The value function approximation structure for today closely follows much of David Silver’s Lecture 6. For additional reading please see SB 2018 Sections 9.3, 9.6-9.7. The deep learning slides come almost exclusively from Ruslan Salakhutdinov’s class, and Hugo Larochelle’s class (and with thanks to Zico Kolter also for slide inspiration). The slides in my standard style format in the deep learning section are my own.
Important Information About Homework 2

- Homework 2 will now be due on Saturday February 10 (instead of February 7)
- We are making this change to try to give some background on deep learning, give people enough time to do homework 2, and still give people time to study for the midterm on February 14
- We will release the homework this week
- You will be able to start on some aspects of the homework this week, but we will be covering DQN which is the largest part, on Monday
- We will also be providing optional tutorial sessions on tensorflow

Tuesday 4-6pm - see pizza for details
Last time: Control (making decisions) without a model of how the world works

This time: Value function approximation and deep learning

Next time: Deep reinforcement learning
Last time: how to learn a good policy from experience

So far, have been assuming we can represent the value function or state-action value function as a vector

- Tabular representation

Many real world problems have enormous state and/or action spaces

- Tabular representation is insufficient
Recall: Reinforcement Learning Involves

- Optimization
- Delayed consequences
- Exploration
- Generalization
Today: Focus on Generalization

- Optimization
- Delayed consequences
- Exploration
- Generalization
Table of Contents

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2 VFA for Prediction

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Value Function Approximation (VFA)

- Represent a (state-action/state) value function with a parameterized function instead of a table.

\[ f(w, s) \]

\[ \hat{v}(S, w) \]

\[ \hat{q}(s, a, w) \]

\[ \hat{q}(s, a_1, w) \]

\[ \hat{q}(s, a_2, w) \]
Motivation for VFA

- Don’t want to have to explicitly store or learn for every single state a
  - Dynamics or reward model
  - Value
  - State-action value
  - Policy
- Want more compact representation that generalizes across state or states and actions
Benefits of Generalization

- Reduce memory needed to store \((P, R)/V/Q/\pi\)
- Reduce computation needed to compute \((P, R)/V/Q/\pi\)
- Reduce experience needed to find a good \(P, R/V/Q/\pi\)
Value Function Approximation (VFA)

- Represent a (state-action/state) value function with a parameterized function instead of a table.

- Which function approximator?
Many possible function approximators including
- Linear combinations of features
- Neural networks
- Decision trees
- Nearest neighbors
- Fourier / wavelet bases

In this class we will focus on function approximators that are differentiable (Why?)

Two very popular classes of differentiable function approximators
- Linear feature representations (Today)
- Neural networks (Today and next lecture)
Consider a function $J(w)$ that is a differentiable function of a parameter vector $w$.

Goal is to find parameter $w$ that minimizes $J$.

The gradient of $J(w)$ is

$$
\nabla_w J(w) = \left[ \frac{\partial J(w)}{\partial w_1}, \frac{\partial J(w)}{\partial w_2}, \ldots, \frac{\partial J(w)}{\partial w_n} \right]
$$

Local optimum of $J$

$$
w \leftarrow w - \alpha \nabla_w J(w)
$$

Shorthand

$$
\Delta w = \alpha \nabla_w J(w)
$$

$$
= -\frac{1}{2} \alpha J(w)
$$
to min $J$.
Value Function Approximation for Policy Evaluation with an Oracle

First consider if could query any state $s$ and an oracle would return the true value for $v^\pi(s)$.

The objective was to find the best approximate representation of $v^\pi$ given a particular parameterized function.
Stochastic Gradient Descent

- Goal: Find the parameter vector $\mathbf{w}$ that minimizes the loss between a true value function $v_\pi(s)$ and its approximation $\hat{v}$ as represented with a particular function class parameterized by $\mathbf{w}$.

