Introduction to Modern AI Week 2: Supervised Learning I - Differentiable Parametric Models

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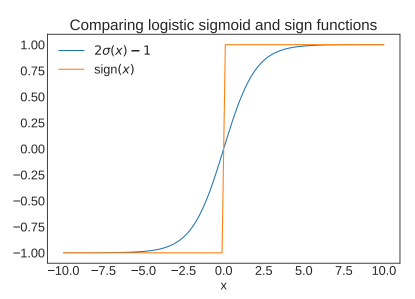
Topics we will cover this week

- In this week we explore more complex models of the form $f(x; \theta)$
- Mainly in supervised learning context, but many results will apply to other learning paradigms
- As we move through the course $f(x; \theta)$ will become more complicated (expressive), but it will retain two important properties
 - ullet it will be described in terms of parameters $oldsymbol{ heta}$
 - it will be differentiable, i.e. $\nabla_{\theta} f(x; \theta)$ will exist and be computable
- Main goal of this week is to introduce neural networks and gradient descent
- First, we will warm-up with the Perceptron
- Next week we will look at non-parametric models such as k-nearest neighbors and decision trees

- Historically significant model + algorithm for binary classification
- Useful to formulate it using the convention that $y \in \{-1, +1\}$
- Precursor to the sophisticated neural networks we use today
- Similar to logistic regression
- Linear model (absorbing bias into w):

$$f(\mathbf{x}; \mathbf{w}) = \operatorname{sign}(\mathbf{w}^T \mathbf{x}) = \begin{cases} -1, & \mathbf{w}^T \mathbf{x} < 0 \\ 1, & \mathbf{w}^T \mathbf{x} > 0 \end{cases}$$

 Unlike logistic regression, the perceptron just predicts a class, not a probability



Algorithm 1 Perceptron Learning Algorithm

- 1: initialize weights w
- 2: while not converged do
- 3: **for** i = 1, ..., N **do**
- 4: compute model prediction $\hat{y}_i = f(\mathbf{x}_i; \mathbf{w})$
- 5: $\mathbf{w} = \frac{\lambda}{2}(\hat{y}_i y_i)\mathbf{x}_i$
- 6: end for
- 7: end while

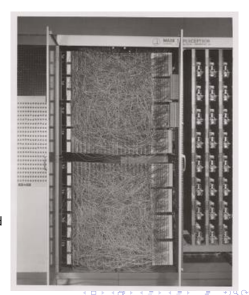
Interpretation:

- if prediction is correct, don't change weight vector
- if prediction is wrong, increase/decrease weight vector proportional to \mathbf{x}_i

Show Example

- The Perceptron learning algorithm is trying to find a separating hyper-plane
- If the data is not linearly separable, then the algorithm will not converge
- There are many solutions when the data is separable, and the algorithm doesn't necessarily pick the best one

- The Perceptron was actually built as a piece of specialized hardware (Mark I Perceptron)
- That's cool... but why are you telling us this?
 - Non-linear generalizations of Perceptrons are the simplest Artificial Neural Networks (ANNs)
 - The learning algorithm is actually stochastic gradient descent
 - They are closely related to a powerful set of models called Support Vector Machines (SVMs)



(Artificial) Neural Networks

The Need For More Expressive Models

- Both linear regression and logistic regression are linear (or log-linear) models
- This property severely constrains the types of functions they can fit
- It would be desirable to have models capable of learning arbitrarily complicated functions
 - Requires flexible families of models (i.e., NNs)
 - Requires general learning algorithm/framework (i.e., gradient descent)

Basis Expansions

Recall linear regression model:

$$f(\mathbf{x}; \mathbf{w}, b) = \mathbf{w}^T \mathbf{x} + b$$

- How can this be improved to incorporate non-linearities?
 - Suppose we knew that the probability explicitly depended on $(x_4)^2$, in addition to a linear dependence on x?
 - Just add it in!:

$$\mathbf{x}' = (x_1, x_2, ..., x_p, (x_4)^2)$$

 $\mathbf{w}' = (w_1, w_2, ..., w_p, w_{p+1})$

- Model remains linear in enlarged x' space!
- Suppose it also depends on $sin(x_2)$?
- Just add it in!:

