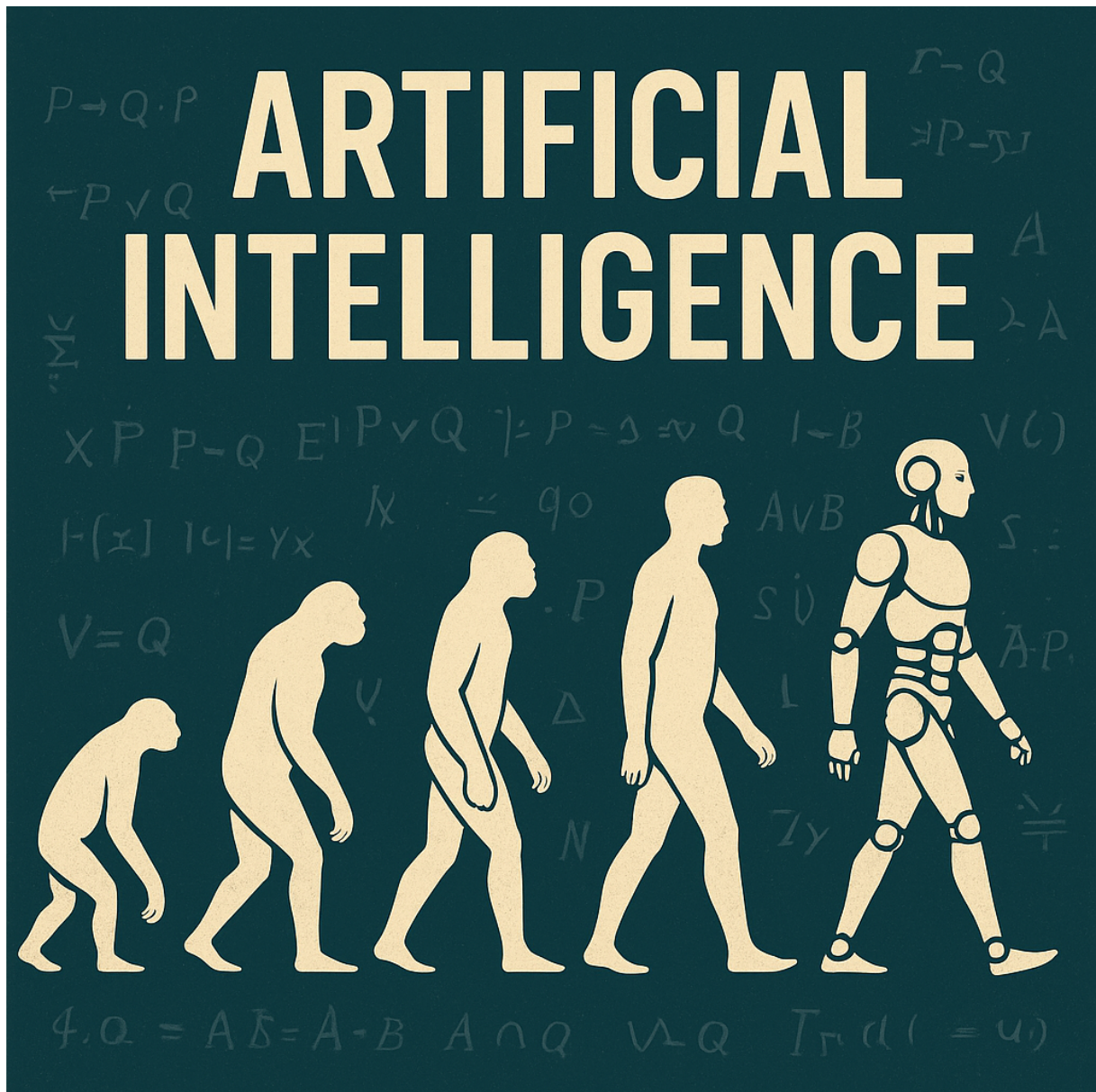


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Master's degree in Artificial Intelligence and Robotics

This document summarizes and presents the topics for the Artificial intelligence course for the Master's degree in Artificial Intelligence and Robotics at Sapienza University of Rome. The document is free for any use. If the reader notices any typos, they are kindly requested to report them to the author.



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# CHAPTER

## 1

# INTRODUZIONE

## 1.1 Basic Definitions

In the context of the artificial intelligence, an **agent** is an entity that can

- Perceive the environment through *sensors* (percepts)
- Act upon the environment through *actuators* (actions).

We say that an agent is **rational** if he selects the action that maximize a given *performance measure*, informally, he attempts to do "the right thing". The best case is hypothetical and often unattainable, because the agent usually can't perform all the actions needed, and can't perceive all the information about the environment.

An agent has a performance measure  $M$  and a set  $A$  of all possible actions, given percept a sequence  $P$  and knowledge  $K$  (data), he has to select the next action  $a \in A$ , is a map

$$M \times P \times K \longrightarrow A. \quad (1.1)$$

An action  $a$  is optimal if it maximize the expected value of  $M$ , given the sequence  $P$  and the knowledge  $K$ . An agent is rational if he always chooses the optimal action. More specifically, an agent consists in two components:

- an architecture which provides an interface to the environment
- a program executed on that architecture.

There are some limitation that we aren't considering, such as the fact that determining the optimal choice could take too much time or memory on the architecture.

### 1.1.1 Types of Agents

There are different kinds of agents, a **Table Driven Agent** is the simplest form of agent architecture. It's essentially a look-up table that maps every possible sequence of percepts (what the agent has sensed so far) to a corresponding action the agent should take. His behavior can be resumed in the algorithm 1.

A **Reflex Agent** consists in three components:

- sensors to get information from the environment

**Algorithm 1** Table Driven Agent**Require:** *percepts***persistent:** *percepts*, a sequence, initially empty

table, a table of actions, indexed by percept sequences, initially fully specified

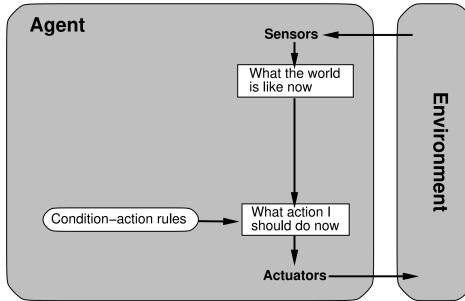
append *percept* to the end of *percepts**action*  $\leftarrow$  LookUp(*percepts*, *table*)**return** *action*

Figure 1.1: Reflex agent diagram

- a decision making process, in form of a *condition-action rules*, typically looks like IF (condition) THEN (action).
- actuators, the outputs that allow the agent to affect or change the environment.

A **Model-Based Reflex Agent** is an enhanced version of the previous one, the key enhancement here is the inclusion of an *Internal State* and a *Model of the World* to make up for the agent's limited view of the environment. The internal state cannot simply be the last thing the agent saw; it needs to be updated to reflect reality. This is done using a Model of the World, which contains two key pieces of knowledge:

- How the world evolves independently of the agent, his accounts for changes in the environment that occur regardless of the agent's actions (e.g., a clock ticking, an external event).
- How the agent's own actions affect the world, this is the effect of the agent's previous action (e.g., if the agent drove forward, its position changed).

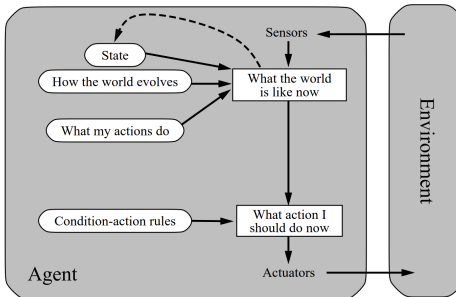


Figure 1.2: Model Based Reflex agent diagram

If a model based reflex agent consider the future prospective, is a **Goal Based Agent**, as shown in figure 1.3.

A **Utility Based Agent** is equipped with a *utility function* that maps a state to a number which represents how desirable the state is. Agent's utility function is an internalization of the performance function.

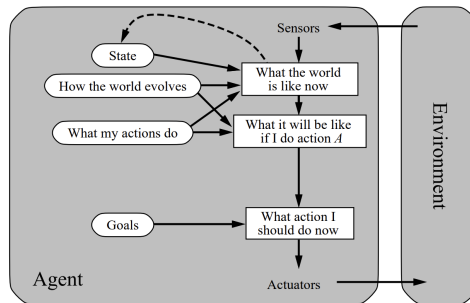


Figure 1.3: Goal Based agent diagram

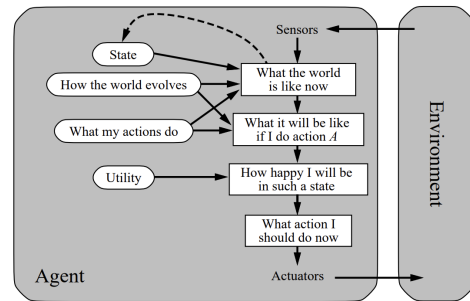


Figure 1.4: Utility Based agent diagram

A **Learning Agent** is an architecture designed to improve its efficiency over time by separating four functions:

- the performance element selects actions
- the critic provides feedback on those actions against a standard
- the learning element uses this feedback to update the agent's internal knowledge
- the problem generator suggests exploratory actions to gain new knowledge. This structure enables the agent to continuously adapt and improve its decision-making.

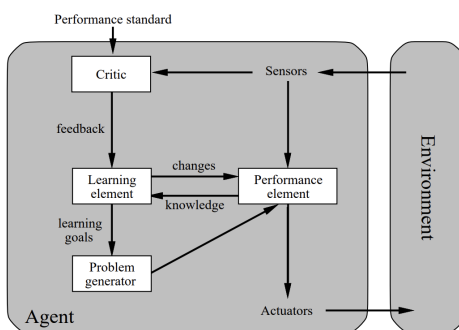


Figure 1.5: Learning agent diagram

An agent can be classified in one of the following groups:

- a **domain specific agent** is a solver specific to a particular problem (such as playing chess), is usually more efficient.
- a **general agent** is a solver for general problems, such as learning the rule of any board game, is usually more intelligent but less efficient.



### 1.1.2 The Environment

An environment can be classified in terms of different attributes:

- An environment can be **fully observable** if all the relevant information are accessible to the sensors, otherwise is **partially observable**.
- If there are no uncertainty, the environment is **deterministic**. An environment is **stochastic** if uncertainty is quantified by using probabilities, otherwise is **non deterministic** if uncertainty is managed as actions with multiple outcomes.
- An environment is **episodic** if the correctness of an action can be evaluated instantly, otherwise if are evaluated in the future developments, is **sequential**.
- An environment can be **static** or **dynamic**, if it does not change, but the agent's performance score changes, the environment is called **semi-dynamic**.
- An environment can be perceived as **discrete** or **continuous**.
- In a single environment there may be multiple agent, that can be **competitive** or **cooperative**.

Many sub-areas of AI can be classified by:

- Domain-specific vs. general.
- The environment.
- Particular agent architectures sometimes also play a role, especially in Robotics.

It follows a classification of some areas in terms of the attributes we discussed:

- **Classical Search**, the environment is

- fully observable
- deterministic
- static
- sequential
- discrete
- single-agent

and the approach is *domain specific*.

- **Planning**, the environment is

- fully observable
- deterministic
- static
- sequential
- discrete
- single-agent

and the approach is *general*.

- **Adversarial Search**, the environment is

- fully observable
- deterministic
- static
- sequential
- discrete
- multi-agent



and the approach is *domain specific*.

- **General Game Playing**, the environment is

- fully observable
- deterministic
- static
- sequential
- discrete
- multi-agent

and the approach is *general*.

- **Constraint Satisfaction & Reasoning**, the environment is

- fully observable
- deterministic
- static
- episodic
- discrete
- single-agent

and the approach is *general*.

- **Probabilistic Reasoning**, the environment is

- partially observable
- stochastic
- static
- episodic
- discrete
- single-agent

and the approach is *general*.

## CHAPTER

# 2

## SEARCH PROBLEMS

### 2.1 Classical Search

Let's consider two basic example of classical search problems, the first one is the following:



Starting from Zurigo, we would like to find a route to Zagabria. We have an initial state (Zurigo), and we have to apply actions (drive) to reach the goal state (Zagabria). Another example is the following, we want to solve the tiles-puzzle game, shown in figure 2.1, to reach the left state, starting from the right one, the actions to perform is the move of the tiles. A performance measure could be to minimize the summed-up action costs.

The diagram illustrates a 4x4 tile puzzle transformation. On the left, a 4x4 grid contains the following values:

9	2	12	6
5	7	14	13
3	4	1	11
15	10	8	Black

A horizontal arrow points to the right, indicating a transformation. On the right, the same 4x4 grid is shown with different values, representing the goal state:

1	2	3	4
5	6	7	8
9	10	11	12
13	14	15	Black

Figure 2.1: The tile game

In the classical search context, we restrict the agent's environment to a very simple setting, with a finite number of states and actions, a single agent, a fully observable state environment that doesn't evolve, given that assumption, the classical search problems are the simplest one, despite that, are very important problems in practice.

Every problem specifies a state space.

**Definition 1** A **State Space** is a 6-tuple  $\Theta = (S, A, c, T, I, S^G)$  where:

- $S$  is a finite set of the states.
- $A$  is a finite set of actions.
- $c : A \rightarrow \mathbb{R}^+$  is the cost function.
- $T \subseteq S \times A \times S$  is the transition relation, that describes how an action on a given state make the agent evolve to the next state. We assume that the problem is deterministic, so for all  $s \in S$ ,  $a \in A$ , if  $(s, a, s') \in T$  and  $(s, a, s'') \in T$  then  $s' = s''$ .
- $I \in S$  is the initial state
- $S^G \subseteq S$  is the set of the goal states, where we want to end.

A transition  $(s, a, s')$  can be denoted  $s \xrightarrow{a} s'$ , we say that  $s \rightarrow s'$  if  $\exists a$  such that  $(s, a, s') \in T$ . We say that  $\Theta$  has **unit costs** if  $\forall a \in A, c(a) = 1$ . A state space can be illustrated as a directed labeled graph.

