

Linear Algebra:

- Vector Multiplication: $x^T y = \sum_{i=1}^n x_i y_i$
- Matrix Multiplication: $C = AB \rightarrow C_{ij} = \sum_{k=1}^n A_{ik} B_{kj}$
- Gradients: $\nabla f = \left[\frac{\partial f}{\partial x_1}, \frac{\partial f}{\partial x_2}, \dots, \frac{\partial f}{\partial x_n} \right]^T$
- Gradients of Common Functions: $\nabla(a^T x) = a$, $\nabla(x^T A x) = (A + A^T)x$
- **Machine Learning**
- Supervised Learning: Given labelled data, attempt to learn function that assigns labels to unlabelled data
- Unsupervised Learning: Given unlabelled data, attempt to find structure
- K-nearest Neighbours: Classify based on majority label of k nearest points
- Confusion matrix:

	Predicted: yes	Predicted: no	Actual: yes	Actual: no
TP				
FP				
FN				
TN				
- Precision = $\frac{TP}{TP+FP}$ Recall = $\frac{TP}{TP+FN}$ Specificity = $\frac{TN}{TN+FP}$
- Cross Validation: Split data into k subsets, train on k-1, test on 1, repeat k times
- Overfitting: Model fits training data too closely, fails to generalise
- Underfitting: Model too simple to capture underlying structure
- Decision Trees: Create best-split feature node, for all split data repeat
- Binary Classification: $f(x) \rightarrow y \in \{1, -1\}$
- Regression: $f(x) \rightarrow y \in \mathbb{R}$
- Hypothesis class: Function our algo will produce
- Loss function: How accurate is our predictor?
- Optimization Algo: How do we minimize loss?
- Linear Regression:
 - Hyp: $f(x) = w_1 + w_2 x$ or $f_w(x) = w \cdot \phi(x) : w = [w_1, w_2], \phi(x) = [1, x]$
 - Loss: $Loss(x, y, w) = (f_w(x) - y)^2$
 - $TrainLoss(w) = \frac{1}{|D_{train}|} \sum_{(x,y) \in D_{train}} Loss(x, y, w)$
 - Optimization: $\nabla_w TrainLoss(w) = \frac{1}{|D_{train}|} \sum_{(x,y) \in D_{train}} 2(w \cdot \phi(x) - y) \phi(x)$
 - Gradient Update: $w \leftarrow w - \eta \nabla_w TrainLoss(w) : \eta = 0.1$
- Binary Classification: Create a linear function representing a boundary
 - Hyp: $f(x) = sign(w \cdot \phi(x)) : sign(z) = 1 : z > 0, -1 : z < 0, 0 : z = 0$
 - Zero-One Loss: $Loss_{0-1}(x, y, w) = 1[f_w(x) \neq y]$
 - Hinge Loss: $Loss_{hinge}(x, y, w) = \max\{1 - (w \cdot \phi(x))y, 0\}$
 - Logistic Regression: $Loss_{logistic}(x, y, w) = \log(1 + e^{-(w \cdot \phi(x))y})$
- Stochastic Gradient Descent: Gradient descent is slow!
- Neural Networks
 - Hyp: $f(x) = V\sigma(W\phi(x))$ where σ is activation function
 - Activation functions: $\sigma(z) = \max\{0, z\}$ (ReLU), $\sigma(z) = \frac{1}{1+e^{-z}}$ (sigmoid)
 - Forward propagation: Compute layer outputs from input to output
 - Backpropagation: Compute gradients using chain rule, update weights
 - Architecture: Input layer \rightarrow Hidden layers \rightarrow Output layer
 - Layers represent multiple layers of abstraction
- How to prevent overfitting?
 - Reduce Dimensionality by removing features
 - Regularize: $\min_w TrainLoss(w) + \lambda/2 \cdot ||w||^2$