- Generally use mean squared error and define the loss as

$$J(\mathbf{w}) = \mathbb{E}_\pi [(v_\pi(S) - \hat{v}(S, \mathbf{w}))^2]$$ (1)

- Can use gradient descent to find a local minimum

$$\Delta \mathbf{w} = -\frac{1}{2} \alpha \nabla \mathbf{w} J(\mathbf{w})$$ (2)

- Stochastic gradient descent (SGD) samples the gradient:

$$\nabla \mathbf{w} J(\mathbf{w}) = \mathbb{E}_\pi \left[ 2(v_\pi(S) - \hat{v}(S, \mathbf{w})) \cdot \nabla \mathbf{w} \hat{v}(s, \mathbf{w}) \right]$$

$$\Delta \mathbf{w} = \alpha \left( v_\pi(S) - \hat{v}(S, \mathbf{w}) \right) \nabla \mathbf{w} \hat{v}(s, \mathbf{w}) \quad \text{in particular,}$$

- Expected SGD is the same as the full gradient update
Don’t actually have access to an oracle to tell true $v_\pi(S)$ for any state $s$

Now consider how to do value function approximation for prediction / evaluation / policy evaluation without a model

Note: policy evaluation without a model is sometimes also called **passive reinforcement learning** with value function approximation

”passive” because not trying to learn the optimal decision policy
Recall model-free policy evaluation (Lecture 3)
- Following a fixed policy $\pi$ (or had access to prior data)
- Goal is to estimate $V^\pi$ and/or $Q^\pi$

Maintained a look up table to store estimates $V^\pi$ and/or $Q^\pi$

Updated these estimates after each episode (Monte Carlo methods) or after each step (TD methods)

Now: in value function approximation, change the estimate update step to include fitting the function approximator

\[
S \rightarrow \omega \rightarrow V \\
V = \omega
\]
Feature Vectors

- Use a feature vector to represent a state

\[ x(s) = \begin{pmatrix} x_1(s) \\ x_2(s) \\ \vdots \\ x_n(s) \end{pmatrix} \]  

(3)
Linear Value Function Approximation for Prediction With An Oracle

- Represent a value function (or state-action value function) for a particular policy with a weighted linear combination of features:
  \[ \hat{v}(S, \mathbf{w}) = \sum_{j=1}^{n} x_j(S) w_j = \mathbf{x}(S)^T \mathbf{w} \]

- Objective function is
  \[ J(\mathbf{w}) = \mathbb{E}_\pi[(v_\pi(S) - \hat{v}(S, \mathbf{w}))^2] \]

- Recall weight update is
  \[ \Delta \mathbf{w} = -\frac{1}{2} \alpha \nabla \mathbf{w} \ J(\mathbf{w}) \] (4)

- Update is:
  \[ \Delta \mathbf{w} = \alpha \left( v_\pi(s) - \hat{v}(s, \mathbf{w}) \right) \cdot \nabla \hat{v}(s, \mathbf{w}) \cdot \mathbf{x}(s) \]

- Update = step-size \times prediction error \times feature value
Monte Carlo Value Function Approximation

- Return $G_t$ is an unbiased but noisy sample of the true expected return $v_\pi(S_t)$
- Therefore can reduce MC VFA to doing supervised learning on a set of (state,return) pairs: $<S_1, G_1>, <S_2, G_2>, \ldots, <S_T, G_T>$
  - Susbstituting $G_t(S_t)$ for the true $v_\pi(S_t)$ when fitting the function approximator
- Concretely when using linear VFA for policy evaluation
  \[
  \Delta \mathbf{w} = \alpha(\mathbf{G}_t - \hat{v}(S_t, \mathbf{w})) \nabla \mathbf{w} \hat{v}(S_t, \mathbf{w}) \\
  = \alpha(G_t - \hat{v}(S_t, \mathbf{w})) \mathbf{x}(S_t) \tag{5}
  \]
- Note: $G_t$ may be a very noisy estimate of true return
MC Linear Value Function Approximation for Policy Evaluation

1: Initialize $w = 0, \text{Returns}(s) = 0 \forall (s, \pi), k = 1$
2: loop
3: Sample $k$-th episode $(s_k1, a_k1, r_k1, s_k2, \ldots, s_kL_k)$ given $\pi$
4: for $t = 1, \ldots, L_k$ do
5: \hspace{1em} if First visit to $(s)$ in episode $k$ then
6: \hspace{2em} Append $\sum_{j=t}^{L_k} r_{kj}$ to Returns($s_t$)
7: \hspace{1em} Update weights
8: \hspace{2em} $\Delta w = \alpha \left( \text{Returns}(s_t) - \hat{v}(s_t, w) \right) \cdot x(s_t)$
9: \hspace{1em} end if
10: end for
11: $k = k + 1$
12: end loop
Recall: Temporal Difference (TD(0)) Learning with a Look up Table