**Definition 2** Let  $\Theta = (S, A, c, T, I, S^G)$  to be a state space, we say that

- $s'$  is a **successor** of  $s$  if  $s \rightarrow s'$
- $s'$  is a **predecessor** of  $s$  if  $s' \rightarrow s$
- we say that  $s'$  is **reachable from**  $s$  if

$$\exists(a_1 \dots, a_n) \subseteq A \quad (2.1)$$

$$\exists(s_2 \dots, s_{n-1}) \subseteq S \quad (2.2)$$

$$(s, a_1, s_2) \in T \quad (2.3)$$

$$(s_2, a_2, s_3) \in T \quad (2.4)$$

$$\vdots \quad (2.5)$$

$$(s_{n-1}, a_n, s') \in T \quad (2.6)$$

we can write the sequence as follows

$$s \xrightarrow{a_1} s_2, \dots, s_{n-1} \xrightarrow{a_n} s'. \quad (2.7)$$

- We say that  $s$  is **reachable** (without reference state) if is reachable from  $I$ .
- $s$  is **solvable** if there exists  $s' \in S^G$  such that  $s'$  is reachable from  $s$ , otherwise  $s$  is **dead end**.

**Definition 3** Let  $\Theta = (S, A, c, T, I, S^G)$  to be a state space, and let  $s \in S$ . A **solution** for  $s$  is a path from  $s$  to some goal state  $s' \in S^G$ . The solution is **optimal** if it's cost is minimal, let  $H$  to be the set of all possible solution (sequence of states) for  $s$

$$H = \{\text{paths from } s \text{ to } s' \in S^G\} = \{(s_{i0}, s_{i1}, s_{i2}, \dots, s_{in}) : s_{in} \in S^G, s_{i0} = s\} \quad (2.8)$$

where  $n^i$  is the length of the  $i$ -th solution. The optimal solution is

$$\arg \min_{(s, s_{i1}, \dots, s_{in}) \in H} \sum_{j=0}^{n^i} c(s_{ij}). \quad (2.9)$$

A solution for  $I$  is called **solution for**  $\Theta$ , if such that solution exists,  $\Theta$  is **solvable**.

### 2.1.1 Vacuum Cleaner Example

Let's consider a vacuum cleaner, that is the agent of our problem, the goal is to clean a room, the vacuum cleaner can be in two possible points (left and right), this points can be clean or dirty. The agent can perform the following actions

- move right
- move left
- suck the dust on the floor

there are 8 possible states

- left point clean, right point clean, vacuum cleaner is on right point
- left point dirty, right point clean, vacuum cleaner is on right point
- left point clean, right point dirty, vacuum cleaner is on right point
- left point dirty, right point dirty, vacuum cleaner is on right point
- left point clean, right point clean, vacuum cleaner is on left point
- left point dirty, right point clean, vacuum cleaner is on left point
- left point clean, right point dirty, vacuum cleaner is on left point
- left point dirty, right point dirty, vacuum cleaner is on left point

the initial state is the one with the left point dirty, right point dirty, and the vacuum cleaner on the left point, we denote the actions  $R$  (move right),  $L$  (move left),  $S$  suck. The state space of the problem is show in figure 2.2.

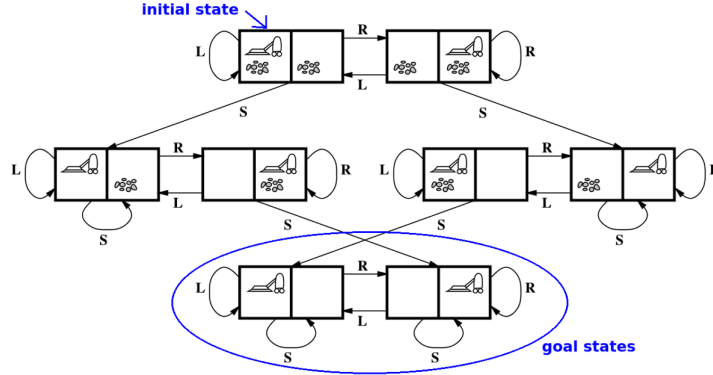


Figure 2.2: The vacuum cleaner state space

Some example of set of actions that can lead from the initial state to a goal state are

$$\begin{aligned} S &\rightarrow R \rightarrow S \\ S &\rightarrow R \rightarrow R \rightarrow S \\ R &\rightarrow S \rightarrow L \rightarrow S \end{aligned}$$

Typically, the state space is exponentially large in the size of its specification, search problems are typically computationally hard and/or NP-complete. We say that we can give an *explicit description* of a search problem if we can define his state space as a graph.

## 2.2 Problem Descriptions

**Definition 4** We have a **black box description** of the problem if we can't describe the state space explicitly but we can

- Know which is the initial state
- Check if a given state is a goal state
- Check the cost of a given action  $a$
- Given a state  $s$ , check all the actions that are applicable to state  $s$
- Given a state  $s$  and an applicable action  $a$ , we can get the successor state.

We can think about it in a programming-way, given a problem described by  $\Theta = (S, A, c, T, I, S^G)$ , we can't check directly  $\Theta$ , but we have an *API* of the problem that provide the following functions

- `InitialState()` : return the initial state of the problem
- `GoalTest( $s$ )` : return true if and only if  $s \in S^G$
- `Cost( $a$  : return  $c(a)$ )`
- `Actions( $s$ )` : return the set  $\{a : \exists s \xrightarrow{a} s' \in T \text{ for some } s' \in S\}$
- `ChildState( $s, a$ )` : return  $s'$  if  $s \xrightarrow{a} s' \in T$ .

We **specify** a search problem if we can program/access to such an *API*. There are a declarative description too.

### Definition 5

We have a **declarative description** of the problem if is described by the following sets:

- $P$  is a set of boolean variables (*propositions*)
- $I \subseteq P$  is the subset of  $P$  indicating which propositions are true in the initial states.
- $G \subseteq P$  is the subset of  $P$  describing the goal states in the following way
  - a state  $s$  is a set of propositions
  - $s \subseteq G \iff s$  is a goal state
- $A$  is a set of actions, each action  $a$  is described by
  - a set  $pre_a \subseteq P$  of *precondition*,  $a$  can be performed if and only if the conditions in  $pre_a$  are true.
  - a set of propositions  $add_a$
  - a set of propositions  $del_a$
  - the outcome of each action is the state  $(\{s\} \cup add_a) \setminus del_a$
  - $c : A \rightarrow \mathbb{R}$  is the cost function.

Declarative descriptions are strictly more powerful than black box ones. In this section we assume the black box description. In principle, the search strategies we will discuss can be used with any problem description that allows to implement the black box *API*.

### 2.2.1 Missionaries and Cannibals Example

The problem is the following

- there are a river and a boat that can carry the people from the left bank to the right
- there are 6 people, 3 missionaries and 3 cannibals
- the boat can carry 0, 1 or 2 people at the same time, not 3
- the goal is to get everybody to the left bank
- if at any time, there are more cannibals than missionaries in one bank, the missionaries get killed and the game is lost.

We can model the problem as follows

- the state space  $S$  is

$$S = \{(M, C, B) : M + C = 6, 0 \leq M \leq 3, 0 \leq C \leq 3, B \in \{0, 1\}\} \quad (2.10)$$

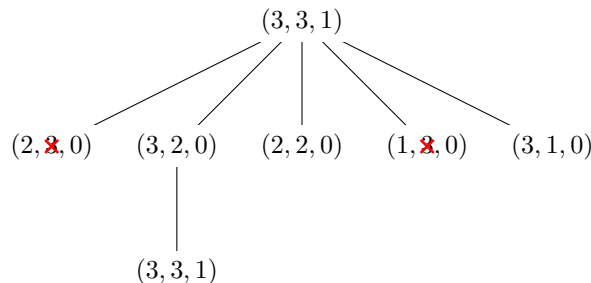
$M$  represents the current number of missionaries on the right bank,  $S$  represents the current number of cannibals on the right bank,  $B = 1$  if the boat is on the right bank, otherwise is on the left.

- the initial state is  $(3, 3, 1)$
- the goal states are  $S^G = (0, 0, 0), (0, 0, 1)$
- each actions have the same cost, is negligible
- the action that can be performed are the following
  - if  $B = 1$ , we can subtract (in total) 1 or 2 from  $M$  or  $C$  (or both), and set  $B = 0$ .
  - if  $B = 0$ , we can add (in total) 1 or 2 from  $M$  or  $C$  (or both), and set  $B = 1$ .

an action is applicable if and only if the following condition are satisfied after

- $M \geq C$  if  $M > 0$ , this decode the facts that the cannibals can't be more or equals than the missionaries on the right bank if there are missionaries on the left bank
- if  $M < 3$  (some missionaries are on the left bank) then  $(3 - M) > (3 - C)$  the missionaries on the left bank must be greater then the cannibals on the left bank.

To search a solution we can start from the initial state and expand the tree of possible solutions accordingly to the applicable actions.



### 2.2.2 Tree and Graph Search

In the search context the following terminology is used

- Search node  $n$ : Contains a state reached by the search, plus information about how it was reached.
- Path cost  $g(n)$ : The cost of the path reaching  $n$ .
- Optimal cost  $g^*$ : The cost of an optimal solution path. For a state  $s$ ,  $g^*(s)$  is the cost of a cheapest path reaching  $s$ .

- Node expansion: Generating all successors of a node, by applying all actions applicable to the node's state  $s$ . Afterwards, the state  $s$  itself is also said to be expanded.
- Search strategy: Method for deciding which node is expanded next.
- Open list: Set of all nodes that currently are candidates for expansion. Also called frontier.
- Closed list: Set of all states that were already expanded. Used only in graph search, not in tree search (up next). Also called explored set.

When we explore the state space of a problem we can maintain a closed list of all the node that has been already searched, to check for each generated new node if it is already in the list (if so, we discard it). If such list is used, the search is called **graph search**, else, if the same state may appear in many search nodes, is called **tree search**. The tree search doesn't use a list so require less memory.

When we analyze a search algorithm, we are interested in various properties

- **Completeness:** the algorithm is guaranteed to find a solution (if there are one).
- **Optimality:** the returned solution is guaranteed to be optimal.
- **Time Complexity:** How long does it take to find a solution? (Measured in generated states).
- **Space Complexity:** How much memory does the search require? (Measured in states).
- **Branching Factor:** The number  $b$  of how many successor a state may have.
- **Gal depth:** the number  $d$  of action required to reach the shallowest (nearest to the initial state) goal state.

## 2.3 Blind Search

We talk about *blind search* if the problem does not require any input beyond the problem API. Does not require any additional work from the programmer. For each node  $n$  in the search context, we define the following data structure:

- $n.State$  is the state which te node contains
- $n.Parent$  is node in the search tree that generated this node
- $n.Action$  is the action that was applied to the parent to generate the node
- $n.PathCost$ , also denoted  $g(n)$ , is the cost of the path from the initial state to the node (as indicated by the parent pointers).

On a node we can perform the following operations

- $\text{Solution}(n)$  returns the path from the initials state to  $n$
- $\text{ChildNode}(n, a)$  returns the node  $n$  corresponding to the application of action  $a$  in state  $n.State$ .

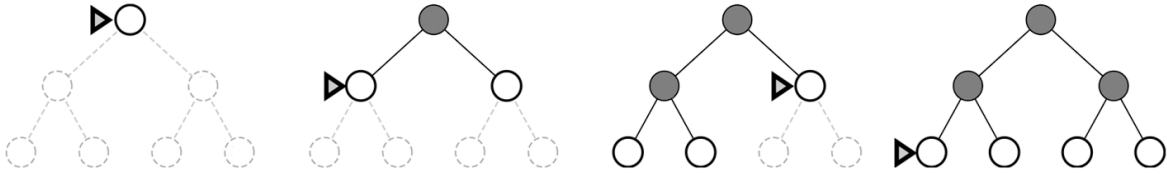
We also have the open list (called frontier) where we can perform the following actions:

- $\text{Empty}(\text{frontier})$  returns true if and only if there are no more elements in the open list.
- $\text{Pop}(\text{frontier})$  returns the first element of the open list, and removes that element from the list.
- $\text{Insert}(\text{element}, \text{frontier})$  inserts an element into the open list.

The insert function can put the element in front, in the last positions, or in other positions, it depends from the implementation (different implementations yield different search strategies).

### 2.3.1 Breadth-First Search

The strategy is to expand nodes in the order they were produced, as a FIFO queue, we expand the shallowest unexpanded node.



This algorithm is complete and optimal (in case of unit cost function). We are using the black box API.

---

**Algorithm 2** Breadth-First Search

---

```

Require: problem
node ← InitialState()
if GoalTest(node.State) then
    return Solution(node)
end if
frontier ← a FIFO queue with node in it
explored ← an empty set
while true do
    if Empty?(frontier) then
        return Failure
    end if
    node ← pop(frontier)
    add node.State in explored
    for each action in Actions(node.State) do
        child ← ChildNode(node, action)
        if child.State is not in explored or frontier then
            if GoalTest(child.State) then
                return Solution(child)
            end if
            frontier ← Insert(child, frontier)
        end if
    end for
end while

```

---

Let  $b$  to be the maximum branching factor and  $d$  the depth of the shallowest goal state, an upper bound (in the worst case) for the number of nodes generated (time complexity) is

$$b + b^2 + b^3 \dots + b^d \in O(b^d) \quad (2.11)$$

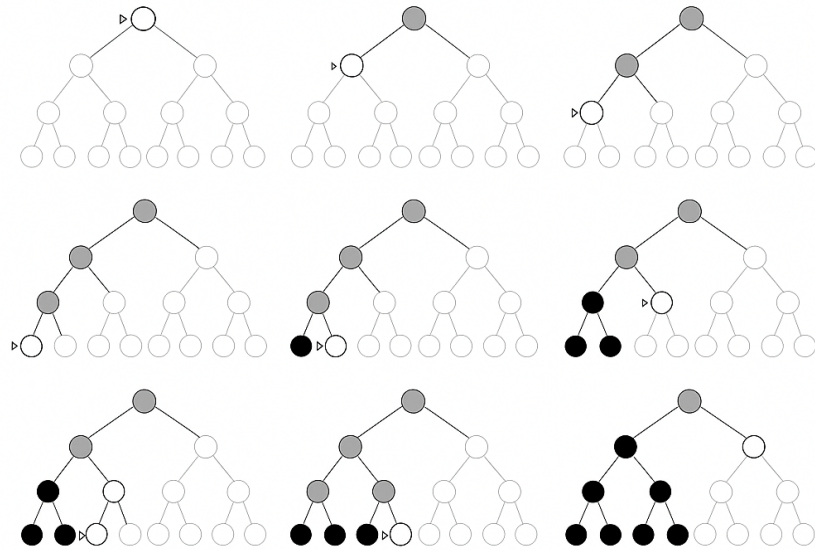
the same for the space complexity since all generated nodes are kept in memory. Let's see an example, assume that  $b = 10$ , the agent can generate  $10^4$  nodes per second, and each node has a size of 1 kilobyte, we have the following data:

Depth	Nodes	Time	Memory
2	110	0.11 milliseconds	107 kilobytes
4	$11,110$	11 milliseconds	10.6 megabytes
6	$10^6$	1.1 seconds	1 gigabyte
8	$10^8$	2 minutes	103 gigabytes
10	$10^{10}$	3 hours	10 terabytes
12	$10^{12}$	13 days	1 petabyte
14	$10^{14}$	3.5 years	99 petabytes

The critical resource for this method is the memory.