$$\mathbf{x}' = (x_1, x_2, ..., x_p, (x_4)^2, \sin(x_2))$$

 $\mathbf{w}' = (w_1, w_2, ..., w_p, w_{p+1}, w_{p+2})$

Basis Expansions

- This basic idea can be extended to incorporate arbitrary non-linearities in a systematic fashion
- Suppose we identify a set of M useful transformations of the original input (or feature) space $\mathbf{x} \colon h_m(\mathbf{x}) \colon \mathbb{R}^p \to \mathbb{R}$, for m = 1, ..., M
- Simply build a linear model in the transformed feature space:

$$f(\mathbf{x};\boldsymbol{\theta}) = \sum_{m=1}^{M} w_m h_m(\mathbf{x}) + b$$

- Issues with this approach
 - ullet Poor scaling: p^d independent terms (and parameters) for d-degree polynomial
 - Inefficient way to make model more flexible unless we have special knowledge of the non-linearities

• Basic idea is very simple: build expressive model $f(x; \theta)$ using simpler modular components called layers:

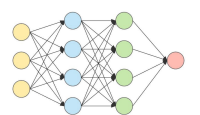
$$f = f^L \circ f^{L-1} \circ \dots \circ f^1$$

- here \circ means "compose", i.e. $(f \circ g)(x) = f(g(x))$
- the output of layer $f^{(\ell)}$ becomes the input to layer $f^{(\ell+1)}$
- As we will see, f has the structure of a bunch of simple processing units connected through a network - loose inspiration from (real) neural networks
- This lecture: simple, "vanilla" neural network, aka multi-layer perceptron (MLP)
- In later lectures we will learn about many sophisticated variants and extensions

• Neural networks are mathematical models, $f(x; \theta)$, where

$$f=f^L\circ f^{L-1}\circ ...\circ f^1$$

- Can also be described graphically
- Circles represent both variables and a computation, such as
 y = g(w₁ x₁ + w₂ x₂ + w₃ x₃ + b)
- Computation involves add/multiply and a possible non-linearity, called the activation function g
- Lines represent flow of information
- Network is organized into layers, input, hidden (×2), and output



Here is how to "translate" the diagram

Input (yellow dots):

$$\boldsymbol{x}^{(0)} = (x_1^{(0)}, x_2^{(0)}, x_3^{(0)})$$

Hidden Layer 1 (blue dots):

$$\mathbf{x}^{(1)} = (x_1^{(1)}, x_2^{(1)}, x_3^{(1)}, x_4^{(1)}) = \mathbf{g}^{(1)} \left(\mathbf{w}^{(1)} \mathbf{x}^{(0)} + \mathbf{b}^{(1)} \right)$$

Hidden Layer 2 (green dots):

$$\mathbf{x}^{(2)} = (x_1^{(2)}, x_2^{(2)}, x_3^{(2)}, x_4^{(2)}) = \mathbf{g}^{(2)} \left(\mathbf{w}^{(2)} \mathbf{x}^{(1)} + \mathbf{b}^{(2)} \right)$$

Output Layer (red dot):

$$\mathbf{x}^{(3)} = (x_1^{(3)}) = g^{(3)} \left(\mathbf{w}^{(3)} \mathbf{x}^{(2)} + \mathbf{b}^{(3)} \right)$$

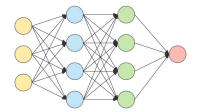


Putting it all together: $y = x_1^{(3)} = f(x; \theta)$, with

$$x_1^{(3)} = g^{(3)} \left(\mathbf{w}^{(3)} \left(g^{(2)} \left(\mathbf{w}^{(2)} \left(g^{(1)} \left(\mathbf{w}^{(1)} \mathbf{x}^{(0)} + \mathbf{b}^{(1)} \right) \right) + \mathbf{b}^{(2)} \right) \right) + \mathbf{b}^{(3)} \right)$$

Some comments

- The input and output dimensions are fixed by the problem
- Hidden layer dimensions are unconstrained
- Number of hidden layers is unconstrained
- Each layer can have a different activation function

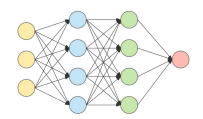


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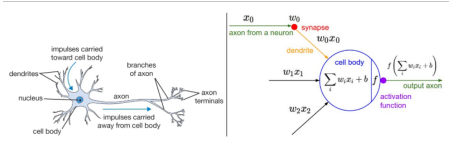
How many parameters does the network contain?

