

# AI Questions & Answers

Marco Bernardi      Andrea Auletta      Aulo

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## 1 Agents

**A.1** Explain in detail the meaning of the acronym PEAS in the context of the definition of an intelligent agent. According to the previous answer, introduce the various types of agents discussed in class.

**Answer:**

**PEAS** = Performance, Environment, Actuators, Sensors

PEAS helps us to define the task environment in which the agent is situated.

- **Performance:** is the criteria that defines how well the agent is doing in the environment. Performance with agents based on their different precepts. Performance measures could be a combination of different criteria. (Ex. Automated taxi driver: safety, legality, comfort, profit, time, etc.)

- **Environment:** It describes the task environment in which the agent is situated. An environment could be fully **observable** (Chess game) because you can see the whole state of the environment, otherwise it will be or **partially observable** (Poker).

It could be **deterministic** (chess game) because the next state of the environment is completely determined by the current state and the action executed by the agent, or **stochastic** (poker) because the next state of the environment can't be predicted in a certain way.

It could be **episodic** if the current action depend on the previous/next actions or **sequential** if the action are independent.

It could have only one agent or multiple agents.

It could be **static** if the environment doesn't change while the agent is deliberating or **dynamic** if the environment can change while the agent is deliberating.

Depending on the environment variables, the environment could be **discrete** or **continuous**.

(Ex. Automated taxi driver: roads and highways, other cars, pedestrians, traffic lights, etc.)

- **Actuators:** They are the mechanism or devices through which the agent acts upon the environment. Actuators translate decisions made by the agent into actions that can be performed on the environment. (Ex. Automated taxi driver: steering wheel, accelerator, brake, signal, etc.)
- **Sensors:** They are the mechanism or devices through which the agent perceives the environment. (Ex. Automated taxi driver: camera, GPS, speedometer, odometer, etc.)

An agent is a perceiving and acting entity, it is considered rational if it tries to achieve the best outcome given the available information.

Exist different types of agents:

- **Simple reflex agents:** they select actions on the basis of the current percept, ignoring the rest of the percept history. A simple reflex agent acts according to a rule whose condition matches the current state, as defined by the percept. The disadvantage of this type of agent is that if the environment is partially observable, then the agent may select a wrong action.
- **Model-based reflex agents:** they select actions on the basis of the current percept and some of the percept history. The disadvantage of this type of agent is that it has hard-coded rules, so it can't learn from experience and it can't be flexible.
- **Goal-based agents:** they select actions based on the goal they are trying to achieve, making the agent more flexible. The disadvantage of this type of agent is that it can have several goals that may conflict with each other.

- **Utility-based agents:** they select actions based on a utility function that measures the performance of the agent. The utility function helps the agent to choose the goal keeping in mind the likelihood of success and the importance of the goals. No remarkable disadvantages.
- **Learning agents:** they select actions based on the knowledge gained from the environment and past experiences, and there is a problem generator component that creates new situations to improve the agent's knowledge. No remarkable disadvantages.

## 2 Uninformed Search

**US.1** Describe the principal uninformed search strategies, and compare them in terms of correctness, completeness (Can we find a solution?), time and space complexity.

**Answer:** Search strategies are used to find a solution to a problem defined by four components: **initial state**, **successor function**, **goal test** and **path cost**. Uninformed search strategies use only the information provided by the problem definition and operate systematically to find a solution. The solution is a sequence of actions that leads from the initial state to a goal state. There are different uninformed search strategies (We're working with trees):

- (a) **Breadth-first search:** it explore the tree level by level, so the fringe is implemented as a FIFO queue. The disadvantage of this strategy is that it requires a lot of memory, because it has to visit all the nodes. It is **complete** (if branching factor  $b$  is finite) and **optimal** if the path costs are all equal. It has a **time complexity** of  $O(b^d)$  and a **space complexity** of  $O(b^d)$ .
- (b) **Uniform cost search:** if the path costs are all equal, it is equivalent to breadth-first search. Otherwise it expands the least-cost unexpanded node (the node with the lowest path cost) and it uses a priority queue (fringe) ordered by path costs. It is **complete** if the path costs are bounded below by a small positive constant and  $b$  is finite. It is **optimal**. It has a **time complexity** of  $O(b^{1+\lceil C^*/\epsilon \rceil})$  (it can be greater than BFS time complexity because

once it found a solution, it has to check if there is a better solution) and a **space complexity** of  $O(b^{1+\lceil C^*/\epsilon \rceil})$ .

- (c) **Depth-first search:** it explores the tree by expanding the deepest node in the current frontier of the search tree. The fringe is implemented as a LIFO queue. It is **not complete** because it can get stuck in an infinite path. It is **not optimal**. It has a **time complexity** of  $O(b^m)$  and a **space complexity** of  $O(bm)$ .

It requires less memory than breadth-first search, but it can get stuck in an infinite path.

- (d) **Iterative deepening search:** it is a combination of depth-first search and breadth-first search. It performs depth-first search up to a certain depth, then it performs breadth-first search up to that depth, and so on increasing the depth limit. It is **complete** if  $b$  is finite and **optimal** if the path costs are all equal. It has a **time complexity** of  $O(b^d)$  and a **space complexity** of  $O(bd)$ .

Its advantage is that it requires less memory than breadth-first search

- (e) **Depth-limited search:** it is a depth-first search with a depth limit that doesn't change. It isn't **complete** and not **optimal**. It has a **time complexity** of  $O(b^l)$  and a **space complexity** of  $O(bl)$ .
- (f) **Bidirectional search:** it performs two simultaneous searches, one forward from the initial state and one backward from the goal. It is **complete** if  $b$  is finite and both searches use breadth-first search. It is **optimal** if the path costs are all equal and both searches use breadth-first search. It has a **time complexity** of  $O(b^{d/2})$  and a **space complexity** of  $O(b^{d/2})$ .

May not find the optimal solution, but if it's applicable it is faster and lighter than breadth-first search.

### 3 Informed Search

- IS.1** Define the concept of informed search, and describe the greedy search and the A\* algorithm. Discuss the properties (optimality, completeness, time and space complexity) and conditions of applicability of the two algorithms.

### Answer:

Informed search algorithms use problem-specific knowledge beyond the definition of the problem itself. Greedy search and A\* algorithm are some special case of best-first search. Best-first search is a search algorithm that explores a graph by expanding the most promising node chosen according to a specified rule. The rule is specific to the problem, defined by an evaluation function  $f(n)$  that estimates the cost of the cheapest path from the node  $n$  to a goal node. Most of the best-first search algorithms include a heuristic function  $h(n)$  in  $f(n)$  that estimates the cost of the cheapest path from the node  $n$  to a goal node.

- **Greedy search:** it expands the node that appears to be closest to goal, according to the heuristic function  $h(n)$ . In this case  $f(n) = h(n)$ . It is **not complete** because it can get stuck in a loop. In a finite space with repeated-state checking it is complete. It is **not optimal**. It has a **time complexity** of  $O(b^m)$  and a **space complexity** of  $O(b^m)$ .
- **A\* algorithm:** The idea at the base of this algorithm is to avoid expanding paths that are already expensive. It expands the node that has the lowest value of  $f(n) = g(n) + h(n)$ . Where:
  - $g(n)$  is the cost of the path from the initial state to the node  $n$ .
  - $h(n)$  is the heuristic function that estimates the cost of the cheapest path from the node  $n$  to a goal node.
  - $f(n)$  is the evaluation function that estimates the cost of the cheapest solution through  $n$ .

A\* uses an admissible heuristic function, that is a heuristic function that never overestimates the cost to reach the goal. It is **optimal** for tree search because it will never select a suboptimal goal  $G_2$  over an optimal goal  $G_1$ , since  $f(G_2) > f(n)$  where  $n$  is an unexpanded node that leads to  $G_1$ . For graph search the previous statement is not true, and we need another proof that use consistent heuristic. A heuristic is consistent if  $h(n) \leq c(n, a, n') + h(n')$  where  $n$  is the current node,  $a$  is the action that leads to  $n'$  and  $n'$  is the successor of  $n$ .  $f(n)$  is non-decreasing along any path. It is **complete**, unless there are infinitely many nodes with  $f(n) \leq$

$f(G)$ . It has a **time complexity** exponential and a **space complexity** of  $O(\text{nodescount})$ .