### 2.3.2 Depth-First Search

The strategy is to expand the most recent explored node, as a LIFO queue, we expand the deepest unexpanded node.



The algorithm is not complete since it may take infinite time, since there is no check for cycles along the branches. It's not optimal since he chooses a direction and looks for a path to a goal state. Is typically implemented as a recursive function, as in algorithm 4.

---

**Algorithm 3** Depth-First Search

---

```

Require: problem, a node n
if GoalTest(n.State) then
    return empty action sequence
end if
for each action in Actions(node.State) do
    n' ← ChildNode(node,action)
    result ← Depth-First Search(problem,n')
    if result ≠ failure then
        return action o result
    end if
end for
return failure

```

---

With *action* o *result* is denoted the concatenation of actions. About the space complexity, this methods stores only a single path of actions, from the root to a leaf node, since once a node has been expanded, it can be removed from memory as soon as all its descendants have been fully explored.

If *m* is the maximal depth reached, the space occupied is in  $O(bm)$ . About the time complexity, in the worst case the nodes generated is in  $O(b^m)$ .

### 2.3.3 Uniform-Cost Search

This methods is equivalent to the well known Dijkstra's algorithm. We expand the node with the lowest path cost  $g(n)$ , the frontier is ordered by the path cost, with the lowest first. It differs from the BFS since a test is added to check if a better path is found to a node currently on the frontier.

**Theorem 1** *The Uniform-Cost search algorithm is optimal, since the Dijkstra's algorithm is optimal, and Uniform-cost search is equivalent to Dijkstra's algorithm on the state space graph.*

The algorithm is complete if we assume that, the costs are strictly positive and the state space is finite. The time and space complexity are

$$O(b^{1+\lfloor g^*/\downarrow \rfloor}) \quad (2.12)$$

**Algorithm 4** Uniform-Cost Search

---

**Require:** *problem*

```

node  $\leftarrow$  InitialState()
frontier  $\leftarrow$  priority queue ordered by ascending g
explored  $\leftarrow$  empty set of states
while true do
    if Empty?(frontier) then
        return failure
    end if
    n  $\leftarrow$  Pop(frontier)
    if GoalTest(n.State) then
        return Solution(n)
    end if
    explored  $\leftarrow$  explored  $\cup$  n.State
    for each action in Actions(node.State) do
        n'  $\leftarrow$  ChildNode(n,a)
        if n'.State  $\notin$  explored  $\cup$  States(frontier) then
            frontier  $\leftarrow$  insert(n',frontier)
        else if  $\exists n'' \in frontier : n''.State = n'.State \wedge g(n') < g(n'')$ 
            replace n'' with n' in frontier
        end if
    end for
end while

```

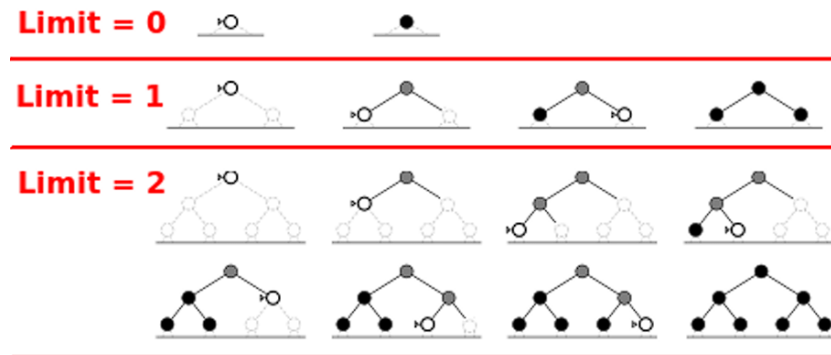
---

where

- $g^*$  is the cost of an optimal solution
- $\epsilon = \min c$  is the positive cost of the cheapest action, the minimum of the function *c*.

### 2.3.4 Iterative Deepening Search

This is an altered version of the Depth-First Search algorithm, where we define a predetermined depth limit, and apply the DFS in function of that limit, iteratively applying this by increasing the depth limit each time.



We split the algorithm in three different function.

**Algorithm 5** Iterative Deepening Search

---

**Require:** *problem*

```

for depth = 0, 1, ...,  $\infty$  do
    result  $\leftarrow$  Depth Limited Search(problem,depth)
    if result  $\neq$  cutoff then
        return result
    end if
end for

```

---

---

**Algorithm 6** Depth Limited Search

---

**Require:**  $problem, limit$   
 $node \leftarrow \text{InitialState}()$   
**return** Recursive DLS( $node, problem, limit$ )

---

The algorithm is complete since we are keep searching until a solution is found, is also optimal if all the cost are unitary, the space complexity is  $O(bd)$ . The time complexity is in  $O(b^d)$ , this methods combines the advantages of breadth-first and depth-first search. It is the preferred blind search method in large state spaces with unknown solution depth.

---

**Algorithm 7** Recursive DLS

---

**Require:**  $n, problem, limit$   
**if** GoalTest( $n.\text{State}$ ) **then**  
    **return** empty action sequence  
**end if**  
**if**  $limit == 0$  **then**  
    **return** cutoff  
**end if**  
 $cutoffOccurred \leftarrow \text{false}$   
**for** each  $action$  in Actions( $n.\text{State}$ ) **do**  
     $n' \leftarrow \text{ChildNode}(n, action)$   
    result  $\leftarrow$  Recursive DLS( $problem, n'$ )  
    **if** result == cutoff **then**  
         $cutoffOccurred \leftarrow \text{true}$   
    **else**  
        **if** result  $\neq$  failure **then**  
            **return**  $action \circ$  result  
            **end if**  
        **end if**  
    **end if**  
**end for**  
**if**  $cutoffOccurred$  **then**  
    **return** cutoff  
**end if**  
**return** failure

---

The following table is a summary of the methods that we considered in this section, confronting the time and space complexity.

Criterion	BFS	Uniform Cost	DFS	Depth Limited	Iterative Deepening
Completeness	Yes, if $a$ is finite	Yes, if $a$ is finite and action costs is positive	No	No	Yes, if $a$ is finite
Optimality	Yes if action costs are 1	Yes	No	No	Yes if action costs are 1
Time Complexity	$O(b^d)$	$O(b^{1+\lfloor g^*/\epsilon \rfloor})$	$O(b^m)$	$O(b^l)$	$O(b^d)$
Space Complexity	$O(b^d)$	$O(b^{1+\lfloor g^*/\epsilon \rfloor})$	$O(bm)$	$O(bl)$	$O(bd)$

where

- $b$ : finite branching factor
- $d$ : goal depth
- $m$ : maximum depth of the search tree

- $l$ : depth limit
- $g^*$  : optimal solution cost
- $\epsilon > 0$ : minimal action cost.

## 2.4 Informed Search

**Definition 6** Let  $\Pi$  to be a problem with  $S$  the set of states, a **heuristic**  $h$  is a function

$$h : S \rightarrow \mathbb{R}^+ \cup \{\infty, 0\}$$

such that, if  $s$  is a goal state, then  $h(s) = 0$ .

A heuristic function  $h^*$  is perfect if  $h^*(s)$  is exactly the cost of a cheapest path from  $s$  to a goal state, and  $h^*(s) = \infty$  if no such path exists.  $h^*$  is also called the *goal distance* of  $s$ .

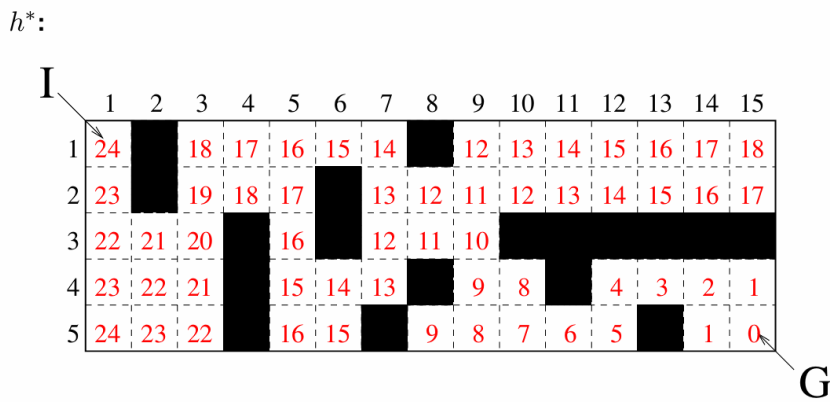
$h$  depend only on the state  $s$  and not on the search node (the path taken to reach  $s$  don't affect  $h(s)$ ). If  $n$  is a search node, we usually write  $h(n)$  instead of  $h(n.State)$  as an abuse of notation.

The purpose of  $h$  is to estimate how far we are from a solution, clearly time is needed to calculate the value of  $h$ , ideally we would like a function  $h$  that is very precise in the estimate, and which is quick to calculate, these two objectives are generally in conflict with each other.

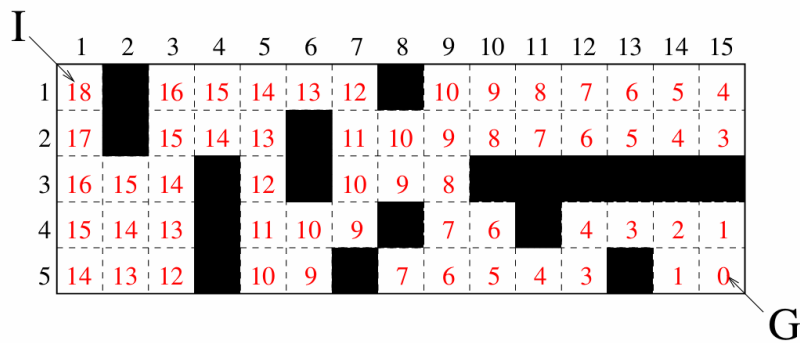
Given a problem  $\Pi$ , a heuristic function  $h$  for  $\Pi$  can be obtained as goal distance within a simplified (relaxed) problem  $\Pi'$ . A classic example is using Euclidean distance as a heuristic function in a graph representing points on a map.



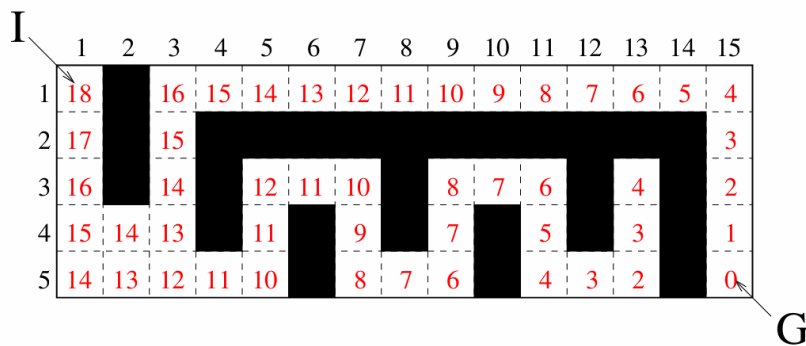
Another example is when we have to reach a goal in discrete grid with some "pitfalls" to avoid.



in each cell is presented the value of the perfect heuristic  $h^*$ . A reasonable and accurate heuristic may be the Manhattan distance.

**Manhattan Distance, “accurate  $h$ ”:**

For some specific grid configurations this heuristic might be inaccurate:

**Manhattan Distance, “inaccurate  $h$ ”:**

**Definition 7** Let  $h^*$  to be a perfect heuristic. A heuristic  $h$  is **admissible** if  $\forall s \in S$  we have  $h(s) \leq h^*(s)$ .

An admissible heuristic function never overestimates the cost to reach the goal.

**Definition 8** A heuristic  $h$  is **consistent** if, for all transition  $s \xrightarrow{a} s'$  in  $\Theta$ , we have  $h(s) - h(s') \leq c(a)$ .

The consistency is important, when applying an action  $a$ , the heuristic value cannot decrease by more than the cost of  $a$ .

**Proposition 1** If  $h$  is consistent then is admissible.

*Proof:* The proof is done by induction.

- Base case: let  $s$  to be the goal state, so  $h(s) = h^*(s) = 0$ , so  $h(s) \leq h^*(s)$  trivially holds.
- Assume that the claim holds for all states  $s'$  where the cheapest path to the goal state have length  $n$ . Let  $s$  to be a state with goal length  $n + 1$ , which have the first transition

$$s \xrightarrow{a} s'$$

so

$$h(s) \leq h(s') + c(a) \quad (2.13)$$

by the inductive hypothesis:

$$h(s') \leq h^*(s') \quad (2.14)$$

by construction:

$$h^*(s) = h^*(s') + c(a) \quad (2.15)$$

by combining the equations:

$$\begin{cases} h(s) \leq h(s') + c(a) \\ h(s') \leq h^*(s') \\ h^*(s) = h^*(s') + c(a) \end{cases} \implies h(s) \leq h^*(s) \quad (2.16)$$

■

The Euclidean distance for the map-travel problem is consistent.

#### Systematic search vs. local search:

- **Systematic search strategies:** No limit on the number of search nodes kept in memory at any point in time.
  - Guarantee to consider all options at some point, thus complete.
- **Local search strategies:** Keep only one (or a few) search nodes at a time.
  - No systematic exploration of all options, thus incomplete.

#### Tree search vs. graph search:

- For the systematic search strategies, we consider graph search algorithms exclusively, i.e., we use duplicate pruning.
- There are tree search versions of these algorithms. These are easier to understand, but aren't used in practice. (Maintaining a complete open list, the search is memory-intensive anyway.)

The informed search uses problem-specific knowledge, by using an evaluation function  $f$  at each node, that function estimate the "desirability" of expanding that node, and we choose to expand the most desirable unexpanded node

$$\arg \max_{\text{unexpanded } n} f(n).$$

We usually implement this with a sorted queue ordered based on the desirability of expansion, this queue is called **frontier**.