Search: No uncertainty

- Definitions: b : branching factor, m : maximum depth
- Components of a search problem:
 - State space: All possible states of an environment
 - Successor function: function that maps actions to consequences
 - Start state and end goal state
- DFS:
 - Idea: Expand one node as much as possible before exploring another path
 - $O(b^m)$ time complexity
 - $O(bm)$ space complexity
 - Complete if infinite tree size prevented and not optimal
- BFS:
 - Idea: prioritize nodes closest to start
 - $O(b^d)$ time complexity where d is depth of solution
 - $O(b^d)$ space complexity
 - Complete and optimal if step costs are equal
- Dijkstra's (Uniform Cost Search):
 - Idea: prioritize expansion of low-cost paths
 - $O(b^{1+\lceil C^*/\epsilon \rceil})$ time complexity where C^* is optimal cost, ϵ is minimum step cost
 - $O(b^{1+\lceil C^*/\epsilon \rceil})$ space complexity
 - Complete if step costs $\geq \epsilon > 0$ and optimal
- Greedy Search
 - Idea: Expand a node that we think is closest to the goal state
 - Utilize a heuristic
- A* Informed Search Algorithm
 - Idea: use a heuristic to improve efficiency and make informed decisions
 - Heuristic: A function that estimates how close a state is to the goal
 - A heuristic is admissible if it never overestimates the cost of reaching a goal: $0 \leq h(n) \leq h^*(n)$ where $h(n)$ is the cost to the nearest goal
 - A heuristic is consistent if $h(A) - h(C) \leq cost(A \rightarrow C)$. If it is consistent then it is optimal
 - $f(n) = g(n) + h(n)$

Markov Decision Processes

- MDPs are defined by:
 - set of states $s \in S$ and actions $a \in A$
 - transition functions $T(s, a, s')$ and reward functions $R(s, a, s')$
 - Start state and terminal state
 - Utility: sum of rewards received
- Expected utility: $EU(a) = \sum_{s'} P(RES(a) = s') U(s')$
- Policies: the plan to get from start to goal: $\pi^* : S \rightarrow A$
- With infinite utility we can mitigate with a finite horizon (limiting depth) or utilizing discounting: $U([r_0, \dots, r_\infty]) = \sum_{t=0}^{\infty} \gamma^t r_t \leq R_{max}/(1-\gamma)$
- Discounting: Sooner rewards have higher utility than later rewards
- The value of a state s is: $V^*(s) = \max_a Q^*(s, a) : Q^*(s, a) =$ expected utility starting out having taken action a from state s and acting optimally
- $\pi^*(s) = \operatorname{argmax}_a Q^*(s, a)$
- Bellman Equations: Take correct first action \rightarrow continue acting optimally
 - $V^*(s) = \max_a Q^*(s, a)$
 - $Q^*(s, a) = \sum_{s'} T(s, a, s')[R(s, a, s') + \gamma V^*(s')]$
- Complete Bellman Equation: $V^*(s) = \max_a \sum_{s'} T(s, a, s')[R(s, a, s') + \gamma V^*(s')]$
- Policy Extraction via Q-Values: $\pi^*(s) = \operatorname{argmax}_a Q^*(s, a)$
- Q-Value Iteration: $Q_{k+1}(s, a) = \sum_{s'} T(s, a, s')[R(s, a, s') + \gamma \max_{a'} Q_k(s', a')]$
- Alternatively, we can perform policy iteration where we calculate values for some fixed policy until convergence, then update the policy using resulting converged values as future values

Reinforcement Learning MDP's without T or R!

- MDPs, but we have to learn T and R
- Agent tries action A, gets $\{s, r\}$
- Model-Based Reinforcement Learning: Learn an approximate model based on experiences
 - STEP 1: Learn empirical MDP Model
 - Count $s' : \forall s, a$
 - Normalize to estimate $\hat{T}(s, a, s')$
 - Discover $\hat{R}(s, a, s')$ when we experience s, a, s'
 - STEP 2: Solve MDP, then run learned policy
- Model-Free Reinforcement Learning: Learn policies/values directly from experience
 - Passive RL: Execute a fixed policy $\pi(s)$ and learn state values
 - Direct Evaluation: Average sample values for each state
 - Temporal Difference Learning: Update $V^\pi(s)$ after each transition
 - TD Update: $V^\pi(s) \leftarrow (1-\alpha)V^\pi(s) + \alpha[R(s, \pi(s), s') + \gamma V^\pi(s')]$
 - Active RL: Learn optimal policy while acting
 - Q-Learning: Learn Q-values without knowing T or R
 - Q-Learning Update: $Q(s, a) \leftarrow (1-\alpha)Q(s, a) + \alpha[R(s, a, s') + \gamma \max_{a'} Q(s', a')]$
 - Exploration vs Exploitation: Balance trying new actions vs using known good actions
 - ϵ -greedy: With probability ϵ choose random action, else choose $\operatorname{argmax}_a Q(s, a)$
 - We can approximate Q-learning with functions f_1, f_2, \dots
 - $V(s) = w_1 f_1(s) + w_2 f_2(s) + \dots + w_n f_n(s)$
 - $Q(s, a) = w_1 f_1(s, a) + w_2 f_2(s, a) + \dots + w_n f_n(s, a)$