- Uses bootstrapping and sampling to approximate $V^\pi$
- Updates $V^\pi(s)$ after each transition $(s, a, r, s')$:

$$V^\pi(s) = V^\pi(s) + \alpha (r + \gamma V^\pi(s') - V^\pi(s))$$  \hspace{1cm} (7)

- Target is $r + \gamma V^\pi(s')$, a biased estimate of the true value $v^\pi(s)$
- Look up table represents value for each state with a separate table entry
Temporal Difference (TD(0)) Learning with Value Function Approximation

- Uses bootstrapping and sampling to approximate true $v^{\pi}$
- Updates estimate $V^{\pi}(s)$ after each transition $(s, a, r, s')$:

$$V^{\pi}(s) = V^{\pi}(s) + \alpha(r + \gamma V^{\pi}(s') - V^{\pi}(s)) \quad (8)$$

- Target is $r + \gamma V^{\pi}(s')$, a biased estimate of the true value $v^{\pi}(s)$
- In value function approximation, target is $r + \gamma \hat{v}^{\pi}(s')$, a biased and approximated estimate of the true value $v^{\pi}(s)$

- 3 forms of approximation:
  1) sampling an expectation
  2) bootstrapping
  3) function approximation
Temporal Difference (TD(0)) Learning with Value Function Approximation

- In value function approximation, target is $r + \gamma \hat{v}^\pi(s')$, a biased and approximated estimate of the true value $v^\pi(s)$.
- Supervised learning on a different set of data pairs:
  $< S_1, r_1 + \gamma \hat{v}^\pi(S_2, w) >, < S_2, r_2 + \gamma \hat{v}(S_3, w) >, \ldots$
Temporal Difference (TD(0)) Learning with Value Function Approximation

- In value function approximation, target is \( r + \gamma \hat{v}^\pi(s') \), a biased and approximated estimate of the true value \( v^\pi(s) \).
- Supervised learning on a different set of data pairs:
  \(< S_1, r_1 + \gamma \hat{v}^\pi(S_2, w) >, < S_2, r_2 + \gamma \hat{v}(S_3, w) >, \ldots \)
- In linear TD(0)

\[
\Delta w = \alpha (r + \gamma \hat{v}^\pi(s', w) - \hat{v}^\pi(s, w)) \nabla_w \hat{v}^\pi(s, w) \tag{9}
\]

\[
= \alpha (r + \gamma \hat{v}^\pi(s', w) - \hat{v}^\pi(s, w)) x(s) \tag{10}
\]
Define the mean squared error of a linear value function approximation for a particular policy $\pi$ relative to the true value as

$$MSVE(w) = \sum_{s \in S} d(s)(v^\pi(s) - \hat{v}^\pi(s, w))^2 \quad (11)$$

where

- $d(s)$: stationary distribution of $\pi$ in the true decision process
- $\hat{v}^\pi(s, w) = x(s)^T w$, a linear value function approximation

---

Define the mean squared error of a linear value function approximation for a particular policy $\pi$ relative to the true value as

$$MSVE(w) = \sum_{s \in S} d(s) (v^\pi(s) - \hat{v}^\pi(s, w))^2$$  \hspace{1cm} (12)$$

where

- $d(s)$: stationary distribution of $\pi$ in the true decision process
- $\hat{v}^\pi(s) = x(s)^T w$, a linear value function approximation

Monte Carlo policy evaluation with VFA converges to the weights $w_{MC}$ which has the minimum mean squared error possible:

$$MSVE(w_{MC}) = \min_w \sum_{s \in S} d(s) (v^\pi(s) - \hat{v}^\pi(s, w))^2$$  \hspace{1cm} (13)$$

---

Define the mean squared error of a linear value function approximation for a particular policy \( \pi \) relative to the true value as

\[
MSVE(w) = \sum_{s \in S} d(s)(v^\pi(s) - \hat{v}^\pi(s, w))^2
\]

where

\[
d(s): \text{stationary distribution of } \pi \text{ in the true decision process}
\]

\[
\hat{v}^\pi(s) = x(s)^T w, \text{ a linear value function approximation}
\]

