# **CS2109S**

AY22/23 Sem 2

github.com/jasonqiu212

#### 01. Introduction

- Agent Anything that can perceive its environment through sensors and acting upon that env. through actuators
- Agent Function Maps from percept histories to actions
- Rational Agent Chooses an action that is expected to maximize its performance measure, given by percept sequence and built-in knowledge
- Autonomous Agent If behavior is determined by its own expereince

#### Performance Measure of Function

Motivation: For an agent to do the right thing, need a measure of goodness

# **Defining the Problem: PEAS**

- 1. Performance measure
- 2. Environment
- 3. Actuators
- 4. Sensors

#### **Characterizing the Environment**

- 1. Fully observable (vs. Partially) Agent's sensors can access complete state of env. all the time
- Deterministic (vs. Stochastic) Next state of env. is determined by current state and action executed by agent
  - Strategic If env. is deterministic except for actions of other agents
- 3. **Episodic** (vs. Sequential) Agent's experience is divided into atomic episodes, where each episode includes perceiving and an action, and action depends on episode itself
- 4. Static (vs. Dynamic) Env. is unchanged while agent is deciding
  - Semi Time does not affect env., but affects performance score
- 5. Discrete (vs. Continuous) Discrete num. of percepts and actions
- 6. Single Agent (vs. Multi-agent) Agent operating by itself in an env.

# Implementing Agents (in ascending complexity)

- 1. Simple Reflex Agents Fixed conditional rules
- 2. Model-based Reflex Agents Stores percept history to make decisions about internal model of world with conditional rules. Eg. Roomba
- 3. Goal-based Agents Keep in mind a goal and action aims to achieve it
- 4. Utility-based Agents Find best way to achieve goal
- 5. Learning Agents Learn from previous experiences

# **Exploitation vs. Exploration**

- Exploitation Maximize expected utility using current knowledge of world
- Exploration Learn more about the world to improve future gains. May not always maximize performance measure.

# 02. Uninformed Search

- Deterministic, fully observable
- Tree Search Can revisit nodes
- Graph Search Tracks visited (Tree Search + Memoization)
- Uninformed Search Uses only information available in problem definition  $| \bullet |$  Space: O(bd)

#### Formulating the Problem

- 1. How to represent state in problem?
- 2. Initial state
- 3. Actions: Successor function
- 4. Goal test
- 5. Path cost
- Abstraction Function Maps abstracted representation to real world state
- Representation Invariant  $I(c) = \text{True} \rightarrow \exists a \text{ s.t. } AF(c) = a$

#### **Breadth-first Search**

- Idea: Expand shallowest unexpanded node using queue
- Given: b: Branching factor and d: Depth of optimal solution
- Complete: Yes (if tree is finite)
- Time:  $O(b^{d+1})$
- Space:  $O(b^d)$
- Optimal: Yes (if cost = 1)
- BFS is Uniform-cost Search with same cost

#### Uniform-cost Search

- Idea: Expand least-cost unexpanded node using priority queue (Dijkstra's)
- Given: C\*: Cost of optimal solution
- Complete: Yes (if step cost  $\geq \epsilon$  where  $\epsilon \geq 0$ )
- Time:  $O(b^{(C^*/\epsilon)})$  ( $C^*/\epsilon$  is approx. number of layers)
- Space:  $O(b^{(C^*/\epsilon)})$
- Optimal: Yes

# Depth-first Search

- Idea: Expand deepest unexpanded node using stack
- Given: m: Maximum depth of tree
- Complete: No (fails with infinite depth or loops)
- Time:  $O(b^m)$
- Space: O(bm) (better than BFS)
- Optimal: No

# **Depth-limited Search**

- Motivation: How to handle infinite depth for DFS?
- Idea: DFS with depth limit I where nodes at depth I have no children
- Time:  $b^0 + b^1 + ... + b^{(d-1)} + b^d = O(b^d)$