- bias vectors
 - $\dim(\mathbf{b}_1) : N_{h_1}$
 - $\dim(\mathbf{b}_2) : N_{h_2}$
 - dim(**b**₃): 1
- weight matrices
 - $\dim(\mathbf{w}^{(1)})$: $N_{h_1} \times p$
 - dim($\mathbf{w}^{(2)}$): $N_{h_2} \times N_{h_1}$
 - dim($\mathbf{w}^{(3)}$): $1 \times N_{h_2}$
- The model can be made arbitrarily large by increasing the number of units in either hidden layer



Activation Functions

- Non-linearities come entirely from the activation functions
- Some common choices for g:
 - Logistic sigmoid: $g(x) = \sigma(x)$
 - ReLU: $g(x) = \max(0, x)$
 - tanh: g(x) = tanh(x)
 - GELU: $g(x) = \frac{x}{2} \left(1 + \operatorname{erf} \left(\frac{x}{\sqrt{2}} \right) \right)$



A cartoon drawing of a biological neuron (left) and its mathematical model (right).

Universal Approximation Theorem

What class of functions can NNs represent?

- By adding more hidden layers (increasing depth) or adding more units to existing layers (increasing width) we can make the model more expressive (contain more parameters)
- More parameters loosely implies a capacity to approximate a larger class of functions
- The universal approximation theorems formally guarantee that, with enough parameters, NNs are capable of approximating any function with arbitrary accuracy
- There are many theorems for different activation functions and assumptions about model architecture (arbitrary width or depth)
- Theorems are not prescriptive they do not tell you have many units/layers you need to achieve a certain accuracy
- In the worst-case, you often need an exponentially large network
- How to train a NN is an entirely separate matter

Gradient Descent

Gradient Descent

- We've already seen that more complex models can be harder to train (fit) then simpler ones
 - The best-fit parameters for linear regression can be solved for in closed-form
 - Fitting logistic regression requires numerical optimization
 - No surprise: NNs will also require numerical optimization
- There are many optimization algorithms, which one(s) should we use?
 - Want a flexible method
 - No assumptions on geometry of loss function (i.e., cannot assume convexity)
 - Want it to be relatively fast/efficient
- Gradient Descent (GD) and variations are the dominant method used for training NNs

Gradient Descent

- GD is a very simple algorithm that only requires local information:
 - Starting at a given point, find the direction of steepest descent (given by the gradient vector)
 - Then take a small step in this direction
 - Repeat
- Think of a blind man trying to get to the bottom of a hill



Algorithm 2 Gradient Descent

1: function to be minimized: f(x)

2: set learning rate α

3: initialize variable x

4: while not converged do

5:
$$\mathbf{x} \leftarrow \mathbf{x} - \alpha \nabla f(\mathbf{x})$$

6: end while

Example: Rosenbrock (banana) function

$$f(x,y) = (1-x)^2 + 10(y-x^2)^2$$

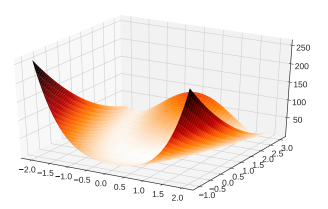
The gradient is

$$\nabla f = \begin{bmatrix} -2(1-x) - 10x(y-x^2) \\ 20(y-x^2) \end{bmatrix}.$$

• The gradient vanishes for the single point (x, y) = (1, 1), which is also the global minimum. At this point the function takes on the value f(1, 1) = 0.

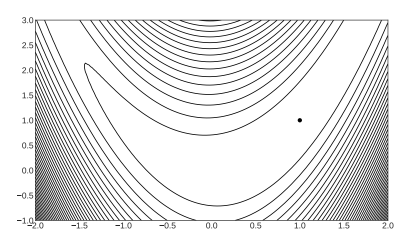
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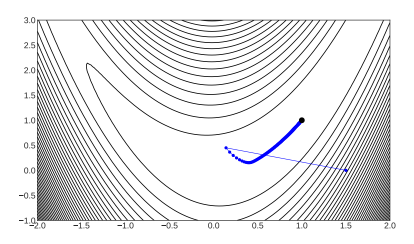
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,