TD(0) policy evaluation with VFA converges to weights \( w_{TD} \) which is within a constant factor of the minimum mean squared error possible:

\[
MSVE(w_{TD}) = \frac{1}{1 - \gamma} \min_w \sum_{s \in S} d(s)(v^\pi(s) - \hat{v}^\pi(s, w))^2
\]
Monte Carlo policy evaluation with VFA converges to the weights $w_{MC}$ which has the minimum mean squared error possible:

$$MSVE(w_{MC}) = \min_w \sum_{s \in S} d(s)(v^\pi(s) - \hat{v}^\pi(s, w))^2$$  \hspace{1cm} (16)$$

TD(0) policy evaluation with VFA converges to weights $w_{TD}$ which is within a constant factor of the minimum mean squared error possible:

$$MSVE(w_{TD}) = \frac{1}{1 - \gamma} \min_w \sum_{s \in S} d(s)(v^\pi(s) - \hat{v}^\pi(s, w))^2$$  \hspace{1cm} (17)$$

Check your understanding: if the VFA is a tabular representation (one feature for each state), what is the MSVE for MC and TD?  

---

4 ibed.
Does TD or MC converge faster to a fixed point?
Not (to my knowledge) definitively understood
Practically TD learning often converges faster to its fixed value function approximation point
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2. VFA for Prediction

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Use value function approximation to represent state-action values
\( \hat{q}^\pi(s, a, w) \approx q^\pi \)

Interleave
- Approximate policy evaluation using value function approximation
- Perform \(\epsilon\)-greedy policy improvement
Action-Value Function Approximation with an Oracle

- \( \hat{q}^\pi(s, a, \mathbf{w}) \approx q^\pi \)

- Minimize the mean-squared error between the true action-value function \( q^\pi(s, a) \) and the approximate action-value function:

  \[
  J(\mathbf{w}) = \mathbb{E}_\pi [(q^\pi(s, a) - \hat{q}^\pi(s, a, \mathbf{w}))^2] \tag{18}
  \]

- Use stochastic gradient descent to find a local minimum

  \[
  -\frac{1}{2} \nabla_{\mathbf{w}} J(\mathbf{w}) = \mathbb{E} [(q^\pi(s, a) - \hat{q}^\pi(s, a, \mathbf{w})) \nabla_{\mathbf{w}} \hat{q}^\pi(s, a, \mathbf{w})] \tag{19}
  \]

  \[
  \Delta(\mathbf{w}) = -\frac{1}{2} \alpha \nabla_{\mathbf{w}} J(\mathbf{w}) \tag{20}
  \]

- Stochastic gradient descent (SGD) samples the gradient

  \((s, a)\) pair
Linear State Action Value Function Approximation with an Oracle

- Use features to represent both the state and action

\[
x(s, a) = \begin{pmatrix}
x_1(s, a) \\
x_2(s, a) \\
\vdots \\
x_n(s, a)
\end{pmatrix}
\]  

(21)

- Represent state-action value function with a weighted linear combination of features

\[
\hat{q}(s, a, w) = x(s, a)^T w = \sum_{j=1}^{n} x_j(s, a) w_j
\]  

(22)

- Stochastic gradient descent update:

\[
\nabla_w J(w) = \nabla_w \mathbb{E}_\pi[(q^\pi(s, a) - \hat{q}^\pi(s, a, w))^2]
\]  

(23)
Similar to policy evaluation, true state-action value function for a state is unknown and so substitute a target value.

In Monte Carlo methods, use a return $G_t$ as a substitute target

$$\Delta w = \alpha (G_t - \hat{q}(s_t, a_t, w)) \nabla_w \hat{q}(s_t, a_t, w)$$ \hspace{1cm} (24)

For SARSA instead use a TD target $r + \gamma \hat{q}(s', a', w)$ which leverages the current function approximation value

$$\Delta w = \alpha (r + \gamma \hat{q}(s', a', w) - \hat{q}(s, a, w)) \nabla_w \hat{q}(s, a, w)$$ \hspace{1cm} (25)
Incremental Model-Free Control Approaches

- Similar to policy evaluation, true state-action value function for a state is unknown and so substitute a target value
- In Monte Carlo methods, use a return $G_t$ as a substitute target

$$\Delta \mathbf{w} = \alpha (G_t - \hat{q}(s_t, a_t, \mathbf{w})) \nabla_{\mathbf{w}} \hat{q}(s_t, a_t, \mathbf{w}) \quad (26)$$