#### 2.4.1 Greedy Best-First Search

We use the heuristic function  $f$  as the evaluation function and we expand the node that is closest to the goal. This algorithm is complete thanks to the assumption that the state space is finite, but is not

---

##### Algorithm 8 Greedy Best-First Search

---

**Require:** A problem  $\Pi$

```

 $n \leftarrow$  a node  $n$  such that  $n.State = \Pi.InitialState$ 
 $frontier \leftarrow$  a priority queue ordered by ascending  $h$ 
 $explored \leftarrow$  empty set of states
while True do
  If  $frontier$  is empty Then Return failure
   $n \leftarrow Pop(frontier)$ 
  If  $n.State$  is the goal state Then Return  $Solution(s)$ 
   $explored \leftarrow explored \cup n.State$ 
  for each action  $a$  in  $\Pi.Actions(n.State)$  do
     $n' \leftarrow ChildNode(\Pi, n, a)$ 
    If  $n'.State \notin explored \cup States(frontier)$  Then insert( $n', h(n')$ ,  $frontier$ )
  end for
end while

```

---

optimal, there exists an algorithm that is optimal and complete.

### 2.4.2 The A\* Algorithm

A\* is an informed search algorithm used for path finding and graph traversal. It finds the shortest path from a starting node to a target goal by combining the benefits of Dijkstra's algorithm and Greedy Best-First Search.

The core of A\* is the evaluation function  $f(n)$ , which determines the priority of each node  $n$  in the search frontier:

$$f(n) = g(n) + h(n)$$

- $g(n)$  (**The Path Cost**): The actual cost of the path from the start node to the current node  $n$ . This ensures the algorithm doesn't ignore the distance already traveled.
- $h(n)$  (**The Heuristic**): An estimated cost to get from node  $n$  to the goal. This "educated guess" allows the algorithm to prioritize nodes that appear to lead directly toward the target.
- $f(n)$  (**The Total Estimated Cost**): The estimated cost of the cheapest solution passing through node  $n$ .

The algorithm maintains a **frontier** (priority queue) of nodes to be explored, ordered by ascending  $f(n)$  values.

1. It always expands the node with the **lowest**  $f(n)$  first.
2. If  $h(n)$  is *admissible* (never overestimates the actual cost), A\* is guaranteed to find the optimal (shortest) path.
3. By balancing  $g(n)$  and  $h(n)$ , A\* avoids expanding paths that are already too expensive while remaining focused on the goal.

---

#### Algorithm 9 A\*

**Require:** A problem  $\Pi$

```

 $n \leftarrow$  a node  $n$  such that  $n.State = \Pi.InitialState$ 
 $frontier \leftarrow$  a priority queue ordered by ascending  $g + h$ 
 $explored \leftarrow$  empty set of states
while True do
    if  $frontier$  is empty Then Return failure
     $n \leftarrow Pop(frontier)$ 
    if  $n.State$  is the goal state Then Return Solution( $s$ )
     $explored \leftarrow explored \cup n.State$ 
    for each action  $a$  in  $\Pi.Actions(n.State)$  do
         $n' \leftarrow ChildNode(\Pi, n, a)$ 
        if  $n'.State \notin explored \cup States(frontier)$  then
            insert( $n', h(n') + g(n')$ ,  $frontier$ )
        else if  $\exists n''$  s.t.  $n''.State = n'.State$  and  $g(n') < g(n'')$ 
            Replace  $n''$  in  $frontier$  with  $n'$ 
        end if
    end for
end while

```

---

Now we want to prove the optimality of A\*.

**Definition 9** Let  $\Pi$  to be a problem with state space  $\Theta = (S, A, c, T, I, S^G)$  and let  $h$  to be a consistent heuristic function for  $\Pi$ . We define the  **$h$ -weighted state space** as

$$\Theta = (S, A^h, c^h, T^h, I, S^G)$$

where

- $A^h = \{a[s, s'] \mid a \in A, s \in S, s' \in S, (s, a, s') \in T\}$
- $c^h : A^h \rightarrow \mathbb{R}^+ \cup \{0\}$  is the function

$$c^h(a[s, s']) = c(a) - [h(s) - h(s')]$$

- $T^h = \{(s, a[s, s'], s') \mid (s, a, s') \in T\}$

In this *weighted state space*, from each action cost, we subtract the "gain in heuristic value". This  $\Theta^h$  is well defined since

$$c(a) - [h(s) - h(s')] \geq 0.$$

**Lemma 1**  $\Theta$  and  $\Theta^h$  have the same optimal solutions.

*Proof:*

- Let  $I$  be a solution path in  $\Theta$  ending in a goal state  $s_n \in S^G$ :

$$I = s_0 \xrightarrow{a_1} s_1, \dots, s_{n-1} \xrightarrow{a_n} s_n$$

- The cost of the same path in  $\Theta^h$  is defined by the sum of the modified transition costs:

$$\begin{aligned} &[-h(s_0) + c(a_1) + h(s_1)] + \\ &[-h(s_1) + c(a_2) + h(s_2)] + \dots + \\ &[-h(s_{n-1}) + c(a_n) + h(s_n)] \end{aligned}$$

- Simplifying the telescoping sum, we see that all intermediate heuristic terms  $h(s_i)$  for  $0 < i < n$  cancel out:

$$\sum_{i=1}^n c(a_i) - h(s_0) + h(s_n)$$

- Substituting constants, where  $h(I)$  denotes the heuristic of the initial state  $h(s_0)$ , and noting that  $h(s_n) = 0$  for goal states:

$$\left[ \sum_{i=1}^n c(a_i) \right] - h(I)$$

- Conclusion: Thus, the costs of solution paths in  $\Theta^h$  are those of  $\Theta$ , minus a constant. The claim follows. ■

**Lemma 2** The search space of  $A^*$  on  $\Theta$  is isomorphic to the search space of the uniform-cost-search on  $\Theta^h$ .

*Proof:*

- Let  $I = s_0 \xrightarrow{a_1} s_1, \dots, s_{n-1} \xrightarrow{a_n} s_n$  be any path in  $\Theta$ .
- The  $g + h$  value, used by  $A^*$ , is:

$$\left[ \sum_{i=1}^n c(a_i) \right] + h(s_n)$$

- The  $g$  value in  $\Theta^h$ , used by uniform-cost search on  $\Theta^h$ , is:

$$\left[ \sum_{i=1}^n c(a_i) \right] - h(I) + h(s_n)$$

(see previous proof).

- The difference  $-h(I)$  is constant, so the ordering of the open list is the same.
- The duplicate elimination is identical. ■

**Theorem 2 (Optimality of  $A^*$ )** Let  $\Pi$  be a problem and  $h$  a heuristic function for  $\Pi$ . If  $h$  is consistent, the solution returned by  $A^*$  (if any) is optimal.

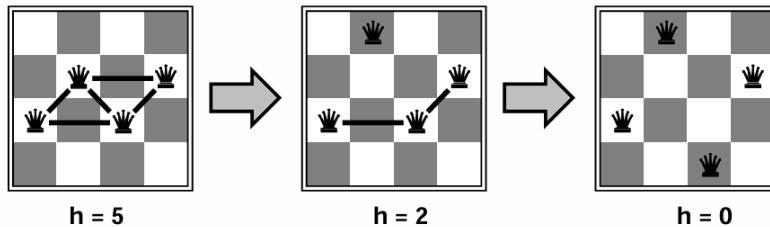
*Proof:* Let  $\vec{s}(A^*, \Theta)$  be the sequence of states (solution) returned by  $A^*$  on  $\Theta$ , we denote  $\vec{S}(UCS, \Theta^h)$  the set of solutions that could in principle be returned by uniform-cost-search run on  $\Theta^h$ . By the Lemma 2, we have

$$\vec{s}(A^*, \Theta) \in \vec{S}(UCS, \Theta^h) \tag{2.17}$$

since the uniform cost search is optimal, every element of  $\vec{S}(UCS, \Theta^h)$  is an optimal solution for  $\Theta^h$ , thus  $\vec{s}(A^*, \Theta)$  is optimal for  $\Theta^h$ , by the Lemma 1, this implies that  $\vec{s}(A^*, \Theta)$  is an optimal solution for  $\Theta$ . ■

## 2.5 Local Search

In local search, we take decisions based on the values of the heuristic function  $h$  only on the immediate neighbor states. In many optimization problems, the path is irrelevant, and the goal state itself is the solution, in these cases the state space is a set of "complete" configurations and the task is to find the optimal configuration. Let's consider the example of the  $n$ -queens:



We have a  $n \times n$  board and we have to put  $n$  queens with no two queens on the same row, column or diagonal. A local search strategy is to move one single queen at time to reduce the number of conflicts. So the algorithm explores all the boards deriving from the movement of one single queen (this simple method is in fact very effective and solve the problem almost always for very large  $n$ ).

Local search algorithms are *not systematic*, the advantages are

- they use very little memory, typically a constant space.
- find reasonable solutions in large or infinite state space where systematic search is unsuitable.

Is useful to solve pure mathematical optimization problems.

### 2.5.1 Hill Climbing

The **hill-climbing** Algorithm 10 keeps choosing actions leading to a direct successor state with best heuristic value. It stops when no more immediate improvements can be made.

---

#### Algorithm 10 Hill-Climbing

**Require:** A problem  $\Pi$

```

 $n \leftarrow$  a node  $n$  with  $n.State = \Pi.InitialState$ 
while True do
    if  $n.State$  is the goal state then return  $n$ 
     $N \leftarrow$  set of all child nodes of  $n$ 
     $n \leftarrow \arg \min_{n \in N} h(n)$ 
    if stopping condition are met then return  $n$ 
end while

```

---

One stopping condition may be: If the neighbor node that minimize  $h$  have a value for  $h$  bigger than the current node, stop. The *Gradient Descent* is an interpretation of this algorithm.

This algorithm is not complete neither optimal. Usually the hill climbing algorithm get stuck in local minima, where all the neighbors look worse than the current node, if all the neighbors have the same value of  $h$ , the next move will be chosen at random.

A possible strategy is to perform different runs of the algorithm, by re-starting it every time we reach a local minimum, or when we have spent a certain amount of time without "making progress" (without finding a node that decreases on  $h$ ). A way to avoid local minima is to do random walks in the hope that these will lead out of them.

A variant of the hill climbing is the **Local Beam Search**, where  $k$  states are kept in memory instead of 1, and at each step, we generate all the successors of these  $k$  states and we update our current state with the best  $k$  among these successors.

### 2.5.2 Simulated Annealing

The idea is to escape local maxima by allowing some "bad" moves but gradually decrease their size and frequency.

The Algorithm 11 shows how the probability of accepting a "bad" move decreases exponentially with the "badness" of the move and the "temperature" going down.

But, what is the temperature in this context? We consider a scalar function that assigns to each time step  $t$  a value of "temperature", this function is called *schedule*, and usually decreases over time. Bigger is the temperature, and bigger is the probability to accept bad moves, so it defines how this probability of doing "bad moves" decreases over time.

---

#### Algorithm 11 Simulated-Annealing

---

**Require:** A problem  $\Pi$ , a map from time to temperature *schedule*  
 $current \leftarrow$  a node *current* with  $current.State = \Pi.InitialState$   
**for**  $t = 1$  to  $\infty$  **do**  
     $T \leftarrow schedule(t)$   
    **if**  $T = 0$  **then return** *current*  
     $next \leftarrow$  a randomly selected successor of *current*  
     $\Delta E = h(next) - h(current)$   
    **if**  $\Delta E > 0$  **then**  $current \leftarrow next$  /\*the successor is actually better\*/  
    **else**  $current \leftarrow next$  with probability  $\exp(\Delta E/T)$  /\*do a bad move\*/  
**end for**

---

### 2.5.3 Genetic Algorithms

The key idea of a genetic algorithm is the following

- we generate a starting **population** of  $k$  randomly selected states
- each individual state is represented by a string on a finite alphabet
- the next generation of states is generated as follows:
  1. we consider a **fitness function** that should assigns higher values to better states
  2. we select random pairs of states from the population, and we generate a new state by making these two **reproduce**
  3. the probability of being chosen is proportional to the fitness value
- we add some **random mutations** with small probability.

Making two states reproduce: Create a new state by selecting some characteristics from the parent states.  
 An example is

$$\begin{aligned} S1 &= \langle \textcolor{blue}{a} \ s \ \textcolor{blue}{w} \ j \rangle \\ S2 &= \langle f \ \textcolor{blue}{h} \ d \ \textcolor{blue}{e} \rangle \\ \text{the results of their reproduction is:} \\ S3 &= \langle a \ h \ w \ e \rangle \end{aligned}$$

---

#### Algorithm 12 GENETIC-ALGORITHM

---

**Require:** A starting *population* of states

---

- Genetic algorithms combine an uphill tendency with random exploration and exchange of information among parallel search threads.
- Their primary advantage comes from the crossover operation.

- Crossover helps iff substrings are meaningful components: we combine large blocks of letters that have evolved independently to perform useful functions.
- Much work remains to be done to identify the conditions under which genetic algorithms perform well.

## 2.6 Adversarial Search

One of the oldest sub-areas of AI is *Game Planning*, we model games as search problems that takes in account the competition between two opponents (such as chess). We consider simple games that satisfies the following restrictions:

- the set of game states is discrete
- the number of possible moves at each step is finite
- the game state is fully observable and each move's outcome is deterministic
- there are only two players
- the players playing in alternating turns
- the utility function  $u$  must be maximized from one player and minimized from the other
- there are no infinite runs of the game, after a finite number of steps, the game must end.

We consider only zero-sum games, where the two players play with the same conditions, without favor. We denote *Max* and *Min* the two players.

**Definition 10** A *game state space* is a 6-tuple  $\Theta = (S, A, T, I, S^T, u)$  where

- $S$  is the set of states, can be partitioned in  $S = S^{Max} \cup S^{Min} \cup S^T$ , denoting the state where *Max* or *Min* plays.
- $A$  is the set of actions, can be partitioned in  $A = A^{Max} \cup A^{Min}$ 
  - $A^{Max}$  is the set of actions that *Max* can take,  $A^{Min}$  is the set of actions that *Min* can take.
  - for  $a \in A^{Max}$ , if  $s \xrightarrow{a} s'$  then  $s \in S^{Max}$  and  $s' \in S^{Min} \cup S^T$
  - for  $a \in A^{Min}$ , if  $s \xrightarrow{a} s'$  then  $s \in S^{Min}$  and  $s' \in S^{Max} \cup S^T$
- $T$  is the set of deterministic transition relation
- $I$  is the initial state
- $S^T$  are the set of terminal states
- $u : S^T \rightarrow \mathbb{R}$  is the utility function

**Definition 11** A *strategy* for *Max* is a function  $\sigma^{Max} : S^{Max} \rightarrow A^{Max}$  such that,  $a$  is applicable to  $s$  if  $\sigma^{Max}(s) = a$ . Analogous for *Min* with  $\sigma^{Min}$ .