# **Iterative Deepening Search**

- Motivation: How to determine depth limit? We don't.
- Idea: Try different depths for depth-limited search
- BFS pretending to be DFS to save space
- Complete: Yes
- Time:  $(d+1)b^0 + db^1 + ... + b^d = O(b^d)$  (More overhead than DLS)

#### Summary

	BFS	Uniform Cost	DFS	DLS	IDS
Complete	Yes	Yes	No	No	Yes
Time	$O(b^d)$	$O(b^{C^*/\epsilon})$	$O(b^m)$	$O(b^l)$	$O(b^d)$
Space	$O(b^d)$	$O(b^{C^*/\epsilon})$	O(bm)	O(bl)	O(bd)
Optimal	Yes	Yes	No	No	No

#### **Bidirectional Search**

- Idea: Search both forwards from initial state and backwards from goal state. Stop when searches meet.
- Time:  $O(2b^{d/2})$
- Operators must be reversible
- Can have many goal states
- How to check if node intersects with other half?

#### 03. Informed Search

#### Heuristic

- Heuristic Estimated cost from n to goal
- Admissible h(n) is admissible if, for every node n,  $h(n) < h^*(n)$  where  $h^*$  is the true cost
- if h is admissible, then A\* using tree search is optimal
- Consistent h(n) is consistent if, for every node n and every successor n'of n generated by action  $a, h(n) \le c(n, a, n') + h(n')$
- Triangle inequality
- If h is consistent, f(n) is non-decreasing along any path (f(n') > f(n))
- If h is consistent, then h is admissible
- if h is admissible, then A\* using graph search is optimal

#### **Dominance**

- If  $h_2(n) \geq h_1(n)$  for all n, then  $h_2$  dominates  $h_1$
- If  $h_2$  dominates  $h_1$  and both are admissible, then  $h_2$  is better for search

#### How to invent admissible heuristic?

- Set fewer restrictions on actions
- E.g. Number of misplaced tiles, Total manhattan distance

#### Best-first Search

- Idea: Expand most desirable node using priority queue
- Evaluation Function: f(n) = h(n)
- Complete: No. Possible to be stuck in loop
- Time and space:  $O(b^m)$
- Optimal: No

#### A\* Search

- Idea: Take note of cost so far and heuristic
- Evaluation Function: f(n) = g(n) + h(n) where g(n) is cost to reach n
- Complete: Yes, unless non-increasing, since cost is factored in
- Time and space: Same as BFS
- Optimal: Yes, depending on the heuristic

# Iterative Deepening A\* Search (IDA\*)

- Motivation: How can we save space?
- ullet Idea: Have a cutoff for f and remember the best f that exceeds cutoff
- Similar to IDS. Linear space complexity.
- Optimal and complete

# Simplified Memory A\* Search (SMA\*)

- Motivation: How can we save space?
- Idea: Do normal A\*. If memory is full, drop node with worst f.
- Lose completeness

#### **Local Search**

- Motivation: What if the goal state is the solution? The path is irrelevant.
- Idea: Keep single current state and try improving it
- Formulating the problem:
  - 1. Initial state
  - 2. Actions: Successor function
  - 3. Good heuristic
  - 4. Goal test

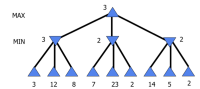
#### Hill-climbing

- Idea: Generate successors from current and pick the best using heuristic
- What if stuck into local minima?
- Introduce randomness
- Simulated Annealing Search Allow some bad moves and gradually decrease frequency
- Why only keep 1 best state?
- Beam Search Perform k hill-climbing searches in parallel
- How to generate successors?
- Genetic Algorithms Successsor is generated by combining 2 parent states

# 04. Adversarial Search

- Assumption: Opponents reacts rationally
- Formulating the problem:
- Initial state
- Successor function
- Terminal test
- Utility function: Measures how good the move is for a player