Games

- Types of games: Zero-sum: Competition (agents have opposite utilities), General-sum games: Agents have independent utilities
- We have states $s \in S$, players $p \in P$, actions $a \in A$, functions $Sx A \rightarrow S$, terminal test $S \rightarrow \{true, false\}$, terminal utilities: $Sx P \rightarrow R$
- Minimax: Time: $O(b^m)$, Space: $O(bm)$
- Alpha-Beta Pruning: Prune branches in a minimax tree to increase efficiency. Time: $O(b^{m/2})$.
- Multi-agent minimax: Terminals have utility tuples, node values are also utility tuples, each player maximizes its own component.
- Monte-Carlo Tree Search
 - Evaluation by rollouts: play multiple games to termination from a state s and count wins and losses
 - Selective search: explore parts of the tree that will help improve the decision at the root, regardless of depth
- Rollout: for each rollout, repeat until terminal: play a move according to a fixed, fast rollout policy. The fraction of wins correlates with the true value of the position
- MCTS V1: Allocate rollouts to promising and uncertain nodes.
- Promising and uncertain are defined by: $UCB1(n) = \frac{U(n)}{N(n)} + C \times \sqrt{\frac{\log N(PARENT(n))}{N(n)}}$
 - $N(n)$ number of rollouts from node n
 - $U(n)$ total utility of rollouts for Player(Parent(n))
- Theorem: As $N \rightarrow \infty$ UCT selects the minimax move

Bayesian Networks

- $P(a|b) = \frac{P(a,b)}{P(b)}$
- $P(Y|e) = \frac{P(Y,e)}{P(e)} = \alpha P(Y, e) = \alpha \sum_{h \in H} P(Y, e, h) \rightarrow \alpha = \sum_{y \in Y} P(Y, e)$
- Normalization: multiple each entry by $\alpha = 1/(\text{Sum of all entries})$
- Chain Rule: $P(x_1, x_2, \dots, x_n) = \prod_i P(x_i | x_1, \dots, x_{i-1})$
- Bayesian Statistics: Probabilities represent my uncertainty about the world
- Frequentist Statistics: Probabilities represent frequencies of real random outcomes. Properties of the world are fixed
- Probabilistic Inference: compute a desired probability from a probability model:
 - Enumerate options with sum
 - Sum out irrelevant variables
 - Normalize
- Time, Space, Data points: $O(d^n)$
- Bayes Rule: $p(H = h | Y = y) = \frac{P(H=h)P(Y=y|H=h)}{P(Y=y)}$, $P(a|b) = \frac{P(b|a)P(a)}{P(b)}$
- $P(Y = y) = \sum_{h' \in H} p(H = h') P(Y = y | H = h')$
- Two variables X and Y are independent if $\forall x, y : P(x, y) = P(x)P(y)$
- This implies that $P(x|y) = P(x), P(y|x) = P(y)$
- Conditional Independence: X is conditionally independent of Y given Z iff:
 - $\forall x, y, z : P(x|y, z) = P(x|z)$ or $\forall x, y, z : P(x, y|z) = P(x|z)P(y|z)$
- Bayes' Nets: Complex joint distributions using conditional distributions
 - Nodes: Variables(with domains). Can be assigned (observed) or unassigned (unobserved)
 - Arcs: Interactions that indicate direct influence between variables
 - For a Bayesian net, we have a set of nodes: X_i for each variable, a directed acyclic graph, and a conditional distribution for each node given its parent variables in the graph
 - A conditional probability table is a table where each row is a distribution for the child given values of its parents
 - $P(X_1, \dots, X_n) = \prod_i P(X_i | Parents(X_i))$

Markov Models

- Markov Models have an initial distribution $P(x_0)$, transition model $P(X_t | X_{t-1})$
- Hidden Markov Models have an additional sensor model with observations $P(E_t | X_t)$
 - Filtering $P(X_t | e_{1:t})$
 - Prediction $P(X_{t+k} | e_{1:t})$
 - Smoothing: $P(X_k | e_{1:t}) : k < t$
 - Explanation: $P(X_{1:t} | e_{1:t})$
- Markov Assumption: $P(X_t | X_0, \dots, X_{t-1}) = P(X_t | X_{t-1})$
- Stationary Assumption: Transition probabilities don't change over time
- Joint Distribution: $P(X_0, X_1, \dots, X_T) = P(X_0) \prod_{t=1}^T P(X_t | X_{t-1})$
- Filtering: $P(X_t | e_{1:t}) = \alpha P(e_t | X_t) \sum_{x_{t-1}} P(X_t | x_{t-1}) P(x_{t-1} | e_{1:t-1})$
- Prediction: $P(X_{t+1} | e_{1:t}) = \sum_{x_t} P(X_{t+1} | x_t) P(x_t | e_{1:t})$
- Most Likely Explanation: $\operatorname{argmax}_{x_{1:t}} P(x_{1:t} | e_{1:t})$ using Viterbi algorithm
- Viterbi: $m_{t+1}[x_{t+1}] = P(e_{t+1} | x_{t+1}) \max_{x_t} P(x_{t+1} | x_t) m_t[x_t]$
- Time steps: Elapse time (predict forward), observe (weight particles), resample