No other optimal algorithm is guaranteed to expand fewer nodes than  $A^*$ .

$A^*$  could be exponential in space  $\Rightarrow$  Solutions:

- **Iterative deepening  $A^*$  ( $IDA^*$ )**: it acts like  $A^*$  but it use a cutoff value instead of a depth limit. During an iteration if  $f(n) > \text{cutoff}$  then the node is not expanded. When the queue is empty, the cutoff is increased to the lowest value of  $f(n)$ .
- **Recurisve Best-First Search**: imitates a deep search, using only linear space. It keeps track of the best alternative path available from any ancestor of the current node. When a node is expanded, the algorithm updates the value for the best alternative path. If the value of the best alternative path is smaller than the value of the current node, recursion goes back to the alternative path. On the return from the recursion, the algorithm updates the f-value of the best child node.

It is **optimal** if the heuristic is consistent. Space complexity is  $O(bd)$  and time complexity is exponential in the worst case. RBFS has a problem, it use too little memory.

- **Memory-bounded  $A^*$  ( $MA^*$ )**:
- **Simplified memory-bounded  $A^*$  ( $SMA^*$ )**: it expand the best node until the memory is full. It removes the worst node from the memory to make space for a new node and backup the f-value of the removed node on the parent node. The parent node will eventually be expanded if the other paths are worse.

It is **complete** only if solution can be kept in memory. It is **optimal** if any optimal solution is reachable otherwise it returns the best solution found.

**IS.2** Introduce the  $A^*$  algorithm in detail and exhaustively. Formally prove its optimality prop-erties. Finally, discuss the memory usage issues of  $A^*$  and how this can be solved/reduced.

**IS.3** Describe the concept of admissible heuristic function, and give the formal definition of heuristic admissible and consistent. Choose a domain and give an example of two heuristic functions  $h_1$  and  $h_2$ , they are both admissible and  $h_1$  dominates  $h_2$ . Discuss how a heuristic function can be systematically constructed.

**Answer:**

A heuristic function is a function that estimates the cost of the cheapest path from the node  $n$  to a goal node. A heuristic function is **admissible** if it never overestimates the cost to reach the goal ( $h(n) \leq h^*(n)$ ). A heuristic function is **consistent** if  $h(n) \leq c(n, a, n') + h(n')$  where  $n$  is the current node,  $a$  is the action that leads to  $n'$  and  $n'$  is the successor of  $n$ . Given two heuristic functions  $h_1$  and  $h_2$ ,  $h_1$  dominates  $h_2$  if  $h_1(n) \geq h_2(n)$  for all nodes  $n$ ; for research purposes, it's better to use the dominant heuristic function. (but should not be too expensive to compute)

Example 8-puzzle:

- $h_1(n)$  = number of misplaced tiles
- $h_2(n)$  = total Manhattan distance

An heuristic function can be systematically constructed by relaxing the problem. In the 8-puzzle example, the relaxed problem is to move the tiles without considering the other tiles. The optimal solution cost of a relaxed problem is no greater than the optimal solution cost of the real problem.

## 4 Iterative Improvement Algorithms

**II.1** Introduce hill-climbing search, appropriately placing it among the various categories of problem solving approaches we discussed in class. Present the variants, the properties, and in the case of a search with restart, formally demonstrate the result relative to the number of expected searches before finding an optimal solution.

**Answer:** Hill-climbing search is an iterative improvement algorithm making part of the local search algorithms. Hill-climbing follow the

iterative improvement paradigm, where the goal state itself is the solution and the path to reach it is irrelevant. As a local search algorithm, it doesn't keep track of the search tree, but only of the current state and try to improve it. Although local search algorithms are not systematic, they have two key advantages:

- (a) they use very little memory, usually a constant amount;
- (b) they can often find reasonable solutions in large or infinite (continuous) state spaces for which systematic algorithms are unsuitable.

Hill-climbing search at each iteration selects the best successor among the neighbors of the current state, and replaces the current state with it. (or lowest heuristic cost if a heuristic function is used) Hill-climbing is **not complete** because it never makes downhill moves, so it can get stuck in a local maximum.

Problems:

if the neighbors solutions are equivalent it can get stuck in a shoulder, otherwise if the neighbors solutions are worse it can get stuck in a local maximum. If we've ridges (a sequence of local maxima not directly connected to each other) it will be difficult to move from one local maximum to another. In continuous space, it is difficult to pick the "right" step size and convergence can be very slow.

For outlast this problems, we can use:

- plateaux (flat values for  $h$ ):
  - allow sideways moves (moves to neighbors with equal  $h$  value), but if there are no uphill moves (flat maximum), an infinite loop will occur; so for outlast this side-problem we can put a limit on the number of sideways moves.
- local maxima:
  - **Stochastic hill climbing**: choose at random from among the uphill moves; the probability of selection can vary with the steepness of the uphill move. This usually converges more slowly than regular hill climbing, but it can drive to better solutions.
  - **random restart hill climbing**: it conducts a series of hill-climbing searches from randomly generated initial states. If  $p$



is the probability to find an optimal solution in a single search, the expected number of searches before finding an optimal solution is  $1/p$ .

We can derive the probability  $1/p$  as follows:

$$x_i = \begin{cases} 0 & \text{if } i\text{-th search doesn't find an optimal solution} \\ 1 & \text{if } i\text{-th search finds an optimal solution} \end{cases} \quad (1)$$

We know that  $\forall i, p = P(x_i = 1) = p$ , and  $P(x_i = 0) = 1 - p$ . Variables  $x_i$  are mutually independent, we have a Bernoulli Process for which we can use the geometric distribution to calculate the expected number of searches before finding an optimal solution. The probability that the  $k$ -th search finds an optimal solution is

$$P\left(\left(\sum_{j=1}^{k-1} x_j = 0\right) \wedge (x_k = 1)\right) = (1 - p)^{k-1} p \quad (2)$$

The expected value of the quantity we are interested in is then:

$$\begin{aligned} \sum_{k=1}^{\infty} k(1 - p)^{k-1} p &= p \sum_{k=1}^{\infty} k(1 - p)^{k-1} \\ &= -p \sum_{k=1}^{\infty} d \frac{(1 - p)^k}{dp} \\ &= -p \frac{d}{dp} \left( \sum_{k=0}^{\infty} (1 - p)^k - 1 \right) = -p \frac{d}{dp} \left( \frac{1}{1 - p} - 1 \right) \\ &= p \frac{1}{p^2} = \frac{1}{p} \end{aligned} \quad (3)$$

For random walk algo is **complete** but extremely inefficient.

## II.2 Local search algorithms:

**Answer:**

- **hill climbing:** refer to previous question ??.

- **simulated annealing**(gradient descent): it's a combination between hill climbing and random walk. It allows "bad" moves, but gradually decreases their size and frequency. The probability of selecting a "bad" move is controlled by a parameter  $T$  called "temperature",  $T$  is gradually decreased. The algorithm halts when a certain criterion is met (e.g.  $T$  is less than a certain threshold). Bad moves are more likely to be selected at the beginning, and less likely as the temperature decreases. If  $T$  decreased slowly enough  $\Rightarrow$  always reach best state  $x^*$ .
- **local beam search**: it keeps track of  $k$  states instead of one. It starts with  $k$  randomly generated states, and at each iteration it generates all the successors of the  $k$  states and selects the best  $k$  successors. It halts when a goal state is found or when a certain criterion is met, otherwise it selects the best  $n$  (beam width) successors and repeat the process.  
**Problem**: often all  $k$  states end on same local hill.  
**Idea**: choose  $k$  successors randomly, biased towards better states: stochastic beam search.
- **genetic algorithms**: it's a variant of stochastic beam search in which successors states are generated by combining two parent states rather than by modifying a single state.
  - (a) are considered the best states of the current population
  - (b) they are combined to generate a new population
  - (c) there is a mutation probability  $p_m$  that a state is randomly modified to explore new areas of the search space
  - (d) the algorithm try to converge to an acceptable solution.