- For SARSA instead use a TD target $r + \gamma \hat{q}(s', a', \mathbf{w})$ which leverages the current function approximation value

$$\Delta \mathbf{w} = \alpha (r + \gamma \hat{q}(s', a', \mathbf{w}) - \hat{q}(s, a, \mathbf{w})) \nabla_{\mathbf{w}} \hat{q}(s, a, \mathbf{w}) \quad (27)$$

- For Q-learning instead use a TD target $r + \gamma \max_{a'} \hat{q}(s', a', \mathbf{w})$ which leverages the max of the current function approximation value

$$\Delta \mathbf{w} = \alpha (r + \gamma \max_{a'} \hat{q}(s', a', \mathbf{w}) - \hat{q}(s, a, \mathbf{w})) \nabla_{\mathbf{w}} \hat{q}(s, a, \mathbf{w}) \quad (28)$$
Convergence of TD Methods with VFA

- TD with value function approximation is not following the gradient of an objective function.
- Informally, updates involve doing an (approximate) Bellman backup followed by best trying to fit underlying value function to a particular feature representation.
- Bellman operators are contractions, but value function approximation fitting can be an expansion.

\[
\begin{bmatrix}
V_1(s_1) \\
\vdots \\
V_1(s_N)
\end{bmatrix}
= 
\begin{bmatrix}
V_2(s_1) \\
\vdots \\
V_2(s_N)
\end{bmatrix}
\]

\[
\|BV_1 - BV_2\|_\infty \leq \|V_1 - V_2\|_\infty
\]

\[
\sum V_1 - \sum V_2
\]
## Convergence of Control Methods with VFA

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Tabular</th>
<th>Linear VFA</th>
<th>Nonlinear VFA</th>
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<tr>
<td>Monte-Carlo Control</td>
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<td>✓</td>
<td>✓</td>
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<tr>
<td>Sarsa</td>
<td>✓</td>
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<tr>
<td>Q-learning</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

(✓) checks ✔️

deep learning
Other Function Approximators

- Linear value function approximators often work well given the right set of features.
- But can require carefully hand designing that feature set.
- An alternative is to use a much richer function approximation class that is able to directly go from states without requiring an explicit specification of features.

- Local representations including Kernel based approaches have some appealing properties (including convergence results under certain cases) but can’t typically scale well to enormous spaces and datasets.
- Alternative is to leverage huge recent success in using deep neural networks.
Today: a brief introduction to deep neural networks

Definitions

Power of deep neural networks

- Neural networks / distributed representations vs kernel / local representations
- Universal function approximator
- Deep neural networks vs shallow neural networks

How to train neural nets
Deep Neural Networks

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How to train neural nets
Feedforward Neural Networks

- Definition of Neural Networks
  - Forward propagation
  - Types of units
  - Capacity of neural networks

- How to train neural nets:
  - Loss function
  - Backpropagation with gradient descent

- More recent techniques:
  - Dropout
  - Batch normalization
  - Unsupervised Pre-training
Artificial Neuron

• Neuron pre-activation (or input activation):

\[ a(x) = b + \sum_i w_i x_i = b + w^\top x \]

• Neuron output activation:

\[ h(x) = g(a(x)) = g(b + \sum_i w_i x_i) \]

where

- \( W \) are the weights (parameters)
- \( b \) is the bias term
- \( g(\cdot) \) is called the activation function
Single Hidden Layer Neural Net

- Hidden layer pre-activation:
  \[ a(x) = b^{(1)} + W^{(1)}x \]
  \[ (a(x)_i = b^{(1)}_i + \sum_j W^{(1)}_{i,j} x_j) \]

- Hidden layer activation:
  \[ h(x) = g(a(x)) \]

- Output layer activation:
  \[ f(x) = \sigma\left( b^{(2)} + W^{(2)^T} h^{(1)}x \right) \]

Output activation function
Artificial Neuron

• Output activation of the neuron:

\[ h(x) = g(a(x)) = g(b + \sum_i w_i x_i) \]

Range is determined by \( g(\cdot) \)

Bias only changes the position of the riff

(from Pascal Vincent's slides)
Single Hidden Layer Neural Net

- Hidden layer pre-activation:
  \[ a(x) = b^{(1)} + W^{(1)}x \]
  \[ (a(x)_i = b^{(1)}_i + \sum_j W^{(1)}_{i,j} x_j) \]

- Hidden layer activation:
  \[ h(x) = g(a(x)) \]

- Output layer activation:
  \[ f(x) = o \left( b^{(2)} + W^{(2)\top} h^{(1)}x \right) \]

\[ \text{Output activation function} \]
Activation Function

• Sigmoid activation function:
  - Squashes the neuron’s output between 0 and 1
  - Always positive
  - Bounded
  - Strictly Increasing

\[
g(a) = \text{sigm}(a) = \frac{1}{1 + \exp(-a)}
\]
Activation Function

• Rectified linear (ReLU) activation function:

  - Bounded below by 0 (always non-negative)
  - Tends to produce units with sparse activities
  - Not upper bounded
  - Strictly increasing

\[ g(a) = \text{recln}(a) = \max(0, a) \]
Consider a network with L hidden layers.