$\sigma^{Max}$  ( $\sigma^{Min}$ ) is defined for all states in  $S^{Max}$  ( $S^{Min}$ ), since the agent don't know how the opponent will react he needs to prepare for all possibilities. A strategy is optimal if it yields the best possible utility assuming perfect opponent play. For an adversarial search problem there are three types of solutions:

- **ultra weak:** Prove whether the first player will win, lose or draw from the initial position, given perfect play on both sides.
- **weak:** Provide a strategy that is optimal from the beginning of the game for one player against any possible play by the opponent.
- **strong:** Provide a strategy that is optimal from any valid state, even if imperfect play has already occurred on one or both sides.

Computing a strategy is often unfeasible, the number of reachable states are big, in chess, there are  $10^{40}$  possible board states. We will consider a *Black Box* description of the games.

### 2.6.1 Minimax Search

This is the canonical algorithm to solving games, by computing an optimal strategy. Remember that the player *Max* attempts to maximize the utility function  $u(s)$  of the terminal state that will be reached during play, when *Min* attempts to minimize it.

We describe the algorithm playing as *Max*, and we assume that the opponents will play by trying always to minimize the utility function. Starting from the actual state, the algorithm explores all the possible outcomes from all possible feasible sequences of moves, expanding a tree.

---

**Algorithm 13** Minimax

---

**Require:**  $s$   
 $v \leftarrow \text{MaxValue}(s)$   
**return** an action  $a \in \text{Actions}(s)$  yielding value  $v$

---



---

**Algorithm 14** MaxValue

---

**Require:**  $s$   
**if** TerminalTest( $s$ ) **then**  
    **return**  $u(s)$   
**end if**  
 $v \leftarrow -\infty$   
**for** each  $a \in \text{Actions}(s)$  **do**  
     $v \leftarrow \max(v, \text{MinValue}(\text{ChildState}(s, a)))$   
**end for**  
**return**  $v$

---



---

**Algorithm 15** MinValue

---

**Require:**  $s$   
**if** TerminalTest( $s$ ) **then**  
    **return**  $u(s)$   
**end if**  
 $v \leftarrow +\infty$   
**for** each  $a \in \text{Actions}(s)$  **do**  
     $v \leftarrow \min(v, \text{MaxValue}(\text{ChildState}(s, a)))$   
**end for**  
**return**  $v$

---

Consider the tree in figure 2.3, starting from the root, the player consider all possible outcomes, the leaf are terminating states, assuming that the opponents will always tries to minimize  $u$ :

- if *Max* choose the left branch, *Min* will choose the action that leads to the terminal state with  $u(s) = 3$
- if *Max* choose the middle branch, *Min* will choose the action that leads to the terminal state with  $u(s) = 2$
- if *Max* choose the right branch, *Min* will choose the action that leads to the terminal state with  $u(s) = 2$

So the best moves is the one that leads to the left branch.

This is the simplest possible game search algorithm, but in practice is unfeasible since the search tree are way too large to expand. The minimax algorithm is:

- Complete, if the tree is finite.
- Optimal against an optimal opponent.

Since it is a depth first search, the time complexity is  $O(b^m)$ , where  $b$  is the branching factor and  $m$  is the depth of the solution. The space complexity is  $O(bm)$ . In chess,  $b \simeq 35$  (possible moves approximately) and a reasonable game terminates in 80 turns, so  $m \simeq 80$ , in such a case the state space is  $O(35^{80})$  (it's impossible in practice to expand the tree).

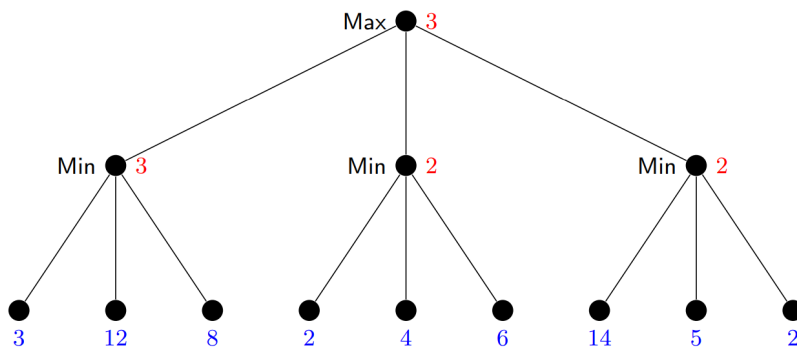
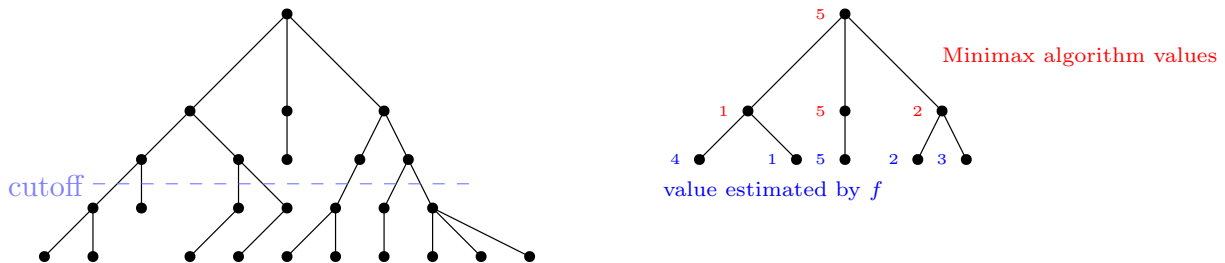


Figure 2.3: Game's tree

### 2.6.2 Evaluation Functions

Since the Minimax game tree are too big, we impose a depth limit  $d$  (called horizon) on the search, and we apply an evaluation function to the non-terminal states in that horizon. An evaluation function  $f : S \rightarrow \mathbb{R}$  should work to estimate the actual value reachable from a state such as in the unlimited-depth Minimax. If a state is terminal, we use the actual value  $u$ . We want  $f$  to be accurate and fast to compute.

While applying this algorithm, we can consider a depth limit  $d$ , evaluate  $f$  on all the states at that depth, and then considering these cut-off tree and apply the Minimax algorithm.



Usually, the evaluation function is a linear weighted function, such as

$$f(s) = \sum_{i=1}^n w_i f_i(s) \quad (2.18)$$

where  $f_1 \dots f_n$  are features extracted from the state  $s$ , designed by human experts of the considered game/domain, and  $w_i \in \mathbb{R}$  are real weights, that can be learned automatically (with machine learning algorithms). For example, features in the tetris game could be the number of holes in the block grid, or the maximum height reached by a placed block, as shown in figure 2.4.

This algorithm is critical since critical aspects of the game may be cut-off by the horizon. Since we want to search *as deeply as possible in a given time*, we could apply the *iterative deepening search*, until the time's up, and then return the solution of the deepest completed search.

A better solution is called *Quiescence search*. Instead of using a fixed search depth  $d$ , quiescence search dynamically adapts the depth to handle "unquiet" positions where the evaluation function's value is likely to fluctuate quickly—typically due to immediate, forcing moves like captures or checks.

For example, in Chess, if a piece exchange is underway, the search continues past the standard depth limit until a "quiet" state is reached, ensuring the final evaluated position is stable and doesn't miss an immediate, significant change in material or safety, thus leading to a much more accurate evaluation.

### 2.6.3 Alpha-Beta Pruning

There is a way to save time during the computation, we can cut-off some branches of the search for which we know at priori that will not lead to the solution. Consider the tree in figure 2.5.

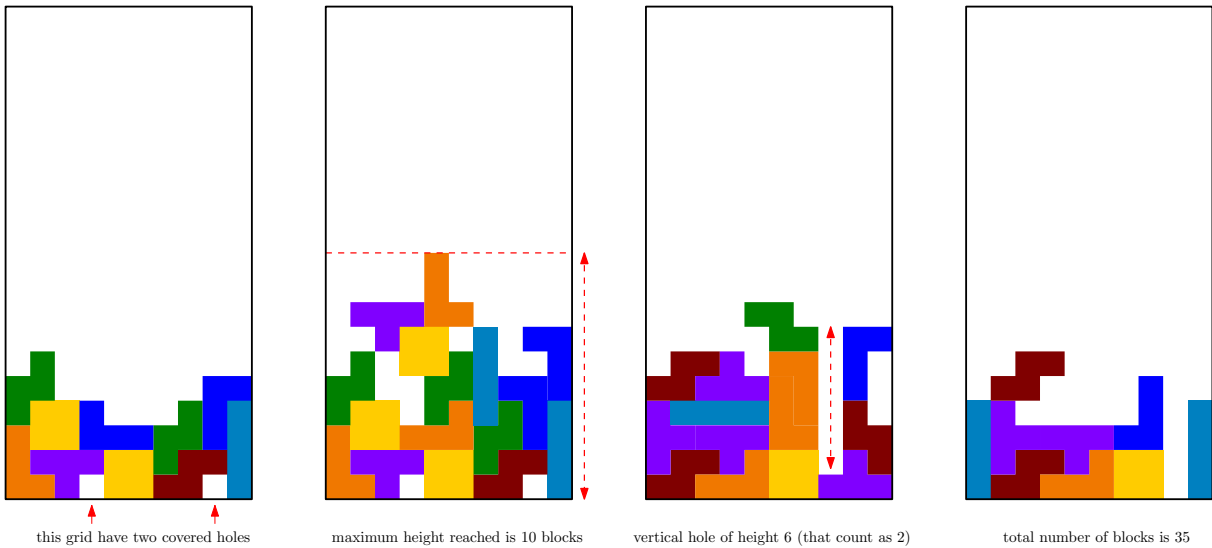
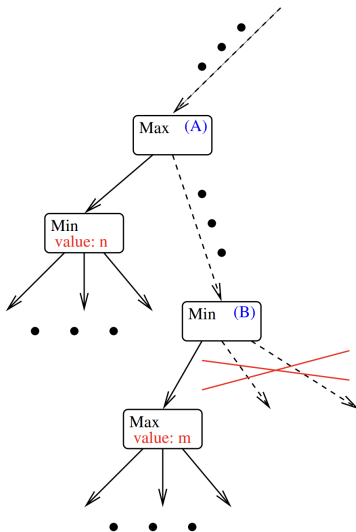


Figure 2.4: Grid's properties

Figure 2.5: Let's say  $n > m$ 

The left branch leads to a utility at least  $n$ , the right branch have one sub-branch with utility value  $m < n$ , so we can say for sure that the left tree will not provide values for the utility greater than  $m$  (since it's the *Min* turn), at this point we can discard the left branch and continue with the right one.

We consider two additional variables  $\alpha, \beta$  defined on each node during the search, such that:

- for each node  $n$ ,  $\alpha$  is the highest utility that search has already found for *Max* on its path to  $n$ .
  - In a *Min* node, if one of the successors has a utility value less or equal than  $\alpha$ , we can stop considering  $n$  and cutting it off (pruning out its remaining successors).
- We can consider a dual method for *Min*, for each node  $n$ ,  $\beta$  is the lowest utility that search has already found for *Min* on its path to  $n$ .
  - in a *Max* node, if one of the successors has a utility value greater or equal than  $\beta$ , we can stop considering  $n$  and cutting it off (pruning out its remaining successors).

---

**Algorithm 16** AlphaBetaPruning

---

**Require:**  $s$ 

```
 $v \leftarrow \text{MaxValue}(s, -\infty, +\infty)$ 
return an action  $a \in \text{Actions}(s)$  yielding value  $v$ 
```

---



---

**Algorithm 17** MaxValue

---

**Require:**  $s, \alpha, \beta$ 

```
if TerminalTest( $s$ ) then
    return  $u(s)$ 
end if
 $v \leftarrow -\infty$ 
for each  $a \in \text{Actions}(s)$  do
     $v \leftarrow \max(v, \text{MinValue}(\text{ChildState}(s, a)), \alpha, \beta)$ 
     $\alpha \leftarrow \max(\alpha, v)$ 
    if  $v \geq \beta$  then return  $v$ 
end for
return  $v$ 
```

---



---

**Algorithm 18** MinValue

---

**Require:**  $s, \alpha, \beta$ 

```
if TerminalTest( $s$ ) then
    return  $u(s)$ 
end if
 $v \leftarrow +\infty$ 
for each  $a \in \text{Actions}(s)$  do
     $v \leftarrow \min(v, \text{MaxValue}(\text{ChildState}(s, a)), \alpha, \beta)$ 
     $\beta \leftarrow \min(\beta, v)$ 
    if  $v \leq \alpha$  then return  $v$ 
end for
return  $v$ 
```

---

### 2.6.4 Monte-Carlo Tree Search

The Alpha-Beta search have two issues

- it needs an accurate and fast evaluation function
- it doesn't explore so much in problems with a large branching factor like *Go*.
- **State Value Estimation:** The value of a state is estimated by calculating the average utility across multiple **playouts** (complete game simulations starting from that specific state).
- **Playout Policy:** During a playout, a policy is used to bias move selection toward "good" moves rather than purely random ones. For complex games like Go, these policies are often learned via **self-play** using neural networks.
- **Pure Monte Carlo Search:** This approach runs simulations from the current game state and tracks which possible moves yield the highest win percentage.
- **Selection Policy:** This policy optimizes computation by focusing on the most important branches of the game tree. Its primary function is to balance the trade-off between:
  - **Exploration:** Searching less-visited parts of the tree to find potentially better moves.
  - **Exploitation:** Deepening the search in branches already known to have high win rates.

In the Monte Carlo Sampling we evaluate actions through sampling:

- Exploitation: play in the machine that returns the best reward.
- Exploration: play machines that have not been tried a lot yet.