## 5 Online Search

**OS.1** Introduce the concept of online search, and describe the algorithms we discussed in class.

**Answer**: Online search algorithms are used when the environment is partially observable, or it is dynamic/semidynamic the agent needs to interact with the environment to get information about it. It's not possible to compute a complete solution before starting to act, so the

agent has to interleave computation and action. Online search work well for exploration problems.

**Problem:** if some actions are irreversible, the agent may get stuck in a dead end (THIS IS NOT AVOIDABLE).

The performance of an online search algorithm is measured by competitive ratio:

$$\frac{\text{total path cost online search}}{\text{total path cost knowing state space in advance}} \quad (4)$$

Type of online search algorithms:

- **Depth-first online search:** it's a depth-first search that keep track of the current path. When the agent reaches a dead end, it backtracks until it finds a node with an unexplored successor. For the backtracking the algorithm uses a table that maps each node to its unbacktracked parent node.

It iterates this process until it reaches the goal state. It is **complete** if the state space is finite and **optimal**. It has a **time complexity** of  $O(b^m)$  and a **space complexity** of  $O(bm)$ .

- **Random search:** it's like hill climbing, but it chooses a random successor instead of using random restart. With this approach, the agent could escape from local maxima. Random walk will eventually find a solution if it exists (if finite state spaces), but it could take a long time.
- **LRTA\* search (Learning Real-Time A\*):** it's a combination of hill climbing with memory + a strategy to overcome local optima. It store a "current best estimate"  $H(s)$  of the cost of reaching the goal from each state that has been visited.  $H(s)$  starts out being just the heuristic estimate  $h(s)$ , but it is updated as the agent moves around the state space. Based on the current best estimate, the agent chooses the action that appears to lead most quickly to the goal. When the agent reaches a new state, it updates the current best estimate of the cost of reaching the goal from the previous state.

This optimism under uncertainty encourages the agent to explore new, possible promising paths. It is **complete** if the state space is

finite. It can explore an environment in  $O(n^2)$  steps in the worst case.

## 6 Adversarial Search

**AS.1** In the context of adversarial search, explain in detail how games with elements of chance can be deal with. In the case of resources limit, explain what is the property of the evaluation function that allows the search to preserve optimal decision (i.e. obtained with no resources limit). Finally, explain why the same approach is not returning an optimal strategy in the case of partially observable games.

**Answer:** In the context of adversarial search, non deterministic games are games with elements of chance introduced by some events like dice rolls or card-shuffling. There are two types of non deterministic games: **perfect information** (backgammon or monopoli) and **imperfect information** (card games). We can deal with non deterministic games by using  $\alpha - \beta$  pruning, that is a generalization of the minimax algorithm.

**Minimax algorithm** is a recursive algorithm for choosing the next move in an n-player game, it achieve the best payoff assuming that the opponent is also playing optimally. It is based on the idea of **minimizing the maximum loss**, that is the best we can do is to choose the move that minimize the maximum loss. Minimax it's **complete** if tree is finite and **optimal** if both players play optimally. It has a **time complexity** of  $O(b^m)$  and a **space complexity** of  $O(bm)$  (depth-first exploration).

**Minimax** can explore the whole tree also if it's not necessary, so we can use  $\alpha - \beta$  pruning to avoid this problem: if it find a value  $n$  less than the upper bound of another path, it halts the exploration of the current path.

$\alpha$  = best choice for the max player along the path to the root.

$\beta$  = best choice for the min player along the path to the root.

**Properties** of  $\alpha - \beta$  pruning:

- pruning doesn't affect final result
- a good ordering of the moves improves the effectiveness of pruning

- perfect order (MAX: from highest to lowest, MIN: from lowest to highest)  $\Rightarrow$  time complexity (almost)  $O(b^{m/2})$
- doubles the depth of exploration

$\alpha - \beta$  pruning suite well for games with deterministic games; it work also for non-deterministic games. In the case of non-deterministic games, the states doesn't have definite values, we can compute only the expected value of a state. **Expectiminimax algorithm** help us to handle this problem, that works like minimax except that it also handle chance nodes. It compute the expected value of a state by multiplying the probability of each outcome by the utility of that outcome. The upper bound of a node is the sum of the expected values of its children. Stronger pruning can be obtained if possible to bound the leaves values.

**Problem:** Resources limit, in real world we have limited resources, so we can't explore the whole tree.

**Solution:** evaluation function & cutoff test.

An evaluation function is used to estimate the expected utility of the game from a given state. An EV function should order the terminal states: states that are wins must evaluate higher than states that are draws or losses. The computation must not take too long. Most EV function work by calculating various features of the states. With a good EV function, we can explore the tree to a limited depth (CUTOFF), obtaining a good approximation of the true minimax value. Cutoff should be apply only to quiescent states (states in which the game is unlikely to change drastically in the near future). Non quiescent states can be expanded further until a quiescent state is reached.

With this approach, there is still a difficult problem to outlast: the horizon effect. The horizon effect is the problem that the agent can't see beyond a certain depth, so it can't see a threat or an opportunity. This cannot be avoided, but we can temporarily avoid it by delaying tactics; for mitigate it we can use the **SINGULAR EXTENSION** (Find and remember move that is clearly better than others, and consider it if is it legal when the search reaches the normal depth limit).

The previous approaches are not returning an optimal strategy in the case of partially observable games; in this types of games, the agent doesn't know the initial state of the game. We can try to predict the

optimal move with  $N$  deals, during this process all the possible moves are considered and weighted by their probability to get the expected value of the state, will be picked the action that wins most tricks on average; this will lead most of the time to a non-optimal strategy. In this type of games, value of an action depends on the information state or belief state the agent is in.

**AS.2** In the context of adversarial search, discuss how huge search spaces, such as those generated by the game of chess or Go, can be computationally managed. Also explain how the algorithm  $\alpha - \beta$  pruning can help in such cases.

**Answer:**

**AS.3** Describe the  $\alpha - \beta$  pruning algorithm in detail. Prove its complexity in the best case.

**Answer:**

## 7 Propositional Logic

**PL.1** Give the abstract definition of knowledge-based agent, discussing its various components. In addition, in the case of propositional logic, introduce how inference can be made by enumeration, discussing its algorithmic and computational complexity aspects.

**Answer:** A knowledge-based agent is an agent that uses an internal representation of the world to decide what actions to perform. The central component of a knowledge-based agent is the **knowledge base**, that is a set of sentences in a knowledge representation language and it represents the agent's knowledge about the world and may initial contain some background knowledge.

A knowledge-based agent has also a **inference engine**, a component that uses the knowledge base to infer new information. Usually you can communicate with the agent through these operations: *tell* and *ask*.

A knowledge-based agent must be able to:

- Represent states, actions, etc.
- Incorporate new percepts

- Update internal representations of the world
- Deduce hidden properties of the world
- Deduce appropriate actions

In the case of propositional logic, inference can be made by enumeration, that is a general algorithm for enumerating all the models of a sentence. The algorithm consists in enumerating all models, and check that sentence  $\alpha$  is true in every model in which  $KB$  is true ( $KB \models \alpha$ ). Models are assignments of *true* or *false* to each propositional symbol. The algorithm is **sound**<sup>1</sup> because it implements directly the definition of entailment, and it is **complete**<sup>2</sup> because it works for any  $KB$  and  $\alpha$  and always terminates.

If  $KB$  and  $\alpha$  contain  $n$  symbols in all, then there are  $2^n$  models. Thus, the time complexity of the algorithm is  $O(2^n)$  and the space complexity is  $O(n)$  because the enumeration is depth-first. Unfortunately, this algorithm is co-NP-complete, so every known inference algorithm for propositional logic has a worst-case time complexity that is exponential in the size of the input.

**PL.2** In the context of propositional logic, explain in detail the Forward-chaining algorithm, discussing its algorithmic and computational complexity aspects.

**Answer:**

Given a knowledge base  $KB$  the idea is to fire any rule whose premises are satisfied in  $KB$ , and add the conclusion to  $KB$  until query is found.