- Layer pre-activation for $k > 0$
  \[ a^{(k)}(x) = b^{(k)} + W^{(k)} h^{(k-1)}(x) \]

- Hidden layer activation from 1 to L:
  \[ h^{(k)}(x) = g(a^{(k)}(x)) \]

- Output layer activation ($k = L + 1$):
  \[ h^{(L+1)}(x) = o(a^{(L+1)}(x)) = f(x) \]
  \[ (h^{(0)}(x) = x) \]
Deep Neural Networks

- Today: a brief introduction to deep neural networks
- Definitions
- **Power of deep neural networks**
  - Neural networks / distributed representations vs kernel / local representations
  - Universal function approximator
  - Deep neural networks vs shallow neural networks
- How to train neural nets
Local vs. Distributed Representations

- Clustering, Nearest Neighbors, RBF SVM, local density estimators
  - Parameters for each region.
  - # of regions is linear with # of parameters.

- RBMs, Factor models, PCA, Sparse Coding, Deep models

(Bengio, 2009, Foundations and Trends in Machine Learning)
Local vs. Distributed Representations

- Clustering, Nearest Neighbors, RBF SVM, local density estimators
  - Parameters for each region.
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Learned prototypes
Local vs. Distributed Representations

- Clustering, Nearest Neighbors, RBF SVM, local density estimators
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Learned prototypes
Local vs. Distributed Representations

- Clustering, Nearest Neighbors, RBF SVM, local density estimators
  - Parameters for each region.
  - # of regions is linear with # of parameters.

- RBMs, Factor models, PCA, Sparse Coding, Deep models
  - Each parameter affects many regions, not just local.
  - # of regions grows (roughly) exponentially in # of parameters.
Capacity of Neural Nets

• Consider a single layer neural network
Capacity of Neural Nets

• Consider a single layer neural network

(from Pascal Vincent's slides)
Universal Approximation

• Universal Approximation Theorem (Hornik, 1991):
  
  – “a single hidden layer neural network with a linear output unit can approximate any continuous function arbitrarily well, given enough hidden units”

• This applies for sigmoid, tanh and many other activation functions.

• However, this does not mean that there is a learning algorithm that can find the necessary parameter values.
1 hidden layer neural networks are already a universal function approximator.

Implies the expressive power of deep networks are no larger than shallow networks.
- There always exists a shallow network that can represent any function representable by a deep (multi-layer) neural network.

But there can be cases where deep networks may be exponentially more compact than shallow networks in terms of number of nodes required to represent a function.

This has substantial implications for memory, computation and data efficiency.

Empirically often deep networks outperform shallower alternatives.
Today: a brief introduction to deep neural networks

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Power of deep neural networks
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How to train neural nets
Feedforward Neural Networks

- How neural networks predict $f(x)$ given an input $x$:
  - Forward propagation
  - Types of units
  - Capacity of neural networks

- How to train neural nets:
  - Loss function
  - Backpropagation with gradient descent

- More recent techniques:
  - Dropout
  - Batch normalization
  - Unsupervised Pre-training
Training

- **Empirical Risk Minimization:**

\[
\arg \min_{\theta} \frac{1}{T} \sum_{t} l(f(x^{(t)}; \theta), y^{(t)}) + \lambda \Omega(\theta)
\]

- Learning is cast as optimization.