#### Minimax



- Idea: Choose move that yields highest minimax value using DFS
- Complete: Yes (if tree is finite)
- Time:  $O(b^m)$ • Space: O(bm)
- Optimal: Yes (against an optimal opponent)

#### How to Draw a Game Tree?

• Do not have repeated states per level

# **Alpha-Beta Pruning**

- Motivation: How to save time for Minimax?
- Idea: By tracking max. and min. values so far, can prune some paths that we would never choose
  - 1.  $\alpha$  contains max. and  $\beta$  contains min.
  - 2. Initially,  $\alpha = -\infty$  and  $\beta = \infty$
  - 3. When going down, copy  $\alpha$  and  $\beta$
  - 4. Prune if  $\alpha \geq \beta$
  - 5. When going up, depending on MIN/MAX level, update  $\alpha$  or  $\beta$
- With perfect ordering, time complexity:  $O(b^{m/2})$ . Doubles search depth.

#### **Resource Limits**

- $\bullet$  In reality, search space for games can be very large.  $\alpha\text{-}\beta$  pruning also not fast enough.
- Solution: Limit depth (Only see a finite moves ahead) and determine best move using evaluation function to estimate desirability of position (Heuristic)
- Other hacks:
- Transpoisitions Memoize equivalent states
- Pre-computation of opening/closing moves

# 05. Introduction to Machine Learning

A machine learns if it improves performance P on task T based on experience E. Where T must be fixed, P must be measurable, E must exist

#### Types of Feedback

- Supervised Correct answer given for each example
- Regression Predict results within continuous output
- Classification Predict results in discrete output
- **Unsupervised** No answers given
- Weakly supervised Answer given, but not precise
- Reinforcement Occasional rewards given

#### **Decision Trees**

- DT can express any function of input attributes, if data is consistent
- Goal: Make DT compact. How?

#### Information Theory

- Idea: Choose attribute that splits examples into subsets that are ideally 'all
  positive' or 'all negative'
- Entropy Measure of randomness in set of data

$$I(P(v_1), ..., P(v_n)) = -\sum_{i=1}^{n} P(v_i) \log_2 P(v_i)$$

 $\bullet$  For data with p positive examples and n negative examples:



$$I(\frac{p}{p+n}, \frac{n}{p+n}) =$$

$$-\frac{p}{p+n} \log_2 \frac{p}{p+n}$$

$$-\frac{n}{p+n} \log_2 \frac{n}{p+n}$$

- Information Gain (IG) Reduction in entropy from attribute test
- Goal: Choose attribute with largest information gain
- Intuition: IG = Entropy of this node Entropy of children nodes
- ullet Given chosen attribute A with v distinct values:

$$\begin{aligned} \text{remainder}(A) &= \sum_{i=1}^v \frac{p_i + n_i}{p+n} I(\frac{p_i}{p_i + n_i}, \frac{n_i}{p_i + n_i}) \\ IG(A) &= I(\frac{p}{p+n}, \frac{n}{p+n}) - \text{remainder}(A) \end{aligned}$$

- Decision Tree Learning Recursively choose attributes with highest IG
- IG is not the only way. Can use whatever objective function that achieves the criteria we want.

#### **Performance Measurement**

- Correctness Correct if  $\hat{y} = y$
- Accuracy  $-\frac{1}{m}\sum_{j=1}^{m}(\hat{y_j}=y_j)$
- Confusion Matrix:

		Actual Label		
		+ve	-ve	
Predicted Label	+ve	TP True Positive	FP False Positive	
	-ve	FN False Negative	TN True Negative	

- Accuracy =  $\frac{TP+TN}{TP+FN+FP+TN}$
- **Precision**  $\frac{TP}{TP+FP}$  How precise are positive predictions?
- Recall  $\frac{TP}{TP+FN}$  How many actual positives are predicted?
- F1 Score  $\frac{2}{1/P+1/R}$  Harmonic mean of precision and recall
- Type I Error: FP, Type II Error: FN
- FP Rate =  $\frac{FP}{FP+TN}$  TP Rate =  $\frac{TP}{TP+FN}$