- Set the known leaves ( $KB$ )
- Inference propagates up the graph as far as possible
- if conjunction are found, waits until all conjuncts are known before proceeding

The algorithm is **sound** (every inference is an application of modus ponens, we're working with Horn clauses) and **complete** (every entailed atomic sentence will be derived). Proof of completeness:

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<sup>1</sup>Soundness:  $i$  is sound if whenever  $KB \vdash_i \alpha$  (sentence  $\alpha$  can be derived from  $KB$  by inference procedure  $i$ ), it is also true that  $KB \models \alpha$

<sup>2</sup>Completeness:  $i$  is complete if whenever  $KB \models \alpha$ , it is also true that  $KB \vdash_i \alpha$

- (a) FC reaches a fixed point (no new atomic sentences can be inferred)
- (b) Consider the final state as a model  $m$ , assigning true/false to symbols (inferred table: the table contains true for every inferred symbol, false otherwise)
- (c) every clause in the original KB is true in  $m$  (because it's a fixed point) Proof: assume the opposite, then there is a clause that is false in  $m$ :  
 $a_1 \wedge \dots \wedge a_k \Rightarrow b$  is false in  $m$ .  
 $a_1 \wedge \dots \wedge a_k$  in the model must be true, so  $b$  must be false, but  $b$  is in the inferred table, so it must be true  $\Rightarrow$  contradiction: the algorithm has not reached a fixed point.
- (d)  $m$  is a model of  $KB$
- (e) if  $KB \models q$ ,  $q$  is true in every model of  $KB$ , including  $m$

It runs in **linear time**.

Forward-chaining is an example of the general concept of **data-driven** (unconscious processing) reasoning, that is the reasoning is driven by known data. One disadvantage of this approach is that it may do a lot of work that is irrelevant to the goal.

**PL.3** In the context of propositional logic, explain in detail the Backward-chaining algorithm, discussing its algorithmic and computational complexity aspects.

**Answer:** Backward-chaining is a goal-driven algorithm (unlike forward-chaining that is data-driven), that is the reasoning is driven by the query. In fact the idea is to work backward from the query  $q$ . If  $q$  is known true, then no work is needed; otherwise we must find the premises that imply  $q$ . Avoid repeated work by keeping track of sub-goals that have already been proved true, and avoid loops by checking if a subgoal is already on the goal stack. Complexity of BC can be much less than linear in the size of the KB.

**PL.4** In the context of inference in propositional logic, explain what is the role of horns form, discussing its strengths and weaknesses.

**Answer:**

Truth methods (are useful for testing the validity of a sentence) divide into two classes:



- **Model checking:** it checks if a given model satisfies a sentence. (Truth table enumeration is an example of model checking)
- **Application of inference rules:** it uses logical entailment to determine the truth of a sentence. (Generation of new sentences from old ones)

Truth table enumeration holds a problem: it is exponential in the number of propositional symbols. We need to find a more efficient way to determine if  $\alpha$  is entailed by  $KB$ . Basically, we need to find the equivalent sentences written in a different way, so we can avoid them. Some real-world knowledgebases satisfy certain restrictions on the form of sentences they contain, which enables them to use a more restricted and efficient inference algorithm. One such restricted form is the **Horn form**.

With the horn form  $KB$  is a conjunction of Horn clauses<sup>3</sup>:

- proposition symbol
- (conjunction of symbols)  $\Rightarrow$  symbol

Every definite clause can be written as an implication whose premise is a conjunction of positive literals and whose conclusion is a single positive literal.

**Modus Ponens** is a inference rule that can be used with the horn form:

$$\frac{\alpha_1, \dots, \alpha_n, \quad \alpha_1 \wedge \dots \wedge \alpha_n \Rightarrow \beta}{\beta} \quad (5)$$

Horn form simplifies the inference process, lowering the complexity from exponential to polynomial. Lowering the complexity is a huge improvement, but it's come with a cost: less expressive power.

It can be used with the **forward chaining** and **backward chaining** algorithms.

**PL.5** In the context of inference in propositional logic, introduce the resolution rule. Then, explain how the resolution algorithm works. Finally, prove its completeness.

**Answer:**

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<sup>3</sup>Horn clauses are a disjunction of literals with at most one positive literal

Resolution yields a complete inference algorithm when coupled with any complete search algorithm. This rule applies only to clauses that is a disjunction of literals: every sentence of propositional logic can be converted into an equivalent conjunction of clauses  $\Rightarrow$  conjunctive normal form (CNF).

Resolution rule:

$$\frac{\ell_1 \vee \dots \vee \ell_k, \quad m_1 \vee \dots \vee m_n}{\ell_1 \vee \dots \vee \ell_{i-1} \vee \ell_{i+1} \vee \dots \vee \ell_k \vee m_1 \vee \dots \vee m_{j-1} \vee m_{j+1} \vee \dots \vee m_n} \quad (6)$$

Resolution algorithm:

1. Convert  $KB \wedge \neg\alpha$  to CNF
  - i. Eliminate  $\Leftrightarrow$  replacing  $\alpha \Leftrightarrow \beta$  with  $(\alpha \Rightarrow \beta) \wedge (\beta \Rightarrow \alpha)$
  - ii. Eliminate  $\Rightarrow$  replacing  $\alpha \Rightarrow \beta$  with  $\neg\alpha \vee \beta$
  - iii. Move  $\neg$  inwards using De Morgan's laws and double negation
  - iv. Apply distributivity law ( $\vee$  over  $\wedge$ ) and flatten
2. Apply resolution rule to resulting clauses
3. Each pair that contains complementary literals is resolved to produce a new clause
4. The process is repeated until one of two things happens:
  - there are no new clauses that can be added,  $KB$  does not entail  $\alpha$
  - two clauses resolve to yield the empty clause  $\Rightarrow \alpha$  is entailed by  $KB$  (The empty clause is equivalent to false, because disjunction is true only if at least one of the disjuncts is true)

The algorithm is sound because it implements directly the definition of entailment, and it is complete because it works for any  $KB$  and  $\alpha$  and always terminates.

### **Proof of completeness:**

The completeness theorem for resolution in propositional logic is called the ground resolution theorem:

*If a set of clauses is unsatisfiable, then the resolution closure of those clauses contains the empty clause. (Resolution Closure: the set of all*

*clauses that can be derived by repeated application of the resolution rule)*

The proof of this theorem is proved by demonstrating its contrapositive:

*If the resolution closure of a set of clauses does not contain the empty clause, then the set of clauses is satisfiable.*

- (a) Construct a model for  $S$  with suitable truth value for  $P_1, \dots, P_k$ :  
 For  $i$  from 1 to  $k$ :
- if a clause in  $RC(S)$  contains  $\neg P_i$  and all its other literals are false under the assignment chosen for  $P_1, \dots, P_{i-1}$ , then assign false to  $P_i$ .
  - Otherwise, assign true to  $P_i$ .
- (b) Show that the the previous point always ends up with a model for  $S$ , this is done by induction:
- Base case:  $i = 1$ :  
 cannot be than  $\neg P_1$  and  $P_1$  simultaneously appear in  $S$  otherwise the empty clause will appear in  $RC(S)$ . So  $P_1 \leftarrow false$  if  $\neg P_1$  occurs in  $S$ , otherwise  $P_1 \leftarrow true$ .
  - Case  $i$ : Assume a partial model  $m_{i-1}$  for symbols  $P_1, \dots, P_{i-1} \Rightarrow$  it is not possible to assign a truth value to  $P_i$  if  $RC(S)$  contains clauses:
    - $C \equiv B \vee P_i$  and  $C' \equiv B' \vee \neg P_i$ , where  $B$  and  $B'$  only contain symbols in  $\{P_1, \dots, P_{i-1}\}$
    - both  $B$  and  $B'$  are false under  $m_{i-1}$
 In the case we had these clauses the resolution of them will return  $B \vee B'$ , so either  $B$  or  $B'$  should be true by the induction hypothesis. This means that:
    - if  $B$  is true  $\Rightarrow P_i$  is false
    - if  $B'$  is true  $\Rightarrow P_i$  is true

Thus it will be possible to extend model  $m_{i-1}$  to either  $m_i = m_{i-1} \cup \{P_i \leftarrow false\}$  or  $m_i = m_{i-1} \cup \{P_i \leftarrow true\}$ . So it's possible to extend the model up to  $k$  and  $S$  is satisfiable.

