathcal{B}} \left(\mathbf{w}^{\mathsf{T}} \mathbf{x}^{(i)} + b - y^{(i)} \right).$$

Hyperparameter Tuning

- These parameters that are **tunable** but **not updated in the training loop**
- Typically adjust hyperparameters based on the results of the training loop as assessed on a separate validation dataset
- $|\mathcal{B}|$ and η for the simple linear regression model

Stopping Training

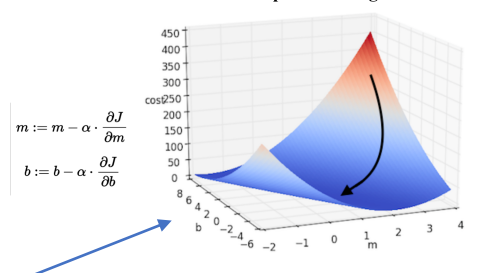
- Finishing training after certain number of iterations or until some stopping criteria met
- The trained parameters will not exact minimizers of the loss because it cannot achieve it exactly in a finite number of steps

Mini Batch Gradient Descent

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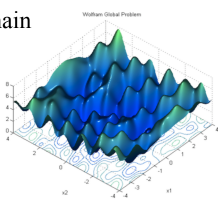
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Loss surface of a simple linear regression model



Loss function of Linear Regression and Deep Neural Networks

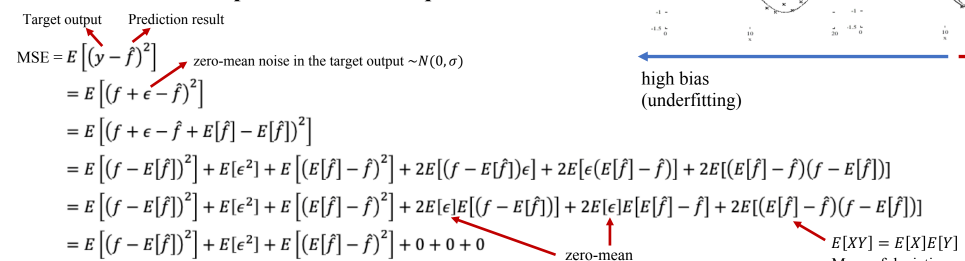
- Linear regression works for a learning problem where there is only one minimum over the entire domain
 - The bowl-shaped loss-function
 - Model with simple inductive bias → model with **high bias error** for complex data
- DNN model contains loss surfaces with many minima
 - The loss function of DNN is not bowl-shaped and not convex (much more complex)
 - Model with complex inductive bias → model with **low bias error** for complex data



Bias-Variance Trade-off

- Kinds of prediction error
 - Bias error: generated due to erroneous hypothesis on a model
 - high bias = underfitting
 - Variance error: from sensitivity to small fluctuations on different input data
 - high variance = overfitting
 - Irreducible error: noise in data

Bias-variance decomposition of mean squared error



Bias² Irreducible Variance Fluctuation (variance) of the prediction results

Error between true result and expected result from model

Variance of the random noise, $E[(\epsilon - 0)^2] = \sigma^2$

Mean of deviation = 0

high variance

(overfitting)

Model complexity

Statistical Interpretation of Linear Regression

• Dose least-squares loss function find actual optimal parameters of a linear regression model?

MLE for weight parameters of a linear regression model with gaussian noise equals Minimizing MSE of the model

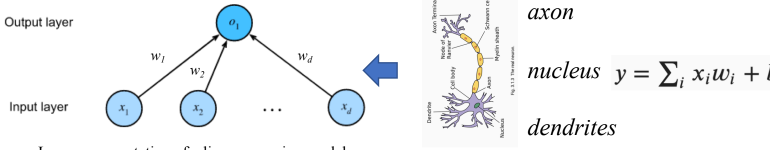
$$y_{i} = \Theta \mathbb{X}_{i} + \underbrace{\epsilon_{i}}_{\text{Gaussian noise}} p(\epsilon_{i}) = \underbrace{\frac{1}{\sqrt{2\pi}\sigma}} \exp\left(-\frac{\epsilon_{i}^{2}}{2\sigma^{2}}\right) \\ p(y_{i}|\mathbb{X}_{i};\theta) = \underbrace{\frac{1}{\sqrt{2\pi}\sigma}} \exp\left(-\frac{(y_{i}-\Theta \mathbb{X}_{i})^{2}}{2\sigma^{2}}\right) \\ \log L(\Theta) = \log \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y_{i}-\Theta \mathbb{X}_{i})^{2}}{2\sigma^{2}}\right) \\ \log L(\Theta) = \log \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y_{i}-\Theta \mathbb{X}_{i})^{2}}{2\sigma^{2}}\right) = \sum_{i=1}^{n} \log \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y_{i}-\Theta \mathbb{X}_{i})^{2}}{2\sigma^{2}}\right) \\ \text{Maximizing log likelihood} = n \log \frac{1}{\sqrt{2\pi}\sigma} - \frac{1}{\sigma^{2}} \frac{1}{2} \sum_{i=1}^{n} (y_{i} - \Theta^{T} \mathbb{X}_{i})^{2} \\ \text{Minimizing Squared error}$$

Interpretation of LR model in a perspective on Deep Neural Network

 σ is some fixed constant

Linear regression model is A kind of fully-connected layer or dense layer

• Every input is connected to every output



Layer representation of a liner regression model

$$o_1 = w_1 x_1 + w_2 x_2 + \dots + w_d x_d$$

Linear Neural Networks

Further material for following chapters

https://github.com/howawindelu/dive-into-deep-learning/blob/master/week3/week3_1_implementation_lukeshin.ipynb

- 3.1.2. Vectorization for Speed
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