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Algorithms:
• K-nearest Neighbours
function KNN(x, data, k):
    distances = []
    for each (x_i, y_i) in data:
        d = distance(x, x_i)
        distances.append((d, y_i))
    distances.sort()
    neighbors = distances[0:k]
    return mode([y for (d, y) in neighbors])

• Learning a Decision Tree
function LearnDecisionTree(data, attributes):
    if all examples have same label:
        return Leaf(label)
    if attributes is empty:
        return Leaf(majority_label(data))
    best_attr = choose_best_attribute(data, attributes)
    tree = Node(best_attr)
    for each value v of best_attr:
        subset = {x in data : x[best_attr] = v}
        if subset is empty:
            tree.add_branch(v, Leaf(majority_label(data)))
        else:
            remaining = attributes - {best_attr}
            subtree = LearnDecisionTree(subset, remaining)
            tree.add_branch(v, subtree)
    return tree

function choose_best_attribute(data, attributes):
    return argmax over attributes of information_gain

• K-Means
function KMeans(data, k):
    centroids = randomly_select_k_points(data)
    repeat until convergence:
        clusters = [[] for i in range(k)]
        for each point x in data:
            closest = argmin_i distance(x, centroids[i])
            clusters[closest].append(x)
        for i in range(k):
            centroids[i] = mean(clusters[i])
    return centroids, clusters

• Stochastic Gradient Descent
function SGD(data, w_init, eta, num_epochs):
    w = w_init # [0,...,0]
    for epoch in range(num_epochs):
        shuffle(data)
        for each (x, y) in data:
            gradient = compute_gradient(Loss(x, y, w), w)
            w = w - eta * gradient
    return w

• DFS
function DFS(problem):
    frontier = Stack()
    frontier.push(start_state)
    explored = set()
    while frontier is not empty:
        node = frontier.pop()
        if node is goal:
            return solution
        explored.add(node)
        for each successor of node:
            if successor not in explored and not in frontier:
                frontier.push(successor)
    return failure

• BFS
function BFS(problem):
    frontier = Queue()
    frontier.enqueue(start_state)
    explored = set()
    while frontier is not empty:
        node = frontier.dequeue()
        if node is goal:
            return solution
        explored.add(node)
        for each successor of node:
            if successor not in explored and not in frontier:
                frontier.enqueue(successor)
    return failure

• Dijkstra's Uniform Cost Search Algorithm
function Dijkstra(problem):
    frontier = PriorityQueue()
    frontier.push(start_state, 0)
    explored = set()
    cost = {start_state: 0}
    while frontier is not empty:
        node = frontier.pop()
        if node is goal:
            return solution
        explored.add(node)
        for each successor of node:
            new_cost = cost[node] + step_cost(node, successor)
            if successor not in explored:
                if successor not in cost or new_cost < cost[successor]:
                    cost[successor] = new_cost
                    frontier.push(successor, new_cost)
    return failure

• A* Algorithm
function AStar(problem):
    frontier = PriorityQueue()
    frontier.push(start_state, h(start_state))
    explored = set()
    g_cost = {start_state: 0}
    while frontier is not empty:
        node = frontier.pop()
        if node is goal:
            return solution
        explored.add(node)
        for each successor of node:
            new_cost = g_cost[node] + step_cost(node, successor)
            if successor not in explored:
                if successor not in g_cost or new_cost < g_cost[successor]:
                    g_cost[successor] = new_cost
                    f_cost = new_cost + h(successor)
                    frontier.push(successor, f_cost)
    return failure

• Expectimax: For games with chance nodes, compute expected values instead of min/max
function EM(state):
    if state is terminal:
        return utility(state)
    if state is max node:
        return max over actions of EM(successor(state, action))
    if state is chance node:
        return sum outcomes of P(outcome) * EM(successor(state, outcome))

• Minimax
function Minimax(state):
    if state is terminal:
        return utility(state)
    if state is max node:
        return max over actions of Minimax(successor(state, action))
    if state is min node:
        return min over actions of Minimax(successor(state, action))