The **UBC** (Upper Confidence Bound) is a formula that automatically balances exploration and exploitation to maximize total gains. The procedure of the search is described in the following steps:

1. **Selection:** Starting at the root, traverse the tree by applying the *selection policy* to choose successors until a leaf node that has not been fully expanded is reached.
2. **Expansion:** Grow the search tree by generating a new child of the selected leaf node by applying a new action.
3. **Simulation:** Perform a complete run of the game (a *playout*) starting from the newly generated child node. Moves for both players are selected according to the *playout policy* to obtain a final reward.
4. **Back-propagation:** Use the result of the simulation to update the value estimates of all search tree nodes along the path back up to the root.

Once the computational time or resources are exhausted, the algorithm chooses the "best" move based on the accumulated statistics and plays it. We have to define a formula that balances exploration and exploitation, a formula Inspired by Multi-Armed Bandit problems is the following

$$UBC1(n) = \frac{U(n)}{N(n)} + C \times \sqrt{\frac{\log N(PARENT(n))}{N(n)}} \quad (2.19)$$

where

- $U(n)$  is the total utility of all playouts that went through the node  $n$
- $N(n)$  is the number of playouts through node  $n$
- $PARENT(n)$  is the parent of  $n$  in the tree.

the left side of the sum represent the exploitation factor and the right side the exploration.

- MTCS is chosen when branching factor becomes very high and the evaluation function is difficult.
- MTCS is more robust than Alpha-Beta search (it is also possible to combine MCTS and evaluation functions).
- MTCS can be applied to unknown games.
- MTCS is a form of reinforcement learning.

---

**Algorithm 19** Monte-Carlo-Tree-Search

---

**Require:** *state*

```
tree  $\leftarrow$  Node(state)
while Is-Time-Remaining() do
    leaf  $\leftarrow$  Select(tree)
    child  $\leftarrow$  Expand(leaf)
    result  $\leftarrow$  Simulate(child)
    Back-Propagate(result, child)
end while
```

**return** the move in Actions(*state*) whose node has highest number of playouts.

---

CHAPTER

---

3

# CONSTRAINT SATISFACTION PROBLEMS

**Definition 12** A *CSP* (*constraint satisfaction problem*) is composed by a set of variables, each associated with its domain, and a set of constraints over these variables, the goal is to find an assignment of variables so that every constraint is satisfied.

In SuDoKu, the variables are the content of the cells, the domain of each cell is  $\{1, 2, \dots, 9\}$ , and the constraints are that, each number should appear only once in each row, column or block. Another CSP problem is the *graph coloring* problem, where we should assign one of the  $k$  possible color to each node, such that two adjacent nodes must have different colors (this problem is NP-hard for  $k = 3$ ).

### 3.1 Constraint Networks

**Definition 13** A *Binary Constraint Network* is a tuple  $\gamma = (V, D, C)$  where:

- $V = \{v_1, v_2 \dots v_n\}$  is a finite set of variables.
- $D = \{D_{v_1}, D_{v_2} \dots D_{v_n}\}$  is a corresponding set of finite domains.
- $C$  is a set of binary relations that models the constraints. A relation is denoted  $C_{uv}$  with  $u, v$  variables in  $V$ .

$$C_{uv} \subseteq D_u \times D_v$$

If  $C_{uv} \in C$  and  $C_{xy} \in C$ , then  $\{u, v\} \neq \{x, y\}$  (no redundancy).

$C_{uv}$  defines the permissible combined assignments to  $u$  and  $v$ . For example, if

$$u, v \in V \tag{3.1}$$

$$v' \in D_v \tag{3.2}$$

$$u' \in D_u \tag{3.3}$$

$$C_{uv} \subseteq D_v \times D_u \in C \tag{3.4}$$

$$(v', u') \notin C_{uv} \tag{3.5}$$

the assignment  $v = v', u = u'$  violates the constraints. Let's see an example, we consider the map of australia show in figure 3.1. We want to colorate each state with one of 3 possible colors, without having two adjacent state with the same color.

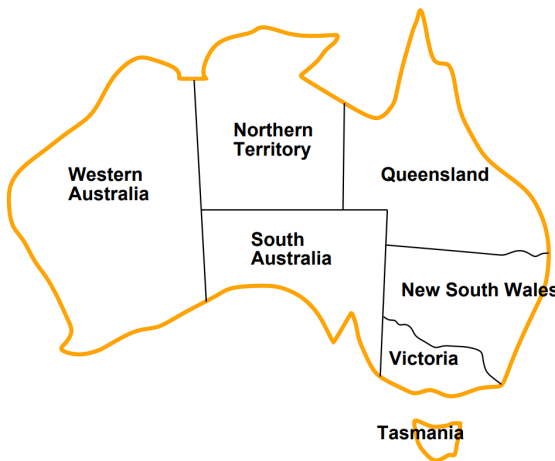


Figure 3.1: Coloring Australia

The variables are the states

$$V = \{WA, NT, SA, Q, NSW, V, T\}$$

The domain for each variable  $v \in V$  is  $D_v = \{\text{red}, \text{green}, \text{blue}\}$ . If all the variables have the same domain

$$\forall v \neq u, D_v = D_u \quad (3.6)$$

we do an notation abuse saying that the domain of the problem is  $D = \{\text{red}, \text{green}, \text{blue}\}$ . For each adjacent couple of states  $u, v$ , there is a constraint

$$C_{uv} = \{(d, d') \in D \times D : d \neq d'\}.$$

Constraint Networks can be extended

- the domains  $D_v$  may be infinite or uncountable, like  $D_v = \mathbb{R}$ .
- the constraints may have an arity higher than 2, with relations over  $k > 2$  variables, like in CNF satisfiability.

A constraint may be **unary**:  $C_v \subseteq D_v$ , in this case the domain of a single variable is restricted, equivalently to setting  $D_v = C_v$ .

There exists CSP solvers, generic algorithms that find assignment for constraints problem, by taking in input a constraint network as the generic language to describe the problem. The core exercise in this context is to model a problem with a constraint network.

**Definition 14** Let  $\gamma = (V, D, C)$  to be a constraint network, a **partial assignment** is a function

$$a : V' \rightarrow \bigcup_{v \in V} D_v$$

where

- $V' \subseteq V$
- $a(v) \in D_v$  for all  $v \in V'$

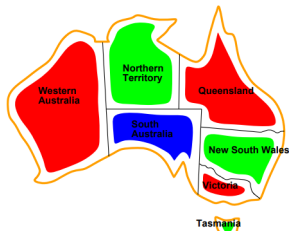
if  $V' = V$  is a **total assignment** (or just assignment). A partial assignment assigns some variables to values from their respective domains. A total assignment is defined on all variables.

**Definition 15** Let  $\gamma = (V, D, C)$  to be a constraint network and let  $a$  to be a partial assignment, we say that  $a$  is **inconsistent** if there exists variables  $u, v \in V$  such that

- $a$  is defined on  $u$  and  $v$

- $C_{uv} \in C$
- $(a(u), a(v)) \notin C_{uv}$ .

If a partial assignment is not inconsistent, is **consistent**. An assignment  $a$  is a **solution** if it's total and consistent. A constraint network  $\gamma$  is **solvable** if there exists a solution  $a$ , otherwise is **unsolvable**.



- **Variables:**  $V = \{WA, NT, SA, Q, NSW, V, T\}$ .
- **Domains:** All  $v \in V$ :  $D_v = D = \{\text{red}, \text{green}, \text{blue}\}$ .
- **Constraints:**  $C_{uv}$  for adjacent states  $u$  and  $v$ , with  $C_{uv} = \{(d, d') \in D \times D \mid d \neq d'\}$ .
- **Solution:**  $\{WA = \text{red}, NT = \text{green}, SA = \text{blue}, Q = \text{red}, NSW = \text{green}, V = \text{red}, T = \text{green}\}$ .

**Definition 16** Let  $\gamma = (V, D, C)$  to be a constraint network, and let  $a$  to be a partial assignment.  $a$  can be **extended** to a solution if there exists a solution  $a'$  that agrees with  $a$  on the variables where  $a$  is defined.

- the domain of  $a$  is  $V' \subset V$
- for each  $v \in V'$ , we have that  $a(v) = a'(v)$ .

**Proposition 2** if  $a$  can be extended to a solution, then it's consistent (the opposite doesn't necessary holds).

Given a constraint network  $\gamma = (V, D, C)$ , if  $n = |V|$  and  $\forall D_v \in D$ ,  $|D_v| = k$ , the number of total assignments is  $k^n$ . There are at most  $n^2$  constraints, each constraint have size at most  $k^2$ , the number of assignments is exponentially bigger than the size of  $\gamma$ .

**Theorem 3** The problem to decide if a constraint network  $\gamma = (V, D, C)$ , is solvable is NP-complete.

## 3.2 Naive Backtracking

The following algorithm is simple and used to find a solution for a constraint network. The method is recursive, the first step given assignment  $a$  is the empty assignment. With *Backtracking*, we mean the

---

### Algorithm 20 NaiveBacktracking

---

```

Require:  $\gamma = (V, D, C), a$ 
  if  $a$  is inconsistent then
    return inconsistent
  end if
  if  $a$  is a total assignment then
    return  $a$ 
  end if
  let  $V'$  to be the domain of  $a$ 
  select some  $v \in V - V'$  (select a variable for which  $a$  is not defined)
  for each  $d \in D_v$  in some order do
     $a' = a \cup \{v = d\}$ 
     $a'' = \text{NaiveBacktracking}(\gamma, a')$ 
    if  $a''$  is not inconsistent then
      return  $a''$ 
    end if
  end for
  return inconsistent

```

---

action of recursively instantiate variables one-by-one, backing up out of a search branch if the current partial assignment is already inconsistent. Note that in the algorithm 20 we are iterating the possible

values in  $D_v$  in a specific **order** that is not described yet, and is not described which variable are we picking at each recursion step.

The Naive Backtracking algorithm is

- simple to implement
- much more efficient than enumerating all possible assignment
- complete, if there is a solution, backtracking will find it.

This algorithm can't predict if an assignment  $a$  can't be extended to a solution unless  $a$  is already inconsistent.

### 3.2.1 About the Order

The ordering in the for loop of algorithm 20 can influences the search space size. If no solution exists below current node, the order doesn't matter, the algorithm will search the whole sub-tree anyway, otherwise, if a solution does exist below current node, by choosing the "correct" value, then no backtracking is needed.

A common strategy is to consider the variable "most constrained", at each recursion step, we pick the variable  $v$  that minimize the size of the set

$$\{d \in D_v : a \cup \{v = d\} \text{ is consistent}\} \quad (3.7)$$

by choosing a most constrained variable  $v$  first, we reduce the branching factor (number of sub-trees generated for  $v$ ) and thus reduce the size of our search tree.

Another common heuristic is to choose the variable that is the "most constraining", we choose  $v$  that maximize the size of the set

$$\{u \in V : a(u) \text{ is undefined} \wedge C_{uv} \in C\} \quad (3.8)$$

by choosing a most constraining variable first, we detect inconsistencies earlier on and thus reduce the size of our search tree.

For the order of the value in the for loop, we can choose the least constraining value first, for a variable  $v$ , we choose the value  $d \in D_v$  that minimize the size of the set

$$\{d' : d' \in D_u \wedge a(u) \text{ is undefined} \wedge C_{uv} \in C \wedge (d', d) \notin C_{uv}\} \quad (3.9)$$

by choosing a least constraining value first, we increase the chances to not rule out the solutions below the current node.

## 3.3 Inference

Given a constraint networks, we would like to obtain an equivalent network with more constraint, without losing any feasible solution, in such case, we are giving a tighter description of the problem, and we obtain a network with a smaller number of consistent partial assignments. Already known constraints may implies additional (unary or binary) constraints.

**Definition 17** Let  $\gamma = (V, D, C)$  and  $\gamma' = (V, D', C')$  to be two constraint networks, sharing the same set of variables. We say that  $\gamma$  are **equivalent** to  $\gamma'$ , and we denote

$$\gamma \equiv \gamma'$$

if every solution (assignment of variables) of  $\gamma$  is a solution of  $\gamma'$  and vice versa.

**Definition 18** Let  $\gamma = (V, D, C)$  and  $\gamma' = (V, D', C')$  to be two constraint networks, sharing the same set of variables. We say that  $\gamma'$  is **tighter** than  $\gamma$  and we denote

$$\gamma' \sqsubseteq \gamma$$

if

- $\forall v \in V, D'_v \subseteq D_v$
- $\forall u, v \text{ such that } u \neq v \text{ either } C_{uv} \notin C \text{ or } C'_{uv} \subseteq C_{uv}$ .

If at least one of these inclusions is strict, we say that  $\gamma'$  is **strictly tighter** than  $\gamma$ :

$$\gamma' \sqsubset \gamma.$$

**Theorem 4** Let  $\gamma$  and  $\gamma'$  to be two constraint networks, if

- $\gamma' \equiv \gamma$
- $\gamma' \sqsubset \gamma$

then  $\gamma'$  has the same solutions as  $\gamma$  but fewer consistent partial assignments than  $\gamma$ . In such case,  $\gamma'$  is a **better encoding** of the problem.

We define *inference* the procedure to derive a tighter equivalent network. We want to use it to define an algorithm that finds a solution for a constraint network. We will incorporate inference in the backtracking algorithm, at every recursive call, a complex inference procedure leads to

- a smaller number of search nodes
- larger runtime needed at each node

We encode partial assignments as unary constraints

- if  $a(v) = d$
- we set the constraint  $D_v = \{d\}$

#### Algorithm 21 BacktrackingWithInference

```

Require:  $\gamma = (V, D, C), a$ 
  if  $a$  is inconsistent then
    return inconsistent
  end if
  if  $a$  is a total assignment then
    return  $a$ 
  end if
   $\gamma' = \text{copy of } \gamma$ 
   $\gamma' = \text{Inference}(\gamma')$ 
  If  $\exists v : D'_v = \emptyset$  then return inconsistent
  let  $V'$  to be the domain of  $a$ 
  select some  $v \in V - V'$  (select a variable for which  $a$  is not defined)
  for each  $d \in D'_v$  in some order do
     $a' = a \cup \{v = d\}$ 
     $D'_v = \{d\}$ 
     $a'' = \text{NaiveBacktracking}(\gamma', a')$ 
    if  $a''$  is not inconsistent then
      return  $a''$ 
    end if
  end for
  return inconsistent

```

With the function **Inference** we denote any procedure deriving a tighter equivalent network. One simple inference procedure called *forward checking* is described in algorithm 22.