#### Pruning

- Motivation: DT overfits to training set, but performs poorly on test set
- Occam's Razor: Simple hypothesis preferred
- Pruning Ignores outliers, which reduces overfitting
- Idea: Go with the majority of T/Fs
- E.g. Min-sample, Max-depth

# 06. Linear Regression

#### Notation

- m = Number of training examples
- $\bullet$  n =Number of features
- $x_i^{(i)} = \text{Input feature } j \text{ of } i \text{th training example}$
- $\bullet y = \text{Output variables}$

#### **Hypothesis**

$$h_w(x): w_0 + w_1 x$$

# **Cost Function (Square Error Function)**

$$J(w_0, w_1) = \frac{1}{2m} \sum_{i=1}^{m} (h_w(x^{(i)}) - y^{(i)})^2$$

- Goal: Minimize cost function. Thus, hypothesis is close to training samples
- Why squared error? Convenience, since we need to differentiate later

### **Gradient Descent**

- 1. Start at some  $(w_0, w_1)$ . Pick nearby point that reduces  $J(w_0, w_1)$ .
- 2. Repeat until convergence:

$$w_j := w_j - \alpha \frac{dJ(w_0, w_1, \dots)}{dw_j}$$

- All updates done at end
- ullet How to do  $rac{dJ(w_0,w_1)}{dw_j}$ ? Partial derivative: Hold everything else constant

• 
$$\frac{dJ(w_0, w_1)}{dw_j} = \frac{d}{dw_j} (\frac{1}{2m} \sum_{i=1}^m (w_0 + w_1 x^{(i)} - y^{(i)})^2)$$

- $\frac{dJ(w_0,w_1)}{dw_0} = \frac{1}{m} \sum_{i=1}^m (w_0 + w_1 x^{(i)} y^{(i)})$  (Note: Chain rule)
- $\frac{dJ(w_0, w_1)}{dw_1} = \frac{1}{m} \sum_{i=1}^{m} (w_0 + w_1 x^{(i)} y^{(i)}) x^{(i)}$
- Time complexity: O(kmn) where k is number of iterations

#### **Learning Rate**

- If  $\alpha$  too small, then descent is too slow. If  $\alpha$  too big, then might overshoot.
- ullet Given constant lpha, descent will grow smaller as we approach minimum

#### Variants of Gradient Descent

- Batch gradient descent: Consider all training examples when updating
- Stochastic gradient descent: Consider 1 random data point at a time (Cheaper and more randomness)
- Mini-batch gradient descent

#### **Using Matrices**

• Given: 
$$w = \begin{pmatrix} w_0 \\ \vdots \\ w_n \end{pmatrix}$$
 and  $x = \begin{pmatrix} x_0 \\ \vdots \\ x_n \end{pmatrix} = \begin{pmatrix} 1 \\ \vdots \\ x_n \end{pmatrix}$ 

 $\bullet$   $h_w(x): w^T x$ 

#### **Feature Scaling**

- Motivation: Gradient descent does not work well if features have different scales
- Mean Normalization  $x_i \leftarrow \frac{x_i \mu_i}{\sigma_i}$

# **Normal Equation**

$$w = (X^T X)^{-1} X^T Y$$

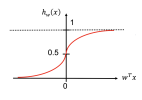
- ullet No need to choose lpha and feature scaling
- $X^TX$  needs to be invertible
- Time complexity:  $O(n^3)$ . Slow if n is big

# 07. Logistic Regression

- Motivation: Classification with continuous input values.
- Idea: Come up with a decision boundary to separate data points.  $h_w(x_1,x_2)=1$ , if  $w_0+w_1x_1+w_2x_2>0$ , or 0, otherwise