  - For classification problems, we would like to minimize classification error.
Stochastic Gradient Descend

- Perform updates after seeing each example:
  - Initialize: \( \theta \equiv \{ W^{(1)}, b^{(1)}, \ldots, W^{(L+1)}, b^{(L+1)} \} \)
  - For t=1:T
    - for each training example \((x^{(t)}, y^{(t)})\)
      \[
      \Delta = -\nabla_\theta l(f(x^{(t)}; \theta), y^{(t)}) - \lambda \nabla_\theta \Omega(\theta)
      \]
      \[
      \theta \leftarrow \theta + \alpha \Delta
      \]

- To train a neural net, we need:
  - Loss function: \( l(f(x^{(t)}; \theta), y^{(t)}) \)
  - A procedure to compute gradients: \( \nabla_\theta l(f(x^{(t)}; \theta), y^{(t)}) \)
  - Regularizer and its gradient: \( \Omega(\theta), \nabla_\theta \Omega(\theta) \)
Computational Flow Graph

- Forward propagation can be represented as an acyclic flow graph

- Forward propagation can be implemented in a modular way:
  - Each box can be an object with an `fprop` method, that computes the value of the box given its children
  - Calling the `fprop` method of each box in the right order yields forward propagation
• Each object also has a `bprop` method

  - it computes the gradient of the loss with respect to each child box.

• By calling `bprop` in the reverse order, we obtain backpropagation
Model Selection

- Training Protocol:
  - Train your model on the Training Set \( D^{\text{train}} \)
  - For model selection, use Validation Set \( D^{\text{valid}} \)
    - Hyper-parameter search: hidden layer size, learning rate, number of iterations/epochs, etc.
  - Estimate generalization performance using the Test Set \( D^{\text{test}} \)

- Generalization is the behavior of the model on unseen examples.
Early Stopping

- To select the number of epochs, stop training when validation set error increases (with some look ahead).
Mini-batch, Momentum

• Make updates based on a mini-batch of examples (instead of a single example):
  - the gradient is the average regularized loss for that mini-batch
  - can give a more accurate estimate of the gradient
  - can leverage matrix/matrix operations, which are more efficient

• **Momentum**: Can use an exponential average of previous gradients:

\[
\nabla_{\theta}^{(t)} = \nabla_{\theta} l(f(x^{(t)}), y^{(t)}) + \beta \nabla_{\theta}^{(t-1)}
\]

  - can get pass plateaus more quickly, by “gaining momentum”
Learning Distributed Representations

• Deep learning is research on learning models with multilayer representations
  - multilayer (feed-forward) neural networks
  - multilayer graphical model (deep belief network, deep Boltzmann machine)

• Each layer learns “distributed representation”
  - Units in a layer are not mutually exclusive
    - each unit is a separate feature of the input
    - two units can be “active” at the same time
  - Units do not correspond to a partitioning (clustering) of the inputs
    - in clustering, an input can only belong to a single cluster
Inspiration from Visual Cortex
Feedforward Neural Networks

- How neural networks predict $f(x)$ given an input $x$:
  - Forward propagation
  - Types of units
  - Capacity of neural networks

- How to train neural nets:
  - Loss function
  - Backpropagation with gradient descent

- More recent techniques:
  - Dropout
  - Batch normalization
  - Unsupervised Pre-training
Why Training is Hard

• First hypothesis: Hard optimization problem (underfitting)
  ➢ vanishing gradient problem
  ➢ saturated units block gradient propagation

• This is a well known problem in recurrent neural networks
Why Training is Hard

• First hypothesis (underfitting): better optimize
  - Use better optimization tools (e.g. batch-normalization, second order methods, such as KFAC)
  - Use GPUs, distributed computing.

• Second hypothesis (overfitting): use better regularization
  - Unsupervised pre-training
  - Stochastic drop-out training

• For many large-scale practical problems, you will need to use both: better optimization and better regularization!
Unsupervised Pre-training

• Initialize hidden layers using unsupervised learning
  - Force network to represent latent structure of input distribution
  - Encourage hidden layers to encode that structure

Why is one a character and the other is not?
Unsupervised Pre-training

- Initialize hidden layers using **unsupervised learning**
  - This is a harder task than supervised learning (classification)
  - Hence we expect less overfitting
Autoencoders: Preview

• Feed-forward neural network trained to reproduce its input at the output layer

\[ h(x) = g(a(x)) = \text{sigm}(b + Wx) \]

\[ W^* = W^\top \]

Decoder

\[ \hat{x} = o(\hat{a}(x)) = \text{sigm}(c + W^*h(x)) \]

For binary units

Encoder

\[ h(x) = g(a(x)) = \text{sigm}(b + Wx) \]
Autoencoders: Preview

• Loss function for **binary inputs**

\[ l(f(x)) = - \sum_k (x_k \log(\hat{x}_k) + (1 - x_k) \log(1 - \hat{x}_k)) \]