The forward checking algorithm tightens the constraints without ruling out any solutions, it guarantees to deliver an equivalent network. The computation can be done incrementally, instead of performing the first loop, we may run only the second loop every time a new assignment  $a(v) = d'$  is added.

- The forward checking procedure is cheap and useful
- In rare situations is not a good idea to use it, there are stronger inference methods, but in many cases, forward checking is the best choice.

---

**Algorithm 22** Forward Checking

---

**Require:**  $\gamma, a$

```

for each  $v$  where  $a(v) = d'$  is defined do
    for each  $u$  where  $a(u)$  is undefined and  $C_{uv} \in C$  do
         $D_u = \{d \mid d \in D_u, (d, d') \in C_{uv}\}$ 
    end for
end for
return  $\gamma$ 

```

---

### 3.3.1 Arc Consistency for Stronger Inference

We now describe a properties between two variables that can be used to describe a stronger inference method than the forward checking.

**Definition 19** Let  $\gamma = (V, D, C)$  to be a constraint network, a variable  $u \in V$  is **arc consistent** to another variable  $v \in V$  if

- either  $C_{uv} \notin C$
- or for every  $d \in D_u$  there exists  $d' \in D_v$  such that  $(d, d') \in C_{uv}$ .

When two variables in a constraint network are arc consistent, it means that the domain of each variable contains at least one value that is compatible with at least one value in the domain of the other variable, according to the constraint between them.

As an example, let's say the variable  $v$  describes the color of your shirt, that can be red, blue, yellow or green, and the variable  $u$  define the color of your pants, that can be black, white, gray or brown. The constraint is that the shirt and the pants cannot be the same color, the variable  $v$  is arc consistent to  $u$  because for every shirt color you can find a pant color and vice versa.

**Note:** Arc consistency is a non commutative relation, if  $v$  is arc consistent to  $u$ , it's not implied directly that  $u$  is arc consistent to  $v$ .

**Definition 20** Let  $\gamma = (V, D, C)$  to be a constraint network, if every variable  $u \in V$  is arc consistent relative to every other variable  $v \in V$ , then we say that the network  $\gamma$  is **arc consistent**.

We can define an inference procedure that aims to make the network  $\gamma$  arc consistent, by removing variable domain values. Let's say that  $AC(\gamma)$  is a procedure that makes  $\gamma$  arc consistent, in such case:

- is guaranteed that  $AC(\gamma)$  returns an equivalent network.
- $AC(\gamma)$  is more powerful (more general) tha forward checking, in details:

$$AC(\gamma) \sqsubseteq \text{ForwardChecking}(\gamma) \quad (3.10)$$

We define the following subroutine:

---

**Algorithm 23** Revise

---

**Require:**  $\gamma, u \in V, v \in V$

```

for each  $d \in D_u$  do
    if  $\neg \exists d' \in D_v$  such that  $(d, d') \in C_{uv}$  then
         $D_u = D_u \setminus \{d\}$ 
    end if
end for
return  $\gamma$ 

```

---

We apply pairwise Revisions in algorithm 24 and define the first version of the inference procedure that makes a constraint network  $\gamma$  arc consistent.

It's not hard to show that this enforce arc consistency. If there are  $n$  variables,  $m$  constraints and  $k$  is the cardinality of the biggest domain, then the time complexity is  $O(mk^2 \cdot nk)$ , this algorithm performs

---

**Algorithm 24 AC\_1**

---

```

Require:  $\gamma$  changes = False
  while changes = False do
    for each  $C_{uv} \in C$  do
      If  $D_u$  after applying Revise( $\gamma, u, v$  reduces) then changes = True
      If  $D_v$  after applying Revise( $\gamma, v, u$  reduces) then changes = True
    end for
  end while

```

---



---

**Algorithm 25 AC\_3**

---

```

Require:  $\gamma$ 
   $M = \emptyset$ 
  for each  $C_{uv} \in C$  do
     $M = M \cup \{(u, v), (v, u)\}$ 
  end for
  while  $M \neq \emptyset$  do
    remove any element  $(u, v)$  from  $M$ 
    Revise( $\gamma, u, v$ )
    if  $D_u$  has changed in the call Revise then
      for each  $C_{wu} \in C$  where  $w \neq v$  do
         $M = M \cup \{(w, u)\}$ 
      end for
    end if
  end while

```

---

redundant computation since  $u$  and  $v$  are revised even if their domains haven't changed since the last time.

There is another version of the arc consistency procedure, described in algorithm 25. At any moment during the while loop, if  $(u, v) \notin M$  then  $u$  is arc consistent to  $v$ .

In the algorithm, we check the constraints where  $w \neq v$  because  $v$  is the reason why  $D_u$  just changed, if  $v$  was arc consistent to  $u$  before, that continues to hold, the values just removed from  $D_u$  did not match any values from  $D_v$  anyway.

**Theorem 5** Let  $\gamma$  to be a constraint network with  $m$  constraint and maximal domain size  $k$ . The algorithm AC3 showed in 25 have a time complexity in  $O(mk^3)$ .

*Proof:* Each Revise call takes  $O(k^2)$ , it is sufficient to prove that at most  $O(mk)$  calls to revise are made. The number of calls to Revise is the number of iterations of the while-loop, which is at most the number of insertions into  $M$ .

- Consider any constraint  $C_{uv}$
- two variables pairs corresponding to the constraint  $C_{uv}$  are inserted into  $M$  in the first for loop
- then beforehand the domain of either  $u$  or  $v$  was reduced, which happens at most  $2k$  times
- thus we have  $O(k)$  insertions per constraint, and  $O(mk)$  insertions overall. ■

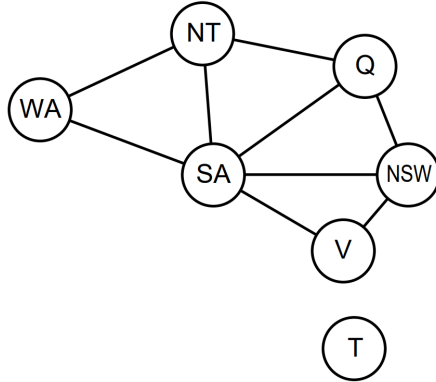
## 3.4 Decomposition

If  $\gamma$  is a network with  $n$  variables and maximal domain size  $k$ , we may search a feasible assignment by checking among all the possible ones, this will require to check  $O(k^n)$  assignments in the worst case. We saw how the inference help to shrink the search space, another useful methods is called *decomposition*.

**decomposition:** exploit the structure of a network to decompose it into smaller (and easier to solve) independent networks.

**Definition 21** Given a constraint network  $\gamma = (V, D, C)$ , the **constraint graph** of  $\gamma$  is the undirected graph, where the vertices are the variables  $V$  and there are an edge  $(u, v)$  if  $C_{uv} \in C$ .

For example, the graph for the "coloring Australia" problem shown in figure 3.1 is the following:



Note how this graph is composed by two connected sub-graphs, since the variable  $T$  (Tasmania) is disconnected from the other variables. The idea is to separate a network in function of his connected sub-graphs.

**Theorem 6** Let  $\gamma = (V, D, C)$  to be a constraint network. Let  $G$  to be the constraint graph, we denote  $\mathcal{V}(G)$  the vertices of  $G$  and  $\mathcal{E}(G)$  the edges of  $G$ , we know that  $\mathcal{V}(G) = V$ . We consider a partition of  $\mathcal{V}(G)$

$$\mathcal{V}(G) = \bigcup_i V_i \quad (3.11)$$

such that each  $V_i$  is connected and

$$\begin{cases} v \in V_i \\ u \in V_j \\ V_i \neq V_j \end{cases} \implies \begin{cases} (u, v) \notin \mathcal{E}(G) \\ (v, u) \notin \mathcal{E}(G) \end{cases} \implies C_{uv} \notin C$$

so the components of the partition are disconnected from each other. Let  $a_i$  to be an feasible assignment (solution) for the variables  $V_i$ , then

$$a = \bigcup_i a_i \quad (3.12)$$

is a solution to  $\gamma$ .

The theorem states that if two parts of  $\gamma$  are not connected (in term of his constraint graph), then they are independent and we can find separately a solution for each connected component.

**Theorem 7** Let  $\gamma = (V, D, C)$  to be a constraint network with  $n$  variables and maximal domain size  $k$ . Let  $G$  to be the constraint graph, if  $G$  is acyclic, we can find a solution for  $\gamma$  (or prove that  $\gamma$  is inconsistent) with a time complexity of  $O(nk^2)$ .

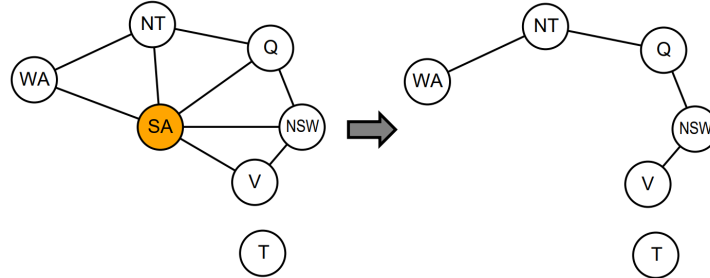
The procedure is the following (assuming that a constraint network is connected, if not, we can apply the procedure to each connected component).

1. We consider the constraint graph of  $\gamma$ , denoted  $G$ . We choose an arbitrary node  $v \in \mathcal{V}(G) = V$ , and we consider the BFS spanning tree  $G'$  starting from  $v$ .
2.  $G'$  is a directed graph, we consider a topological ordering  $v_1, \dots, v_n$  of the variables.
3. we iterate through the variables  $v_i$  in the opposite order respect to the topological ordering: for each  $i = n, n-1, n-2, \dots, 3, 2$  (except for  $v_1$ )
  - (a) we perform the procedure  $\text{Revise}(\gamma, \text{parent}(v_i), v_i)$
  - (b) if  $D_{\text{parent}(v_i)} \neq \emptyset$ , return inconsistent
- at this point, every variable is arc consistent relative to its children
4. We perform the algorithm 21 using the variable order  $v_1, v_2, \dots, v_n$ .

### 3.4.1 Cutsets

**Definition 22** Let  $\gamma = (V, D, C)$  to be a constraint network. Let  $G$  to be the constraint graph, a **cutset**  $V_0 \subseteq V$  is a set such that, the graph  $G'$  where  $\mathcal{V}(G') = V \setminus V_0$  is acyclic.

A cutset is a subset of variables removing which renders the constraint graph acyclic. The graph of the "coloring Australia" problem is *almost acyclic* since, removing *SA* will make the graph a tree.



The following procedure use cutsets to find a solution for  $\gamma$ .

```

 $V_0 :=$  a cutset; return CutsetConditioning( $\gamma, V_0, \emptyset$ )
function CutsetConditioning( $\gamma, V_0, a$ ) returns a solution, or "inconsistent"
   $\gamma' :=$  a copy of  $\gamma$ ;  $\gamma' :=$  ForwardChecking( $\gamma', a$ )
  if exists  $v$  with  $D'_v = \emptyset$  then return "inconsistent"
  if exists  $v \in V_0$  s.t.  $a(v)$  is undefined then select such  $v$ 
    else  $a' :=$  AcyclicCG( $\gamma'$ ); if  $a' \neq$  "inconsistent" then return  $a \cup a'$ 
    else return "inconsistent"
  for each  $d \in$  copy of  $D'_v$  in some order do
     $a' := a \cup \{v = d\}$ ;  $D'_v := \{d\}$ ;
     $a'' :=$  CutsetConditioning( $\gamma', V_0, a'$ )
    if  $a'' \neq$  "inconsistent" then return  $a''$ 
  return "inconsistent"

```

TODO: Riscrivere questo algoritmo, l'indentazione è ambigua

The forward checking operation is required to ensure that  $a \cup a'$  is consistent in  $\gamma$ , the runtime is exponential in  $V_0$ . It is true that finding an optimal cutset for a graph is NP-hard, but practical approximated procedures exists.

## CHAPTER

# 4

# LOGIC AND KNOWLEDGE REPRESENTATION

## 4.1 Propositional Logic

This chapter presents a brief summary of propositional logic. Let  $\Sigma$  to be a set of boolean variables, also called *Atoms*, the definition of a formula over the variables  $\Sigma$  is recursive.

**Definition 23** *Let  $\Sigma$  to be a set of atoms, then*

1.  $\top$  (True) and  $\perp$  (False) are  $\Sigma$ -formulas.
2. Each atom  $P \in \Sigma$  is a  $\Sigma$ -formula.
3. If  $\varphi$  is a formula<sup>1</sup>, then  $\neg\varphi$  is a formula.

If  $\varphi$  and  $\psi$  are formulas then:

1.  $\varphi \wedge \psi$  is a formula (*conjunction*)
2.  $\varphi \vee \psi$  is a formula (*disjunction*)
3.  $\varphi \rightarrow \psi$  is a formula (*implication*)
4.  $\varphi \leftrightarrow \psi$  is a formula (*equivalence*)

We recall that

$$\varphi \rightarrow \psi \equiv \neg\varphi \vee \psi \tag{4.1}$$

$$\varphi \leftrightarrow \psi \equiv (\varphi \wedge \psi) \vee (\neg\varphi \wedge \neg\psi) \tag{4.2}$$

**Definition 24** *An interpretation  $I$  over  $\Sigma$  is an assignment  $I : \Sigma \rightarrow \{0, 1\}$ .*

We say that  $I$  **satisfy** a formula  $\varphi$ , and we write  $I \models \varphi$  if the assignment makes the formula true. We can also say that  $I$  is a **model** of  $\varphi$  and the set of all the models of  $\varphi$  (all the assignments that satisfies  $\varphi$ ) is denoted  $M(\varphi)$ .

**Example:**

$$\varphi = (P \vee Q) \iff (R \vee S) \tag{4.3}$$

---

<sup>1</sup>We abbreviate "Σ-formula" with "formula" leaving it implicit that the formula is on the set of variables defined in the context.

the interpretation

$$I = \begin{cases} P = 1 \\ Q = 0 \\ R = 1 \\ S = 1 \end{cases} \quad (4.4)$$

is a model since  $I \models \varphi$ :

$$(P \vee Q) \leftrightarrow (R \vee S) \implies (1 \vee 0) \leftrightarrow (0 \vee 0) \implies 1 \leftrightarrow 1 \implies \top \quad (4.5)$$

A set of formulas  $KB$  is called **Knowledge Base** and an interpretation  $I$  models  $KB$  if, for all  $\varphi \in KB$  we have that  $I \models \varphi$ . We also say that  $I$  is a model of  $KB$ .