#### **Sigmoid Function**

• Motivation: Step function is discontinuous and not differentiable



$$h_w(x) = g(w^T x)$$

$$g(z) = \frac{1}{1 + e^{-z}}$$

# Hypothesis

$$h_w(x): \frac{1}{1 + e^{-w^T x}}$$

ullet Interpretation: Estimated probability that y=1 given input x

#### **Cost Function**

- Problem: Least square error gives non-convex cost function, which is bad for G.D.
- Solution: Use log



- If y = 1, let cost be  $-log(h_w(x))$
- $h_w(x) \to 0, J(w) \to \infty$
- $h_w(x) \to 1, J(w) \to 0$



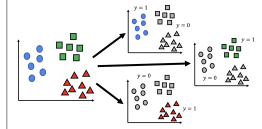
- ullet If y=0, let cost be  $-log(1-h_w(x))$
- $h_w(x) \to 0, J(w) \to 0$
- $h_w(x) \to 1, J(w) \to \infty$

$$J(w) = -\frac{1}{m} \sum_{i=1}^{m} y^{(i)} log h_w(x^{(i)}) + (1 - y^{(i)}) log (1 - h_w(x^{(i)}))$$

#### **Gradient Descent**

- Same algorithm as linear regression
- ullet  $\frac{dJ(w)}{dw_i}=rac{1}{m}\sum_{i=1}^m(h_w(x^{(i)})-y^{(i)})x_n^{(i)}$  (Same as linear regression)

#### Multi-class Classification



- 1. Train binary classifier  $h_w^{(i)}(x)$  for each class i to predict y=i
- 2. For each input x, pick class i is greatest (i.e.  $\max_i h_w^{(i)}(x)$ )

# 08. Model Evaluation and Selection

#### **Linear Regression**

$$J_{\mathsf{test}}(w) = \frac{1}{2m_{\mathsf{test}}} \sum_{i=1}^{m_{\mathsf{test}}} (h_w(x_{\mathsf{test}}^{(i)}) - y_{\mathsf{test}}^{(i)})^2$$

#### Logistic Regression

$$\operatorname{error}(h_w(x), y) = \begin{cases} 1 & \text{if misclassification} \\ 0 & \text{otherwise} \end{cases}$$

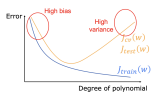
$$\text{Test Error} = \frac{1}{m_{\text{test}}} \sum_{i=1}^{m_{\text{test}}} \operatorname{error}(h_w(x_{\text{test}}^{(i)}), y_{\text{test}}^{(i)})$$

#### Model Selection

- How to choose model? (i.e.  $x^2$ ,  $x^3$ , etc. for hypothesis)
  - 1. Split data into 3 sets: Training set, validation set, and test set
  - 2. Train each model using training set
  - 3. Compute  $J_{cv}(w)$  for each model and pick model with lowest  $J_{cv}(w)$
  - 4. Use  $J_{\text{test}}(w)$  to estimate performance on unseen samples

#### Bias and Variance

- High bias Underfit, High variance Overfit
- As degree of polynomial increases:



Impact of m: High Bias

# Impact of m: High Variance



ullet Observation: If high bias,  $J_{cv}$  and  $J_{train}$  are equally bad (Small gap). If high variance, more samples does not help close the gap.

- How do we address overfitting?
- Reduce number of features
- Regularization Keep features, but reduce respective weights

# 09. Regularization

- Idea: Include w inside J(w) to minimize weights and get simpler hypothesis
- What if  $\lambda$  is too large? Weights tend to 0, resulting in straight line
- How to choose  $\lambda$ ? Use validation set
- Small  $\lambda$ :  $J_{\text{train}}(w)$  low and  $J_{cv}(w)$  high (Overfit)
- Large  $\lambda$ :  $J_{\text{train}}(w)$  high and  $J_{cv}(w)$  high (Underfit)