## 8 First-Order Logic

**FOL.1** Describe the fundamental elements to be able to carry out inference in the first-order logic through the resolution rule. Discuss the various strategies used by the resolution rule, highlighting the strengths and weaknesses. At the end discuss the computational aspects.

**Answer:** The fundamental elements of first-order logic are:

- **Constants:** represent objects in the domain of discourse.
- **Predicates:** represent properties of objects in the domain of discourse.
- **Functions:** represent relations between objects in the domain of discourse.
- **Variables:** represent unspecified objects in the domain of discourse.
- **Connectives:**  $\wedge, \vee, \neg, \Rightarrow, \Leftrightarrow$ .
- **Quantifiers:**  $\forall, \exists$ .
- **Equality:**  $=$ .

Sentences are true with respect to a model and an interpretation.

The idea is to use the quantifiers to do inference and reconduct to the inference in propositional logic, example:

- **Universal quantification:**  $\forall x P(x)$ , it's true in a model  $m \iff P$  is true with  $x$  being each possible object in the model.

**Universal instantiation:** every instance of a universally quantified sentence is entailed by:

$$\frac{\forall v \quad \alpha}{SUBST(\{v/g\}, \alpha)} \quad (7)$$

for any variable  $v$  and ground term  $g$ . It can be applied several times to add new sentences, the new  $KB$  is equivalent to the original one.

- **Existential quantification:**  $\exists x P(x)$ , it's true in a model  $m \iff P$  is true with  $x$  being at least one object in the model.

**Existential instantiation:** for any sentence  $\alpha$ , variable  $v$  and constant symbol  $k$  that does not appear elsewhere in the KB:

$$\frac{\exists v \quad \alpha}{SUBST(\{v/k\}, \alpha)} \quad (8)$$

EI can be applied only once to replace the existential sentence, the new  $KB$  is not equivalent to the original one, but it is satisfiable if the original one is.

**FOL.2** In the context of first order logic, describe in a complete and exhaustive way the Unification algorithm, further explaining why this is particularly useful for logical inference

**Answer:**

Propositional logic seems to generate lots of irrelevant sentences:

With  $p$   $k$  – ary Predicates and  $n$  constants, there are  $p \cdot n^k$  instantiations.

If there is some substitution  $\theta$  that makes each of the conjuncts of the premise of the implication identical to sentences already in the KB, then we can assert the conclusion of the implication. This inference process is called **Generalize Modus Ponens**:

$$\frac{p_1', p_2', \dots, p_n', \quad (p_1 \wedge p_2 \wedge \dots \wedge p_n \Rightarrow q)}{q\theta} \quad (9)$$

Soundness GMP:

For any definite clause  $p$ , we have  $p \models p\theta$  by UI.

- $(p_1 \wedge \dots \wedge p_n \Rightarrow q) \models (p_1 \wedge \dots \wedge p_n \Rightarrow q)\theta = (p_1\theta \wedge \dots \wedge p_n\theta \Rightarrow q\theta)$
- $p_1' \wedge \dots \wedge p_n' \models p_1' \wedge \dots \wedge p_n' \models p_1'\theta \wedge \dots \wedge p_n'\theta$
- From 1 and 2,  $q\theta$  follows by ordinary Modus Ponens.

**Unification** is a process that finds substitutions for variables that make two sentences identical. We can get the inference immediately if we can find a substitution  $\theta$  such that  $\alpha\theta = \beta\theta$ . For avoiding failures, we need to eliminate overlap of variables in  $\alpha$  and  $\beta \Rightarrow$  **Standardizing apart**:

each sentence have to use different variables name (we can rename variables in a sentence without changing its meaning).

**Unification algorithm:** The algorithm works by comparing the structures of the inputs, element by element. The substitution  $\theta$  that is the argument to UNIFY is built up along the way and is used to make sure that later comparisons are consistent with bindings that were established earlier.

**Problem:** there could be more than one such unifier  $\Rightarrow$  for each pair of sentences we need to find the most general unifier (MGU: that one which places the fewest restrictions on the variables), that is unique up to renaming and substitution of variables.

This algorithm is particularly useful for logical inference because it is used in algorithms for logical inference like forward/backward chaining and resolution.

**FOL.3** In the context of inference in first-order logic, introduce the resolution rule. Discuss in what form the clauses need to be represented in order to efficiently enforce the resolution and how that form can be achieved. Finally, present the various strategies that have been proposed regarding the choice of clauses to be used during the inference.

**Answer:**

The resolution rule is an inference procedure based on the resolution for propositional logic and extended to first-order logic. It requires sentences to be in **conjunctive normal form** (CNF: a conjunction of clauses, where a clause is a disjunction of literals). For achieving this form, we must follow these steps:

- (a) **Eliminate implications:**  $p \Rightarrow q$  is equivalent to  $\neg p \vee q$ .
- (b) **Move negation inwards:** we need rules for negated quantifiers:
  - $\neg \forall x P(x)$  is equivalent to  $\exists x \neg P(x)$
  - $\neg \exists x P(x)$  is equivalent to  $\forall x \neg P(x)$
- (c) **Standardize variables:** rename variables to avoid conflicts.
- (d) **Skolemize:** eliminate existential quantifiers by replacing it with Skolem functions. In the existential instantiation rule, we drop the quantifier and substitute the variable with another one creating only ONE new sentence. When the existential instantiation

can't be applied (The sentence doesn't match the pattern  $\exists v \alpha$ ), we replace the existential quantifier variables with Skolem functions which has as arguments the universally quantified variables in whose scope the existential quantifier appears.

- (e) **Drop universal quantifiers:** we can drop universal quantifiers because they are implicit in CNF.
- (f) **Distribute  $\vee$  over  $\wedge$ :**  $p \vee (q \wedge r)$  is equivalent to  $(p \vee q) \wedge (p \vee r)$ .

Once the sentences are in CNF, we can apply the resolution rule:

Two sentences (standing apart) are resolvable if they contain complementary literals: propositional literals are complementary if one unifies with the negation of the other. Thus we have:

$$\frac{\ell_1 \vee \dots \vee \ell_k, \quad m_1 \vee \dots \vee m_n}{SUBST(\theta, \ell_1 \vee \dots \vee \ell_{i-1} \vee \ell_{i+1} \vee \dots \vee \ell_k \vee m_1 \vee \dots \vee m_{j-1} \vee m_{j+1} \vee \dots \vee m_n)} \quad (10)$$

where  $UNIFY(\ell_i, \neg m_j) = \theta$ .

This rule is called binary resolution because it combines two clauses at a time.

The various strategies that have been proposed regarding the choice of clauses to be used during the inference are:

- **Unit preference:** select a unit clause (a clause with only one literal) if possible, the idea is that we're trying to produce an empty clause, so it might be a good idea to prefer inferences that produce shorter clauses.
- **Unit resolution:** is a restricted form of resolution in which every resolution step must involve a unit clause. It is incomplete in general, but it is complete for Horn clauses.
- **Set of support:** select a set of clauses (the set of support) and use at least one element of the set in each step. It is incomplete if the wrong set of support is chosen (Ex of a good set: the negated query)
- **Input resolution:** every resolution combines one of the input sentences (from the KB or the query) with some other sentence. It is complete if KB is in Horn form, but incomplete in the general case.

Complete if modified: allows  $P$  and  $Q$  to be resolved together either if  $P$  is in the original KB or if  $P$  is an ancestor of  $Q$  in the proof tree.

- **Subsumption:** eliminates all sentences that are subsumed by other sentences in the KB.

## 9 Uncertainty

Agents may need to handle uncertainty, whether due to partial observability, nondeterminism, or a combination of the two. Agents that never fail must to know everything about the world and the various exception, but this is not possible in the real world:

- **Laziness:** it's too expensive to enumerate all the possible cases.
- **Theoretical ignorance:** most of the time we don't know the complete domain of the problem.
- **Practical ignorance:** even if we know all the possible cases, there is uncertainty in applying this rules in some specific cases.