• Alpha Beta Pruning
function ABP(state, alpha, beta):
    if state is terminal:
        return utility(state)
    if state is max node:
        v = -infinity
        for each action:
            v = max(v, ABP(successor(state, action), alpha, beta))
            if v >= beta:
                return v # Beta cutoff
            alpha = max(alpha, v)
        return v
    if state is min node:
        v = +infinity
        for each action:
            v = min(v, ABP(successor(state, action), alpha, beta))
            if v <= alpha:
                return v # Alpha cutoff
            beta = min(beta, v)
        return v

• MCTS Version 2.0 UCT
function MCTS(root_state):
    tree = {root_state}
    N = {root_state: 0}
    U = {root_state: 0}
    repeat until out of time:
        # Selection: traverse tree using UCB1
        node = root_state
        path = [node]
        while node is fully expanded and not terminal:
            node = argmax_child UCB1(child)
            path.append(node)
        # Expansion: add new child
        if node is not terminal:
            child = unexplored_child(node)
            tree.add(child)
            N[child] = 0
            U[child] = 0
            path.append(child)
        # Simulation: rollout from new node
        result = rollout(child)
        # Backpropagation: update counts
        for n in path:
            N[n] += 1
            U[n] += result
        # Return action with highest visit count
        return argmax_child N[child]

• Value Iteration
function ValueIteration(S, A, T, R, gamma):
    Initialize V(s) = 0 for all s
    repeat until convergence:
        V_new = {}
        for each state s in S:
            V_new[s] = max_a sum_s' T(s,a,s') [R(s,a,s') + gamma*V(s')]
        V = V_new
    return V, extract_policy(V)

• Policy Iteration
function PolicyIteration(S, A, T, R, gamma):
    Initialize random policy pi
    repeat until policy stable:
        # Policy Evaluation
        V = solve V(s) = sum_s' T(s,pi(s),s') [R(s,pi(s),s') + gamma*V(s')]
        # Policy Improvement
        pi_new(s) = argmax_a sum_s' T(s,a,s') [R(s,a,s') + gamma*V(s')]
        if pi_new == pi: break
        pi = pi_new
    return pi

• Forward Algorithm for Hidden Markov Models
function Forward(observations, T, E, pi):
    alpha_0 = pi * E[obs_0]
    for t = 1 to T:
        alpha_t = E[obs_t] * (T^T * alpha_{t-1})
        normalize(alpha_t)
    return alpha_T

• Backward Algorithm
function Backward(observations, T, E):
    beta_T = [1, 1, ..., 1]
    for t = T-1 down to 0:
        beta_t = T * (E[obs_{t+1}] * beta_{t+1})
        normalize(beta_t)
    return beta_0

Equations
• Exponential Moving Average:  $x_n = \alpha x_{n-1} + (1 - \alpha)x_n$ 
• Log-Based Normalization:  $\frac{1}{n} \log(P(w_0, \dots, w_T)) = \frac{1}{n} \sum_{t=1}^T \log(P(w_t|w_{t-1}))$ 
• HMM Forward Pass:  $\alpha_t(j) = P(O_t|X_t = j) \sum_i \alpha_{t-1}(i) \cdot P(X_t = j|X_{t-1} = i)$ 
• Entropy:  $H(X) = - \sum_i P(x_i) \log_2 P(x_i)$ 
• Information Gain:  $IG(Y|X) = H(Y) - \sum_{x \in X} P(x)H(Y|X = x)$ 
• Learning rate decay:  $\alpha_t = \frac{1}{1+t}$ 
• L2 Regularization:  $\lambda ||w||_2^2 = \lambda \sum_i w_i^2$ 
• Sum of geometric series:  $\sum_{i=0}^{\infty} \gamma^i = \frac{1}{1-\gamma}$  for  $|\gamma| < 1$ 
• Conditional probability expansion:  $P(A, B|C) = P(A|B, C)P(B|C)$ 
• Law of Total Probability:  $P(A) = \sum_i P(A|B_i)P(B_i)$ 
• Marginalization:  $P(X) = \sum_y P(X, Y = y)$ 
• Product Rule:  $P(X, Y) = P(X|Y)P(Y) = P(Y|X)P(X)$ 
• Total Probability:  $P(Y) = \sum_x P(Y|X = x)P(X = x)$ 
• Independence:  $P(X, Y) = P(X)P(Y) \iff P(X|Y) = P(X)$ 
• Conditional Independence:  $P(X, Y|Z) = P(X|Z)P(Y|Z)$ 

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