#### Linear Regression with Regularization

$$J(w) = \frac{1}{2m} \left( \sum_{i=1}^{m} (h_w(x^{(i)}) - y^{(i)})^2 + \lambda \sum_{i=1}^{n} w_i^2 \right)$$

$$w_n := w_n - \frac{\alpha}{m} \sum_{i=1}^m ((h_w(x^{(i)}) - y^{(i)}) x_n^{(i)}) - \frac{\alpha \lambda}{m} w_n$$

#### Logistic Regression with Regularization

$$J(w) = -\frac{1}{m} \sum_{i=1}^{m} y^{(i)} log(h_w(x^{(i)})) + (1 - y^{(i)}) log(1 - h_w(x^{(i)})) + \frac{\lambda}{2m} \sum_{i=1}^{n} w_i^2$$

$$w_n := w_n - \frac{\alpha}{m} \sum_{i=1}^m ((h_w(x^{(i)}) - y^{(i)}) x_n^{(i)}) - \frac{\alpha \lambda}{m} w_n$$

$$w = (X^T X + \lambda I)^{-1} X^T Y$$
 where I's first column is 0

 $\bullet X^T X$  can be non-invertible

# 10. Support Vector Machine

• Idea: Maximize margin between positive and negative samples

#### **Decision Rule**

$$w \cdot x \geq c \rightarrow \mathsf{Positive}; w \cdot x < c \rightarrow \mathsf{Negative}$$

- Dot product:  $u \cdot v = u^T v = p||v||$  (By law of cosine)
- Intuition: Decision boundary  $\bot$  Weight vector.  $w \cdot x = p||w||$ . Sample a is on decision boundary if  $w \cdot a = c$
- Constraints: Let b = -c
- $w \cdot x + b > 1$  if y = 1
- $w \cdot x + b < -1$  if y = 0
- Combined: Let  $\hat{y}^{(i)} = 1$  or -1 for pos. and neg. samples respectively.  $\hat{y}^{(i)}(w \cdot x^{(i)} + b) > 1$
- If x is inside margin, then  $w \cdot x + b = 1$  or -1 respectively
- Why add these constraints? Mathematical convenience for hinge loss

#### Margin

$$\mathsf{Margin}\;\mathsf{width} = \frac{2}{||w||}$$

- Let  $x^+$  and  $x^-$  be closest positive and negative samples
- Margin width  $=(x^+-x^-)\cdot \frac{w}{||w||}$  (i.e. Length of projection of  $x^+-x^-$  on weight vector)  $=\frac{1-b+1+b}{||w||}$  (since  $x^+$  and  $x^-$  are inside margin)  $=\frac{2}{||w||}$

# Objective Function for Hard-Margin

$$\min \frac{1}{2} ||w||^2 \text{ s.t. } \hat{y}^{(i)}(w \cdot x + b \ge 1)$$

- 1. Maximize margin:  $\max \frac{2}{||w||} = \min ||w|| = \min \frac{1}{2} ||w||^2$
- 2. Classify correctly
- Hard-Margin No samples inside margin
- Before, we assume hard margin. What if there exists outliers that causes SVM to overfit?

#### **Objective Function for Soft-Margin**

$$h_w(x) = \begin{cases} 1 & \text{if } w^T x \ge 0\\ 0 & \text{otherwise} \end{cases}$$

- Slack Variable  $(\xi)$  Loss of misclassified point
- ullet Goal: Maximize margin and allow misclassification by tweaking C
- Large C overfits (Basically hard-margin). Small C underfits.
- i.e.  $\min(\frac{1}{2}||w||^2 + C\sum_i \xi^{(i)})$  s.t.  $\forall i, \xi^{(i)} \geq 0$  and  $\hat{y}^{(i)}(w \cdot x^+b) \geq 1 \xi^{(i)} \rightarrow J(w) = C\sum_i \max(0, 1 \hat{y}^{(i)}(w^Tx^{(i)})) + \frac{1}{2}\sum_{i=1}^n w_i^2$
- Hard-margin: Must follow constraint. Soft-margin: Constraint is flexible with slack variable, so can define cost function to minimize.