Probability provides a way of summarizing the uncertainty that comes from our laziness and ignorance, thereby solving the qualification problem (the problem of how to represent all the knowledge that an agent needs to know about the world). In **subjective** (degree of belief) or **Bayesian** (degree of belief given the evidence) probability relate propositions to one's own state of knowledge (it might be learned from past experience of similar situations). Decision under uncertainty  $\Rightarrow$  Decision theory = Probability theory + Utility theory (used to represent and infer preferences).

**U.1** In the context of treatment of uncertainty, discuss what is the main computational challenge when using the joint probability distribution of the main involved variables.

**Answer:**

**Definition:** for a set of random variables (r.v.s)  $X_1, \dots, X_n$ , the joint probability distribution  $P(X_1, \dots, X_n)$  is a table that contains the probability of each possible combination of values for the random variables.



Joint probability distribution is used for inference by enumeration: for any proposition  $\phi$  sum the atomic events where it is true:

$$P(\phi) = \sum_{\omega: \omega \models \phi} P(\omega) \quad (11)$$

It works also for conditional probability:

$$P(a|b) = \frac{P(a \wedge b)}{P(b)} \text{ if } P(b) \neq 0 \quad (12)$$

In the ?? the denominator act as a normalization factor, it is the probability of the event  $b$ .

Inference procedure:

- $X$  is a set of random variables
- $Y$  is the set of query variables
- $E$  is the set of evidence variables given specific values  $e$
- $H$  is the set of hidden variables ( $H = X - Y - E$ )

Then the required summation of joint entries is done by summing out the hidden variables:

$$P(Y|E = e) = \alpha P(Y, E = e) = \alpha \sum_h P(Y, E = e, H = h) \quad (13)$$

The terms in the summation are joint entries because  $Y$ ,  $E$  and  $H$  are together exhaust the set of random variables.

The main computational challenge is that the number of possible worlds grows exponentially with the number of random variables. Worst-case time complexity  $O(d^n)$ , where  $d$  is the maximum number of values that any variable can take on (arity). Space complexity  $O(d^n)$ , because we need to store the probability of each possible combination of values.

**U.2** In the context of the treatment of uncertainty, formally define the concept of conditional independence and how this can be used. In particular, discuss its role in the Bayesian Networks.

**Answer:**

**Definition:** two random variables  $X$  and  $Y$  are independent given a set of random variables  $Z$  if and only if:

$$P(X|Y) = P(X) \text{ or } P(Y|X) = P(Y) \text{ or } P(X, Y) = P(X)P(Y) \quad (14)$$

This means that in a given set of random variables, if we can find independent variables, then the full joint distribution can be factored into smaller separated joint distributions on the subsets composed by the variables that are related to each other (size of the representation from exponential in  $n$  to linear in  $n$ ). In this way the full joint distribution can be represented as the product of the smaller joint distributions. In real world problems it's difficult to find a clean separation of variables by independence. Conditional independence is our most basic and robust form of knowledge about uncertain environments.

Conditional independence:

$$P(X|Y, Z) = P(X|Z)P(Y|Z) \quad (15)$$

Full joint distribution:

$$P(Cause, Effect_1, \dots, Effect_n) = P(Cause) \prod_i P(Effect_i | Cause) \quad (16)$$

Such probability distributions is called **naïve Bayesian** model

In a Bayesian network, conditional independence is used to define the network topology. The topological semantics specifies that each variable is conditionally independent of its nondescendants, given its parents.

**U.3** What does a Bayesian Network consist of? Why is it useful? What is its computational complexity in space and time (also discuss special cases) for the exact inference?

**Answer:**

A Bayesian network is a directed graph in which each node is annotated with quantitative probability information. The full specification is as follows:

- (a) Each node corresponds to a random variable, which may be discrete or continuous.

- (b) A set of directed links or arrows connects pairs of nodes. If there is an arrow from node  $X$  to node  $Y$ ,  $X$  is said to be a parent of  $Y$ . The graph has no directed cycles and hence is a directed acyclic graph, or DAG.
- (c) Each node  $X_i$  has a conditional probability distribution  $P(X_i | \text{Parents}(X_i))$  that quantifies the effect of the parents on the node.

Bayesian networks are useful because they provide a compact representation of a full joint distribution, given from the combination of the topology (conditional independence) and the conditional distributions. In the simplest case, conditional distribution represented as a conditional probability table (CPT) giving the distribution over  $X_i$  for each combination of values for the parents.

In Bayesian networks we can do inference: computing the probability distribution of a set of query variables, given some observed evidence variables. We can also do Inference by stochastic simulation:

- Draw  $N$  samples from a sampling distribution  $S$ .
- Compute an approximate posterior probability  $\hat{P}$ .
- Show this converges to the true probability  $P$ .

Given the intractability of exact inference in large, multiply connected networks, it is essential to consider approximate inference methods.

- Sampling from an empty network
- Rejection sampling: reject samples disagreeing with evidence
- Likelihood weighting: weight samples by likelihood of evidence
- Markov chain Monte Carlo (MCMC): sample from a stochastic process whose stationary distribution is the true posterior

The complexity of exact inference in Bayesian networks depends strongly on the structure of the network.

- In single connected networks (or polytrees), any two nodes are connected by at most one (undirected) path, so the complexity of inference is linear in the size of the network ( $O(d^k n)$ ).

- In multiply connected networks, we can reduce 3SAT to exact inference to NP-Hard that is equivalent to counting 3SAT solutions, thus #P-complete.

## 10 Machine Learning

**ML.1** Introduce the main paradigms of machine learning, describing in particular the fundamental ingredients of the supervised paradigm, and how the complexity of an hypothesis space can be measured in a useful way in the case of a binary classification task.

### Answer:

Machine learning is the study of computer algorithms that is able to learn from data. A learning algorithm must have the following components:

- **Tasks:** how the machine learning algorithm should process an example.
- **Performance measure:** how accurate is the function/model returned by the learning algorithm.
- **Experience:** Dataset.

There are different paradigms of machine learning:

- **Supervised learning:** given pre-classified examples (training set),  $Tr = \{(x^{(i)}, f(x^{(i)}))\}$ , learn a general description  $h(x)$  (hypothesis) which captures the information content of the examples. Then given a new example  $\tilde{x}$ , we can predict the corresponding output  $h(\tilde{h})$ . It's called supervised because we assume that an expert provides the value of  $h$  for the corresponding training instance  $x$ .
- **Unsupervised learning:** given a set of examples  $Tr = \{x^{(i)}\}$ , discover regularities and/or patterns in the data. In this case there is no expert to provide the correct answer.
- **Reinforcement learning:** the agent learns by interacting with the environment. The agent receives a reward that can be positive, negative or neutral for each action and the goal is to maximize the total reward.

The fundamental ingredients of the supervised paradigm are:

- **Training data:** data that are drawn from the Instance Space,  $X$ .
- **Hypothesis space:** it is the set of functions that the learning algorithm can choose the function  $h$  that approximate the function  $f$  (Function to be learned).
- **Learning algorithm:** search algorithm into the hypothesis space.

$H \neq$  set of possible functions, search into  $H$  to be exhaustive  $\rightarrow$  overfitting: the algorithm learns the training data too well, so it doesn't generalize well to new examples.

There is inductive bias on  $H$  and search algorithm: set of assumptions that the learning algorithm uses to predict outputs of new instances.

The complexity of an hypothesis space can be measured in a useful way by the **VC dimension**:

- **Definition:** the VC dimension of a hypothesis space  $H$  is the size of the largest set of points that can be shattered by  $H$ .
- **Shattering:** a set of points  $S$  is shattered by  $H$  if for every possible labeling of the points in  $S$ , there exists a function  $h$  in  $H$  that correctly classifies the points in  $S$ .

In the case of binary classification task, we have only two possible labels. If we work with linear classification, we can classify the points in the plane with the sign of the position with respect to a line (positive or negative). The VC dimension of the hypothesis space of linear classifiers is 3, because we can shatter 3 points, but not 4: it is not possible to find a line that separates 4 points in all the possible ways, there always exist two couples of points such that if we connect the two members by a segment, the two resulting segments will intersect so a curve is needed.

**ML.2** Explain in detail the supervised learning paradigm, describe the role of the training set, the validation set and the test set (how to use data in our hands). Give the definition of true error and empirical error, highlighting the role of them during the learning process.