• Example: Rosenbrock function

$$f(x,y) = (1-x)^2 + 10(y-x^2)^2$$



Some issues to keep in mind

- ullet The algorithm only converges if the learning rate lpha is small enough
- "Small enough" is problem dependent
- GD can get stuck in local minima
- Not well-suited for problems with widely-varying length scales (i.e., some very steep directions and some very shallow directions)

- To better understand GD, let's examine a simple optimization problem that we can solve exactly
- Consider a quadratic loss function:

$$L(\mathbf{w}) = \frac{1}{2} \mathbf{w}^T \mathbf{A} \mathbf{w} - \mathbf{b}^T \mathbf{w}$$

- $w, b \in \mathbb{R}^d$, and $A \in \mathbb{R}^{d \times d}$. also assume A is symmetric and invertible, meaning A^{-1} exists
- Gradient is:

$$\nabla_{\boldsymbol{w}} L = \boldsymbol{A} \boldsymbol{w} - \boldsymbol{b}$$

• Optimal solution is then $\mathbf{w}^* = \mathbf{A}^{-1}\mathbf{b}$

- Quadratic loss function: $L(\mathbf{w}) = \frac{1}{2} \mathbf{w}^T \mathbf{A} \mathbf{w} \mathbf{b}^T \mathbf{w}$
- Gradient: $\nabla_{\boldsymbol{w}} L = \boldsymbol{A} \boldsymbol{w} \boldsymbol{b}$
- Optimal solution: $\mathbf{w}^* = \mathbf{A}^{-1}\mathbf{b}$
- Diagonalize:

$$m{A} = m{O}^{-1} m{\Lambda} m{O} \,, \qquad m{\Lambda} = {\sf diag}(\lambda_1, \lambda_2, ..., \lambda_d) \,.$$
 $m{b}' = m{O} m{b} \,, \qquad m{x} = m{O} m{w} \,.$

Gradient descent update rule:

$$\mathbf{w}^{t+1} = \mathbf{w}^t - \alpha \nabla_{\mathbf{w}} L(\mathbf{w}^t) = \mathbf{w}^t - \alpha (\mathbf{A} \mathbf{w}^t - \mathbf{b})$$

Becomes:

$$\mathbf{x}^{t+1} = \mathbf{x}^t - \alpha \left(\mathbf{\Lambda} \mathbf{x}^t - \mathbf{b}' \right)$$

• In terms of components:

$$x_a^{t+1} = (1 - \alpha \lambda_a) x_a^t + \alpha b_a'$$



• GD update rule in x variable:

$$x_a^{t+1} = (1 - \alpha \lambda_a) x_a^t + \alpha b_a'$$

Can "unroll":

$$x_a^{t+1} = (1 - \alpha \lambda_a) \left((1 - \alpha \lambda_a) x_a^{t-1} + \alpha b_a' \right) + \alpha b_a'$$
$$= (1 - \alpha \lambda_a)^2 x_a^{t-1} + (1 + (1 - \alpha \lambda_a)) \alpha b_a'$$

Continuing:

$$\mathbf{x}_{\mathsf{a}}^{t+1} = (1 - \alpha \lambda_{\mathsf{a}})^{t+1} \, \mathbf{x}_{\mathsf{a}}^{0} + \alpha b_{\mathsf{a}}' \sum_{k=0}^{t} (1 - \alpha \lambda_{\mathsf{a}})$$

• For this to converge as $t \to \infty$, require $|1 - \alpha \lambda_a| < 1$. Then first term vanishes and second term is a geometric series, can be summed to give $\mathbf{x}^* = \mathbf{O}\mathbf{w}^* = \mathbf{\Lambda}^{-1}\mathbf{b}$:

$$x_{a}^{*} = \lim_{t \to \infty} x_{a}^{t} = \alpha b_{a}' \left(\frac{1}{1 - (1 - \alpha \lambda_{a})} \right) = \frac{b_{a}'}{\lambda_{a}}$$

To summarize

- Optimization problem for d-dim vector w decomposes into d separate
 1-dim optimization problems if we work in the eigenbasis
- Convergence requires $|1 \alpha \lambda_a| < 1$
- Corresponds to $0 < \alpha \lambda_{\it a} < 2$
 - requires all eigenvalues to have same sign positive for a (global) minimum
 - then $\alpha > 0$
 - $\alpha <$ 0, $\lambda_a <$ 0 corresponds to gradient ascent
- Convergence in this case is exponential
- But convergence rate is different for each eigen-direction, overall rate is limited by worst eigenvalue