$$J(w) = C \sum_{i}^{m} y^{(i)} \mathsf{cost}_1(w^T x^{(i)}) + (1 - y^{(i)}) \mathsf{cost}_0(w^T x^{(i)}) + \frac{1}{2} \sum_{i=1}^{n} w_i^2$$

# Legistic Loring

#### Hinge Loss

- $\bullet \, \mathsf{cost}_1(z) = \max(0, 1-z)$
- $\bullet \cot_0(z) = \max(0, 1+z)$

#### Kernel

- Motivation: What if data is not linearly separable?
- Idea: Map features to higher dimensions (i.e.  $\phi(x)$ ). High dim. is slow.
- Kernel Function Dot product between two points in transformed dim.

$$K(u, v) = \phi(u) \cdot \phi(v)$$

- Property: Getting K(u,v) does not need  $\phi(u)$  and  $\phi(v)$
- Linear Kernel  $K(u, v) = u \cdot v$
- Polynomial Kernel  $K(u,v) = (u \cdot v)^d$
- Gaussian Kernel  $K(u,v) = e^{-||u-v||^2/2\sigma^2}$  (Note:  $\phi(u)$  can map to infinite dimensions)

#### **Kernel Trick**

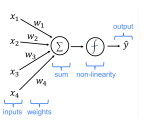
- ullet By manipulating SVM objective func. and decision rule,  $x^{(i)} \cdot x^{(j)}$  emerges
- $\bullet$  If we map features (i.e.  $\phi(x)$ ), can replace above term with kernel (K(u,v))
- Trick: No need to compute transformed features explicitly

#### **Similarity Features**

- Idea: Compute new features based on proximity to landmarks
- ullet Given example x.  $f_i = sim(x, l^{(i)}) = K(x, l^{(i)})$  where K is gaussian kernel
- Transformed x:  $f = [f_0, f_1, \dots, f_m]^T$ . Plug into obj. function.
- $\bullet$  Do feature scaling, else similarity value may be dominated by some features
- $\bullet$  Large  $\sigma$  underfits (Smoother features). Small  $\sigma$  overfits.
- Not all similarity functions get valid kernels. Need to satisfy Mercer's Thm.

#### 11. Neural Networks

• Idea: Models after human brain (Perceptron)



$$\hat{y} = g(\sum_{i=0}^n w_i x_i)$$
 where  $x_0 = 1$ 

- Activation Function (g) Nonlinear function (e.g. Step func. sgn(x),  $\sigma(x)$ , tanh(x), max(0,x))
- Supports both classification and regression. Depends on activation func.
- If activation function is linear, then regression

# Perceptron Learning Algorithm (PLA)

- 1. Initialize weights w
- 2. Select misclassified sample and update w. Repeat until convergence.

$$w \leftarrow w + \eta(y - \hat{y})x$$

- Why this works? Each update gets closer to correct classification
- Different from GD!
- Ends when no more misclassification. Thus, can select any linear model
- Cannot converge on non-linear data

# Single-Layer Neural Network

- Basically logistic regression, if activation function is  $\sigma(x)$
- How to determine weights? Gradient descent
- Many ways to calculate error. e.g. MSE  $(J(w) = \frac{1}{2}(\hat{y} y)^2)$ .
- Given  $\hat{y} = g(f(x)); f(x) = w^T x$ ,  $\frac{dJ(w)}{dw_i} = \frac{dJ}{d\hat{y}} \frac{d\hat{y}}{df} \frac{df}{dw_i} = (\hat{y} y)g'(f)x_i$

$$w_i := w_i - \eta(\hat{y} - y)g'(f)x_i$$

• If activation function  $g = \sigma(x)$ , then  $g'(x) = \sigma'(x) = \sigma(x)(1 - \sigma(x))$ 