**Answer:**

Supervised Learning ref to answer ??.

On learning tasks we have a set of data, this can be splitted into:

- **Training set:** used to train the model.
- **Validation set:** it is a subset of the training set used to tune the hyperparameters of the model (hold-out, cross-validation).
- **Test set:** used to evaluate the selected model.

Model selection is the process of choosing the best model for a given task by selecting the best hyperparameters:

- **Hold-out procedure:** split the training set into two parts, the first one is used to train the model and the second one (validation set) is used to test the trained model with different hyperparameters.
- **Cross-validation:** K-different classifier/regressors are trained on K-different subsets of  $Tr$  ( $Va_1, \dots, Va_k$ ) and then is iteratively applied the Hold-out procedure on the k-pairs ( $Tr_i = Tr - Va_i, Va_i$ ).

The empirical error ( $error_{Tr}(h)$ ) of hypothesis  $h$  with respect to  $Tr$  is the number of examples that  $h$  misclassifies:

$$error_{Tr}(h) = \#\{(x, f(x)) \in Tr | f(x) \neq h(x)\} / |Tr| \quad (17)$$

The true error ( $error_D(h)$ ) of hypothesis  $h$  with respect to target concept  $c$  and distribution  $D$  is the probability that  $h$  will misclassify an instance drawn at random according to  $D$ :

$$error_D(h) \equiv Pr_{x \in \mathcal{D}}[c(x) \neq h(x)] \quad (18)$$

We can say that  $h \equiv \mathcal{H}$  overfits  $Tr$  if  $\exists h' \in \mathcal{H}$  such that  $error_{Tr}(h) < error_{Tr}(h')$  and  $error_D(h) > error_D(h')$ .

The goal of machine learning is to solve a task with the lowest possible true error, but a classifier learn on training data so it generate empirical

error and not true error. It's possible to have bound on the true error from the empirical error with probability  $1 - \delta$ :

$$error_D(h_w^*) \leq \underbrace{error_{Tr}(h_w^*)}_A + \underbrace{\epsilon(n, VC(\mathcal{H}), \delta)}_B \quad (19)$$

B (VC-confidence) depends on the ratio between  $VC(\mathcal{H})$  and  $n$  (number of training examples) and on  $1 - \delta$  (confidence level).

Problem: as the VC-dimension grows, the empirical risk (A) decreases, however the VC confidence (B) increases! For minimizing the right hand of the confidence bound we can use the principle **Structural Risk Minimization**: we get a tradeoff between A and B, we want to select the hypothesis with the lowest bound on the true risk.

**ML.3** In the context of machine learning, explain the fundamental ingredients of perceptron. Provide a brief introduction of how this model can be extended by creating a multi-layer architecture.

**Answer:**

A perceptron given an input vector  $\vec{x}$  and a weight vector  $\vec{w}$  calculates  $f(\sum w_i * x_i)$  that is the activation function of the perceptron. A Neural Network is a system consisting of interconnected units that compute nonlinear functions.

In a Neural Network we can find:

- **Input Units:** represent input variables.
- **Output Units:** represent output variables.
- **Hidden Units:** represent internal variables that codify correlations between input and output variables.
- **Weights:** are associated to connections between units.

Having decided on the mathematical model for individual “neurons,” the next task is to connect them together to form a network. **Feed-forward networks:** the information flows in one direction, from the input units, through the hidden units (if any) and to the output units. Gradient descent for feed-forward: we need to optimize the weights of the network to minimize the error on the training set → backpropagation algorithm. We define the function error with  $c$  output units:

- (a) is defined a loss function which measure the error of the network on the training set.
- (b) the weights are randomly initialized.
- (c) it's executed the forward pass: the input is propagated through the network and the output is computed.
- (d) is calculated the gradient descent with respect to the weights.
- (e) the weights are updated in the opposite direction of the gradient multiplied by a learning rate.
- (f) steps c and e are repeated until a stop condition is reached (for example the error is below a threshold or the number of epochs is reached).

## 11 Reinforcement Learning

**RL.1** Introduce the Reinforcement Learning paradigm. Moreover, present the Q-Learning algorithm explaining its theoretical basis.

**Answer:**

Reinforcement Learning is a paradigm in which the agent learns by interacting with the environment. The agent receives a reward that can be positive, negative or neutral for each action and the goal is to learn to choose actions that maximize the total reward.

$$r_0 + \gamma r_1 + \gamma^2 r_2 + \dots \quad (20)$$

Where:

- $r_i$  is the reward received after the action at  $a_i$  the state  $s_i$ .
- $\gamma$  is the discount factor, it is a value between 0 and 1 that represents the importance of future rewards.

We've a finite set of states  $S$  and a finite set of actions  $A$ ; at each discrete time agent observes state  $s_t \in S$  and selects action  $a_t \in A$ . Then the agent receives a reward  $r_t$  and the environment moves to a new state  $s_{t+1}$ . All this should work under the Markov assumption:  $s_{t+1} = \delta(s_t, a_t)$  and  $r_t = r(s_t, a_t)$ .



- $r_t$  and  $s_{t+1}$  depend only on current state and action.
- functions  $\delta$  and  $r$  may be nondeterministic and not necessarily known to the agent.

The agent's goal is to execute actions in environment, observe results, and: learn action policy  $\pi() : S \rightarrow A$  that maximizes  $E[r_t + \gamma r_{t+1} + \gamma^2 r_{t+2} + \dots]$  from any starting state in  $S$ . (training examples of form  $((s, a)r)$ ).

Considering deterministic environments, for each possible policy  $\pi$  we can define an evaluation function over states:

$$V^\pi(s) \equiv \sum_{i=0}^{\infty} \gamma^i r_{t+i} \quad (21)$$

Here the task is to learn the optimal policy  $\pi^*$  that maximizes  $V^{\pi^*}(s)$  for all  $s \in S$ .

We want to have an agent that learn the evaluation function  $V^{\pi^*}$ , but for doing this the agent need to know the transition function  $\delta$  and the reward function  $r$ .

In the case that the agent doesn't know the transition function  $\delta$  and the reward function  $r$ , we can define the Q-function similar to  $V^*$ :

$$Q^*(s, a) \equiv r(s, a) + \gamma V^*(\delta(s, a)) \quad (22)$$

if agent learns Q, it can choose optimal action even without knowing  $\delta$ .

$$\pi^*(s) = \arg \max_a Q(s, a) \quad (23)$$

Thus, Q is the evaluation function the agent will learn. Note that Q and  $V^*$  are related by:

$$V^*(s) = \max_{a'} Q(s, a') \quad (24)$$

which allows us to write Q recursively as:

$$\begin{aligned} Q^*(s, a) &= r(s_t, a_t) + \gamma V^*(\delta(s_t, a_t)) \\ &= r(s_t, a_t) + \gamma \max_{a'} Q(s_{t+1}, a') \end{aligned} \quad (25)$$

so we can define a training rule to learn  $Q$ :

$$\hat{Q}(s, a) \leftarrow r + \gamma \max_{a'} \hat{Q}(s', a') \quad (26)$$

The Q-learning algorithm is:

- (a) Initialize table  $\hat{Q}(s, a) \leftarrow 0$ .
- (b) Observe current state  $s$ .
- (c) do forever:
  - i. select an action  $a$  and execute it.(it could be randomly selected or using  $\arg \max_a \hat{Q}(s, a)$ )
  - ii. receive immediate reward  $r$ .
  - iii. observe new state  $s'$ .
  - iv. update the table entry for  $\hat{Q}(s, a)$ :

$$\hat{Q}(s, a) \leftarrow r + \gamma \max_{a'} \hat{Q}(s', a') \quad (27)$$

- v.  $s \leftarrow s'$

We can show that  $\hat{Q}$  converges to  $Q$ .