Stochastic Gradient Descent Algorithm

Let's use gradient descent to minimize a loss function

Loss function is averaged over training set:

$$loss(\boldsymbol{\theta}) = \frac{1}{N} \sum_{i=1}^{N} \ell(\mathbf{x}_i, y_i; \boldsymbol{\theta})$$

- ullet could be the sum of squared errors (SSE), or binary cross entropy (BCE), or really any loss
- Gradient of loss will also be averaged:

$$\nabla_{\theta} loss(\theta) = \frac{1}{N} \sum_{i=1}^{N} \nabla_{\theta} \ell(\mathbf{x}_i, y_i; \theta)$$

Stochastic Gradient Descent Algorithm

$$\nabla_{\theta} loss(\theta) = \frac{1}{N} \sum_{i=1}^{N} \nabla_{\theta} \ell(\mathbf{x}_i, y_i; \theta)$$

- This expression is not always convenient to work with
- GD requires many iterations, and each iteration requires that the gradient be computed N times
- N is often large (and we want N to be as large as possible)
- Idea: let's cut a corner and instead of averaging the gradient over all N data points, let's average it over a "mini-batch" of $N_B \ll N$ data points
- Each GD update can use a different, randomly chosen batch
- Note on terminology:
 - $N_B = N$: gradient descent
 - $1 < N_B < N$: mini-batch stochastic gradient descent
 - $N_B = 1$: stochastic gradient descent (aka on-line learning)

Introduction to Backpropagation

- ullet To use SGD, we must be able to compute the gradient of the loss function $abla_{ heta}\ell$
- Let's consider the MSE loss for a simple NN with 1 hidden layer:

$$\ell(\mathbf{x}_i, y_i; \boldsymbol{\theta}) = (y_i - f(\mathbf{x}; \boldsymbol{\theta}))^2,$$

$$f(\mathbf{x}; \boldsymbol{\theta}) = g^{(2)} \left(\mathbf{w}^{(2)} \left(g^{(1)} \left(\mathbf{w}^{(1)} \mathbf{x}^{(0)} + \boldsymbol{b}^{(1)} \right) \right) + \boldsymbol{b}^{(2)} \right)$$

- Need to compute $\nabla_{\mathbf{w}^{(1)}} f$, $\nabla_{\mathbf{w}^{(2)}} f$, $\nabla_{b^{(1)}} f$, $\nabla_{b^{(2)}} f$
- This is do-able, but a bit of a pain
- What if we later want to tweak the network? What if we want to consider a large model with 10's or 100's of layers?
- It's desirable to have an automated procedure to compute the gradient for us

Introduction to Backpropagation

- Suppose we have functions f, g, and the composition $h = g \circ f$
- The chain rule let's use calculate the derivative h':

$$h' = (g \circ f)' = (g' \circ f)f'$$

or,

$$h'(x) = g'(f(x))f'(x)$$

• Example:

$$\frac{d}{dx}\sin(x)^2 = 2\sin x \cos x$$

- $f(x) = \sin(x), g(x) = x^2$
- $f'(x) = \cos(x), g'(x) = 2x$
- $h'(x) = (2\sin x)\cos x$

Introduction to Backpropagation

• What about functions built using more than 1 composition?

$$f = f^{(L)} \circ f^{(L-1)} \circ ... \circ f^{(1)}$$

Apply the chain rule iteratively:

$$f = f^{(L)} \circ F^{(L-1)}, \qquad F^{(L-1)} = f^{(L-1)} \circ \dots \circ f^{(1)}$$

 $f' = (f^{(L)'} \circ F^{(L-1)}) F^{(L-1)'}$

- and so on for $F^{(L-1)'}$, and then $F^{(L-2)'}$, ...
- This "algorithm" can be used to compute the gradient of an arbitrarily complicated neural net
- Q: why is it called backpropagation?
- In week 4 we'll discuss how this is implemented in modern deep learning software libraries using automatic differentiation

Recap

- Vanilla, Feed-Forward Neural Networks
- (Stochastic) Gradient Descent
- Backpropagation