$$w_i := w_i - \eta(\hat{y} - y)\hat{y}(1 - \hat{y})x_i$$

# Forward Propagation

- How to fit non-linear data? Have many layers. 1 layer computes more complex features as next layer's input
- Given layer *l*:
- Vector: Bolded, lower case. Matrix: Bolded, upper case

$$\mathbf{a}^{[l]} = \mathbf{g}^{[l]}(\mathbf{f}^{[l]})$$

$$\mathbf{f}^{[l]} = (\mathbf{W}^{[l]})^T \mathbf{a}^{[l-1]}$$

#### **Backpropagation**

- Big idea: Gradient descent with lots of chain rules
- Goal: How to get  $\frac{d\epsilon}{dw}$ ?
- ullet E.g.  $rac{d\epsilon}{dw_2}=rac{d\epsilon}{d\hat{y}}rac{d\hat{y}}{dw_2}$  and  $rac{d\epsilon}{dw_1}=rac{d\epsilon}{d\hat{y}}rac{d\hat{y}}{da_1}rac{da_1}{dw_1}$
- But, very costly to do this for many layers. Can we vectorize it?

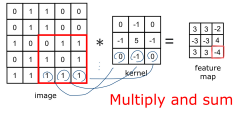
$$\hat{y}'(\mathbf{W}^{[l]}) = \frac{d\mathbf{f}^{[l]}}{d\mathbf{W}^{[l]}} (\delta^{[l]})^T = \mathbf{a}^{[l-1]} (\delta^{[l]})^T$$

$$\delta^{[l]} = \frac{dg^{[l]}}{d\mathbf{f}^{[l]}} \frac{d\mathbf{f}^{[l+1]}}{dg^{[l]}} \delta^{[l+1]} = g'^{[l]} (\mathbf{f}^{[l]}) \mathbf{W}^{[l+1]} \delta^{[l+1]}$$

# 12. Deep Learning

#### **Convolutional Neural Network**

- Idea: Exploit **spatial structure**. Map pixels using kernel to capture pixels as a group.
- Kernel: Like a filter



- Padding Add blank pixels at the side to avoid losing edge pixels
- Stride Skip some pixels by some number of steps for faster computation
- Let  $\mathbf{W}^{[l]}$  be kernels, which is a 3D matrix since we apply lots of kernels

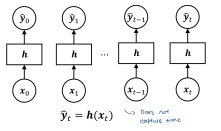
$$\mathbf{A}^{[l]} = q^{[l]}(\mathbf{W}^{[l]} * \mathbf{A}^{[l-1]})$$

- Each layer has many kernels (3D matrix)
- Kernel input: 3D matrix of features maps (Each depth is from diff. kernel)
- Kernel output: 2D matrix feature map
- Pooling Reduces dimensionality of feature maps to avoid number of layers exploding (E.g. Max-Pool, Average-Pool, Sum-Pool)
- Softmax Gets distribution of probabilities to make final classification by taking max. (Similar to multi-class classification)

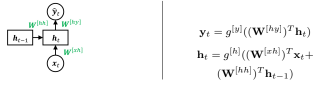
$$P(y = j) = \frac{e^{x_j}}{\sum_{i=0}^{k} e^{x_i}}$$

#### **Recurrent Neural Network**

- Idea: Exploit temporal structure by tracking past values
- Not limited to time. As long as there exists a recurring relationship.
- Try 1: Have many perceptrons running in parallel for each moment



• Try 2: Use past perceptron in new one



#### **Problems with Deep Learning**

- 1. Overfitting
- Drop out Randomly set some activations to 0
- Early stopping Stop training when  $J_{test}$  and  $J_{train}$  diverge
- 2. Vanishing/Exploding Gradient
- ullet Vanishing Gradient If many gradients pprox 0 in backpropagation, then weights won't change
- Exploding Gradient Gradients keep getting larger, causing GD to diverge
- Solutions:
- Proper weight initialization
- Use other activation functions (E.g. ReLU)
- Batch normalization (Feature scaling)
- Gradient clipping