For non-deterministic environments, we redfine  $V$  and  $Q$  by taking expected values

$$V^\pi(s) \equiv E\left[\sum_{i=0}^{\infty} \gamma^i r_{t+1}\right] \quad (28)$$

$$Q(s, a) \equiv E[r(s, a) + \gamma V^*(\delta(s, a))] \quad (29)$$

and the training rule becomes:

$$\hat{Q}_n(s, a) \leftarrow (1 - \alpha_n) \hat{Q}_{n-1}(s, a) + \alpha_n (r + \gamma \max_{a'} \hat{Q}_{n-1}(s', a')) \quad (30)$$

## 12 Constraint satisfaction problems

1. Give the formal definition for a constrain satisfaction system, and discuss the approaches presented in class on how to find solutions.

**Answer:**

A costraint satisfaction systems used to solve constraint satisfaction problems. It is defined by:

- **state**: a set of variables  $X_i$  with values from domains  $D_i$ .
- **goal test**: a function that determines whether a given state is a goal state.

It can be helpful to visualize a CSP as a constraint graph, where each node represents a variable and each arc represents a constraint (in binary CSP at most two vars for each constraint). A solution to a CSP is an assignment of values to all variables such that all constraints are satisfied. A constraint satisfaction problem can have **discrete** or **continuous** variables.

- **Discrete variables**: finite domains and infinite domains (integers, strings, etc). For infinite domains it's needed a constraint language that allows for compact representation of constraints.
- **Continuous variables**: common in real world problems, the most known is the **linear programming** (LP) problem.

The constraints could be:

- **Unary**: involve a single variable.
- **Binary**: involve pairs of variables.
- **Higher-order**: involve 3 or more variables.
- **Preferences**: soft constraints, they are not required to be satisfied  $\rightarrow$  constrained optimization problem.

The first approach to find solutions is **Standard search formulation**  $\rightarrow$  **min-conflicts**:

- **Initial state**: the empty assignment,
- **Successor function**: assign a value to an unassigned variable that does not conflict with current assignment.
- **Goal test**: the current assignment is complete and satisfies all constraints.

It use deep-first search.

There is an important property not considered by the previous approach: **commutativity**. The commutativity property states that the order in

which we assign values to variables does not matter, the result is the same.

The term backtracking search is used for a depth-first search that chooses values for one variable at a time and backtracks when a variable has no legal values left to assign.

It repeatedly chooses an unassigned variable, and then tries all values in the domain of that variable in turn, trying to find a solution. If an inconsistency is detected, then BACKTRACK returns failure, causing the previous call to try another value. We can improve the backtracking search with general-purpose methods:

- **Variable ordering:** choose the next variable to be assigned a value. (most constrained variable, most constraining variable, least constraining).
- **Variable assignment:** forward checking: keep track of remaining legal values for unassigned variables and terminate search when any variable has no legal values.
- **Detecting failure:** constraint propagation (arc and node consistency).
- **Problem structure:** the problem structure can give some advantages in the search:
  - **independent subproblems:** independence can be ascertained by finding the connected components of the constraint graph. Each component can be solved independently of the others in  $O(d^c n/c)$  time, otherwise  $O(d^n)$ .
  - **Tree-structured CSPs:** graph is a tree when any two variables are connected by only one path and can be solved in linear time.  $O(nd^2)$  Algorithms for tree-structured CSPs:
    - (a) Choose a variable as root, order the variables from root to leaves such that every node's parent precedes it in the ordering.
    - (b) For  $i$  from  $n$  down to 2, impose arc consistency (A variable  $X$  is arc-consistent with respect to another variable  $Y$  if, for every value  $x$  in the domain of  $X$ , at least one value exists  $y$  in the domain of  $Y$  such that the pair  $(x,y)$

respects all constraints between  $X$  e  $Y$ .) on the arc from  $X_i$  to its parent.

- (c) For  $i$  from 1 up to  $n$ , assign  $X_i$  consistently with all its ancestors.

The last approach is based on the use of Iterative algorithms that repeatedly improve the quality of the current assignment. (ex: hill climbing, simulated annealing, genetic algorithms). To apply to CSPs:

- allow states with unsatisfied constraints
- operators reassign variable values

Variable selection: randomly select any conflicted variable Value selection by min-conflicts heuristic:

- choose value that violates the fewest constraints i.e., hillclimb with  $h(n)$  = total number of violated constraints

## 13 NLP

NLP (Natural Language Processing) is the field of AI that is concerned with the processing and understanding of natural language, acquiring knowledge and communicating with humans.

**NLP.1** In the field of NLP, explain the difference between the n-gram (in particular unigram and bigram) and bag-of-words models. Show how the n-gram model can be used for classification and information retrieval problems, detailing the relations and possible differences with models such as bag-of-words.

**Answer:**

For NLP related tasks we need language models that predict the probability distribution of linguistic expressions.

**Definition:** a language model is a probability distribution over sequences of words.

One of this models is the **n-gram** model, it is defined as a probability distribution on sequences of characters.

$$P(c_{1:N}) = \text{the probability of a sequence of } N \text{ characters } c_1, \dots, c_N \quad (31)$$

A sequence of written symbols of length  $N$  is called an  $N$ -gram. (ex: unigram, bigram, trigram, etc)

*a pattern  $n$ -gram is a Markov chain of order  $n-1 \rightarrow$  the probability depends only on the previous  $n-1$  symbols*

For trigram:

$$P(c_{1:N}) = \prod_{i=1}^N P(c_i | c_{i-2:i-1}) \quad (32)$$

The main problem on  $n$ -gram model is that the train corpus provide only an estimate of the true probability distribution, for example it could lead to zero probability for unseen  $n$ -grams. There are two main solution to this problem:

- **Laplace Smoothing:** assign non-zero probability to unseen  $n$ -grams. If a random boolean variable  $X$  is false in all  $n$  observations, then  $P(X = \text{true}) = \frac{1}{n+2} \rightarrow$  one test could be true and one false.
- **Backoff:** in which we start by estimating  $n$ -gram counts, but for any particular sequence that has a low (or zero) count, we back off to  $(n-1)$ -grams

We can evaluate a model with cross-validation but we would like to define a task independent quality model.

$$\text{Perplexity}(c_{1:N}) = P(c_{1:N})^{-\frac{1}{N}} \quad (33)$$

Low perplexity means that the model is good at predicting the test set. Perplexity is the inverse probability of the test set, normalized by the number of words.

$N$ -gram models can be also used for words, with the difference that the vocabulary is significantly larger. The objective is to calculate the probability of a sentence  $P(w_1, \dots, w_n)$ . Chain rule can be used to calculate the joint probability of words:

$$P(w_1, \dots, w_n) = \prod_{i=1}^n P(w_i | w_1, \dots, w_{i-1}) \quad (34)$$

We will never see enough data to estimate all these probabilities accurately, so we need to make some approximation. By markov assumption we can approximate the probability of a word with the probability of the previous word:

$$P(w_1, \dots, w_n) \approx \prod_{i=1}^n P(w_i | w_{i-k:i-1}) \quad (35)$$

**Bag-of-words** (also known as vector model) model: it is a simplifying representation used in natural language processing. Represents the frequency of occurrence of each word in a document.

Difference between n-gram and bag-of-words:

- Feature vector are large and scattered in BoW.
- BoWs and unigram return the same result.
- The order of words is lost in BoW, while it is preserved a local notion of order in higher order n-grams.
- in n-gram computational complexity increases with n.

A simple representation of a document is one-hot encoding, where each word of it present in the vocabulary is represented by one or zero in the vector (the word is present or not). The problem is that one-hot vectors are orthogonal, so we can't capture any similarity between words. Some possible solutions are:

- **Knowledge representation way:** rely on a list of synonyms or taxonomies of words to obtain similarity.
- **Machine learning way:** learn to code the similarity in the vector themselves. The idea is to construct a vector space where the distance between vectors is a measure of similarity between the words. Example of this approach is the **word2vec** model: it use context for predict a target word (CBOW) or use a word for predict the context (skip-gram).

N-gram models can be used for classification problems. Given a text of some kind decide which predefined group of classes it belongs to. (ex: spam detection) There are two main approaches:

- **rules-based:** define a set of hard-coded rules that classify the text. We define the rules based on a combination of words (or other features)

**Problem:** rule-based can lead to high accuracy result, but building such rules is very expensive and not always possible.

- **linguistic modeling and machine learning:** we define a n-gram pattern for each class and we calculate the probability of the text to belong to each class with Naive Bayes (or other classifier).