  ➢ Cross-entropy error function \( f(x) \equiv \hat{x} \)

• Loss function for **real-valued inputs**

\[ l(f(x)) = \frac{1}{2} \sum_k (\hat{x}_k - x_k)^2 \]

  ➢ sum of squared differences
  ➢ we use a linear activation function at the output
Pre-training

- We will use a greedy, layer-wise procedure
  - Train one layer at a time with unsupervised criterion
  - Fix the parameters of previous hidden layers
  - Previous layers can be viewed as feature extraction
Fine-tuning

• Once all layers are pre-trained
  - add output layer
  - train the whole network using supervised learning

• We call this last phase fine-tuning
  - all parameters are “tuned” for the supervised task at hand
  - representation is adjusted to be more discriminative
Why Training is Hard

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  ➢ Unsupervised pre-training
  ➢ Stochastic drop-out training

• For many large-scale practical problems, you will need to use both: better optimization and better regularization!
• **Key idea**: Cripple neural network by removing hidden units stochastically

  ➢ each hidden unit is set to 0 with probability 0.5

  ➢ hidden units cannot co-adapt to other units

  ➢ hidden units must be more generally useful

• Could use a different dropout probability, but 0.5 usually works well
Dropout

- Use random binary masks $m^{(k)}$
  - Layer pre-activation for $k > 0$
    \[ a^{(k)}(x) = b^{(k)} + W^{(k)}h^{(k-1)}(x) \]
  - Hidden layer activation (k=1 to L):
    \[ h^{(k)}(x) = g(a^{(k)}(x)) \odot m^{(k)} \]
  - Output activation (k=L+1)
    \[ h^{(L+1)}(x) = o(a^{(L+1)}(x)) = f(x) \]
Dropout at Test Time

• At test time, we replace the masks by their expectation
  ➢ This is simply the constant vector 0.5 if dropout probability is 0.5
  ➢ For single hidden layer: equivalent to taking the geometric average
    of all neural networks, with all possible binary masks

• Can be combined with unsupervised pre-training

• Beats regular backpropagation on many datasets

• Ensemble: Can be viewed as a geometric average of exponential
  number of networks.
Why Training is Hard

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• For many large-scale practical problems, you will need to use both: better optimization and better regularization!
Batch Normalization

• Normalizing the inputs will speed up training (Lecun et al. 1998)
  ➢ could normalization be useful at the level of the hidden layers?

• **Batch normalization** is an attempt to do that (Ioffe and Szegedy, 2014)
  ➢ each unit’s pre-activation is normalized (mean subtraction, stddev division)
  ➢ during training, mean and stddev is computed for each minibatch
  ➢ backpropagation takes into account the normalization
  ➢ at test time, the global mean / stddev is used
Batch Normalization

**Input:** Values of $x$ over a mini-batch: $\mathcal{B} = \{x_1...m\}$; Parameters to be learned: $\gamma, \beta$

**Output:** $\{y_i = \text{BN}_{\gamma, \beta}(x_i)\}$

\[
\begin{align*}
\mu_B & \leftarrow \frac{1}{m} \sum_{i=1}^{m} x_i \quad \text{// mini-batch mean} \\
\sigma^2_B & \leftarrow \frac{1}{m} \sum_{i=1}^{m} (x_i - \mu_B)^2 \quad \text{// mini-batch variance} \\
\hat{x}_i & \leftarrow \frac{x_i - \mu_B}{\sqrt{\sigma^2_B + \epsilon}} \quad \text{// normalize} \\
y_i & \leftarrow \gamma \hat{x}_i + \beta \equiv \text{BN}_{\gamma, \beta}(x_i) \quad \text{// scale and shift}
\end{align*}
\]

Learned linear transformation to adapt to non-linear activation function ($\gamma$ and $\beta$ are trained)
Batch Normalization

• Why normalize the pre-activation?
  - can help keep the pre-activation in a non-saturating regime (though the linear transform $y_i \leftarrow \gamma \hat{x}_i + \beta$ could cancel this effect)

• Use the global mean and stddev at test time.
  - removes the stochasticity of the mean and stddev
  - requires a final phase where, from the first to the last hidden layer
    • propagate all training data to that layer
    • compute and store the global mean and stddev of each unit
  - for early stopping, could use a running average