**Definition 25** Let  $\varphi$  to be a formula:

- $\varphi$  is **satisfiable** if  $\exists I$  such that  $I \models \varphi$
- $\varphi$  is **unsatisfiable** if is not satisfiable.
- $\varphi$  is **falsifiable** if there exists  $I$  that doesn't satisfy  $\varphi$ .
- $\varphi$  is a **tautology** if  $\forall I$ ,  $I \models \varphi$  (it's not falsifiable).

**Definition 26** Two formulas  $\varphi$  and  $\psi$  are **equivalent**

$$\varphi \equiv \psi$$

if  $M(\varphi) = M(\psi)$  ( $I$  models  $\varphi$  if and only if models also  $\psi$ ).

**Definition 27** Let  $\Sigma$  to be a set of atoms, a knowledge base  $KB$  over  $\Sigma$  **entails** a formula  $\varphi$

$$KB \models \varphi$$

if, all the models  $I$  of  $KB$ , are models of  $\varphi$ .

$$M\left(\bigwedge_{\psi \in KB} \psi\right) \subseteq M(\varphi)$$

We say that the formula  $\varphi$  follows from  $KB$ , and would be redundant to include  $\varphi$  in  $KB$ . It's also important to derive new logic formula form a given knowledge base.

**Theorem 8**  $KB \models \varphi$  if and only if  $KB \cup \{\neg\varphi\}$  is unsatisfiable.

*Proof*  $\implies$ : Let  $KB \models \varphi$  for any  $I \models KB$  we have  $I \models \varphi$  so  $I \not\models \neg\varphi$ .

*Proof*  $\impliedby$ : If  $KB \cup \{\neg\varphi\}$  is unsatisfiable then for any  $I$  such that  $I \models KB$  we have  $I \not\models \neg\varphi$  hence  $I \models \varphi$ .

Given a formula with  $n$  variables, the number of possible assignment is  $2^n$ , so it's hard to test the satisfiability of a formula by testing all the interpretations.

**Definition 28** A formula is in **conjunctive normal form (CNF)** if is a conjunction of disjunction of literals.<sup>2</sup>

$$\bigwedge_i \left( \bigvee_j L_{i,j} \right) \quad (4.6)$$

**Definition 29** A formula is in **disjunctive normal form (DNF)** if is a disjunction of conjunction of literals.

$$\bigvee_i \left( \bigwedge_j L_{i,j} \right) \quad (4.7)$$

<sup>2</sup>A literal is an atom  $P$  or the negation of an atom  $\neg P$ .

It is important to know that **every formula** can be expressed in CNF or DNF, or, for each formula  $\varphi$ , there exists an equivalent formula  $\psi$  that is a CNF or a DNF. It is possible to construct the equivalent CNF (or DNF) from  $\varphi$  by exploiting the equivalences such as

- $\varphi \rightarrow \psi \equiv \neg\varphi \vee \psi$
- $\neg(\varphi \vee \psi) \equiv \neg\varphi \wedge \neg\psi$
- and so on...

$$\begin{array}{ll}
 ((P \vee H) \wedge \neg H) \rightarrow P & \text{Eliminate "}" \rightarrow "} \\
 \neg((P \vee H) \wedge \neg H) \vee P & \text{Move "}" \neg " \text{ inwards} \\
 (\neg(P \vee H) \vee H) \vee P & \text{Move "}" \neg " \text{ inwards} \\
 ((\neg P \wedge \neg H) \vee H) \vee P & \text{Distribute "}" \vee " \text{ over "}" \wedge "} \\
 ((\neg P \vee H) \wedge (\neg H \vee H)) \vee P & \text{Distribute "}" \vee " \text{ over "}" \wedge "} \\
 (\neg P \vee H \vee P) \wedge (\neg H \vee H \vee P) &
 \end{array}$$

### 4.1.1 Resolution

We can construct a simple algorithm that, given a CNF formula  $\psi$ , tests if is satisfiable or not, by trying to derive from  $\psi$  new formulas: If it finds a derived formula  $\psi$  that is unsatisfiable then  $\psi$  is unsatisfiable.

We use a method called *Calculus* that, given a single *rule* (we will define a rule later), allows us to derive new formulas from a given one. If  $\varphi$  is derived from  $\psi$ , we write

$$\psi \vdash \varphi.$$

**Definition 30** A clause  $C$  is a disjunction of literals, like:

$$\neg P \vee Q \vee R \vee \neg Q \quad (4.8)$$

To shrink the notation, we identify  $C$  with the set of the literals included:

$$P \vee \neg Q \text{ became } C = \{P, \neg Q\} \quad (4.9)$$

Since a CNF is a conjunction of disjunction we can identify each CNF with a set of clauses, usually denoted  $\Delta$ . For example, the CNF

$$\Delta = \{\{P, \neg Q\}, \{R, Q\}\} \quad (4.10)$$

is

$$(P \vee \neg Q) \wedge (R \vee Q) \quad (4.11)$$

We denote the empty clause  $\square$ . An interpretation  $I$  satisfies a single clause  $C$

$$I \models C$$

if there exists a literal  $l \in C$  such that  $I \models l$ .  $I$  satisfies  $\Delta$

$$I \models \Delta$$

if, for all  $C \in \Delta$  we have that  $I \models C$ .

**Definition 31** A *inference rule* is a rule that derive a new formula from a given one, the **resolution rule** is the following:

$$\frac{C_1 \dot{\cup} \{l\}, C_2 \dot{\cup} \{\neg l\}}{C_1 \cup C_2} \quad (4.12)$$

You may wonder what this formula means and how it is applied, the symbol  $\dot{\cup}$  identify a disjoint union, so  $C_1 \dot{\cup} \{l\}$  is a union and is implied that  $l \notin C_1$ .  $l$  is a literal. The formula is written as a fraction:

- the term over the fraction line represents the clauses that we already have in our CNF
- the term under the fraction line represents a new formula that we derive

In practice, if we have two clauses  $C'_1 = C_1 \dot{\cup} \{l\}$  and  $C'_2 = C_2 \dot{\cup} \{\neg l\}$  in  $\Delta$ , we can derive a disjunction  $\phi = C_1 \cup C_2$ .

if you have two clauses where the same literal appears, but in one it is affirmative ( $l$ ) and in the other it is negated ( $\neg l$ ), you can "add them" by eliminating  $l$  and  $\neg l$  and merging everything else.

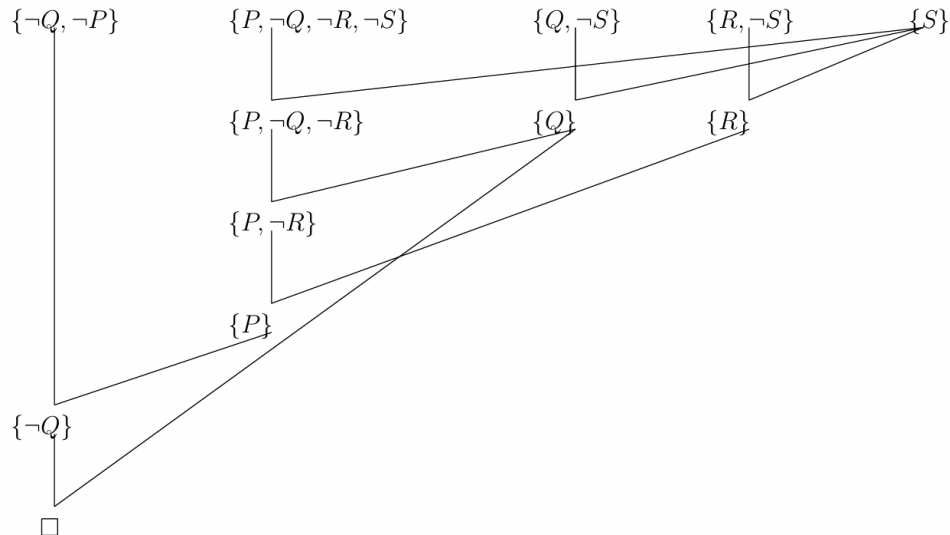
For example, if you have  $\{P, \neg R\}$  and  $\{R, Q\}$ , you can resolve them and find a new formula  $\{P, Q\}$ .

**Theorem 9** *The soundness theorem affirms that, if  $\Delta \vdash D$  (the formula  $D$  can be derived by some clause in  $\Delta$  with the resolution rule) then  $\Delta \models D$  (the CNF  $\Delta$  entails  $D$ ).*

In the definition of *entailment*, is a knowledge base that "entails" a formula, and not a formula that "entails" another formula, but a knowledge base  $KB$  is equivalent to a CNF by putting in conjunction all the formulas in  $KB$ .

The resolution algorithm takes as an input a knowledge base  $KB$  and a formula  $\phi$  and guess if  $KB \models \phi$  by the following steps:

1. transform  $KB$  in a CNF
2. write the CNF as a set of clauses  $\Delta$ , by adding to  $\Delta$  also the clause  $\{\neg\phi\}$
3. apply the resolution rule on  $\Delta$  until it's derived the empty clause  $\square$ .



It is important to notice that  $\Delta \models \varphi$  doesn't implies that  $\Delta \vdash \varphi$ , but, by the contradiction theorem, if we run the resolution on  $KB \cup \{\neg\varphi\}$  and we derive the empty clause  $\square$  ( $KB \cup \{\neg\varphi\}$  is unsatisfiable) then  $KB \models \varphi$ .

**Theorem 10** *A CNF  $\Delta$  is unsatisfiable if and only if  $\Delta \vdash \square$ .*

#### 4.1.2 The DPLL procedure

The SAT problem is to decide whenever a given propositional formula  $\varphi$  is satisfiable or not, as we well know, this is the most classic example of a problem NP-complete. The SAT problem is a "subset" of the constraint satisfaction problem:

- SAT can be viewed as a CSP problem in which all variable domains are Boolean, and the constraints have unbounded arity.

Every CSP problem can be encoded as a SAT problem:

- Given any constraint network  $\gamma$ , we can in low-order polynomial time construct a CNF formula  $\varphi(\gamma)$  that is satisfiable if and only if  $\gamma$  is solvable.

This and many other problems can be "encoded" as a SAT formula through reduction.

A **SAT solver** is an algorithm that, given a CNF  $\Delta$ , returns an interpretation  $I$  (if exists) such that  $I \models \Delta$ . We consider the DPLL procedure: A complete SAT solver. A SAT solver can be used in different contexts:

- We can use it to entail a new formula  $\varphi$  from  $\Delta$ .
- We need an assignment  $I$  that models  $\Delta$ .

---

**Algorithm 26** Davis-Putnam-Logemann-Loveland (DPLL)

---

```

1: function DPLL( $\Delta, I$ )
2:   /* Unit Propagation (UP) Rule: */
3:    $\Delta' :=$  a copy of  $\Delta$ ;  $I' := I$ 
4:   while  $\Delta'$  contains a unit clause  $C = \{l\}$  do
5:     extend  $I'$  with the respective truth value for the proposition underlying  $l$ 
6:     /* remove false literals and true clauses */
7:     simplify  $\Delta'$ 
8:   end while
9:   /* Termination Test: */
10:  if  $\square \in \Delta'$  then
11:    return "unsatisfiable"
12:  end if
13:  if  $\Delta' = \{\}$  then
14:    return  $I'$ 
15:  end if
16:  /* Splitting Rule: */
17:  select some proposition  $P$  for which  $I'$  is not defined
18:   $I'' := I'$  extended with one truth value for  $P$ ;  $\Delta'' :=$  a copy of  $\Delta'$ ; simplify  $\Delta''$ 
19:  if  $I''' :=$  DPLL( $\Delta'', I''$ )  $\neq$  "unsatisfiable" then
20:    return  $I'''$ 
21:  end if
22:   $I'' := I'$  extended with the other truth value for  $P$ ;  $\Delta'' := \Delta'$ ; simplify  $\Delta''$ 
23:  return DPLL( $\Delta'', I''$ )
24: end function

```

---

The Unit Propagation (UP) section of the algorithm correspond to the application of the following rule.

**Definition 32** *The unit resolution is the inference rule:*

$$\frac{C \cup \{\neg l\}, \{l\}}{C} \quad (4.13)$$

That is, if  $\Delta$  contains parent clauses of the form  $C \cup \{\neg l\}$  and  $\{l\}$  the rule allows to add the resolvent clause  $C$ .

Does the UP rule have the soundness property of Theorem 9? Yes, we have to show that if  $\Delta'$  can be derived from  $\Delta$  then  $\Delta \models \Delta'$ , this is true since any derivation made by unit resolution can also be done by the full resolution, which we already know has this property.

Is the UP rule complete? To be complete, we have to show that, if  $\Delta \models \Delta'$ , then  $\Delta'$  can be derived from  $\Delta$  by the UP rule, this is false, a counter example is given by the following CNF

$$\Delta = \{\{P, Q\}, \{P, \neg Q\}, \{\neg P, Q\}, \{\neg P, \neg Q\}\}$$

is unsatisfiable but the UP rule cannot derive the empty clause  $\square$ .

### 4.1.3 Conflict Analysis and Clause Learning

The DPLL is a systematic way of searching for a resolution proof. We define the *number of decisions* of a DPLL run as the total number of times a truth value was set by either unit propagation or the splitting rule, the following relationship between the DPLL algorithm and the resolution procedure holds:

If DPLL returns "unsatisfiable" on  $\Delta$ , then  $\Delta \models \square$  with a resolution derivation whose length is at most the number of decisions.

So, DPLL is an effective practical method for conducting resolution proofs. The DPLL is a "tree resolution", this is a problem since there are some  $\Delta$  whose shortest tree resolution proof is exponentially longer than their shortest (general) resolution proof. In tree resolution, every time a clause is needed, it must be derived from scratch because results aren't stored, this makes DPLL "make the same mistakes over and over again", leading to proofs that are exponentially longer than they need to be. Modern SAT solvers use Clause Learning (CDCL). When the solver hits a conflict, it analyzes the cause, creates a new clause to "remember" that mistake, and adds it to the database so it never explores that specific failing path again.

During the execution of the DPLL procedure, there are many branches of the recursive run, where for each branch we compute a partial interpretation, a literal during a branch can be

- *choice literals*: the value of an atom  $P$  is set by the splitting rule, assigning  $P = 0$  or  $P = 1$ .
- *implied literal*: the value is set automatically by the UP rule.
- *conflict literal*: when the UP rule derive an empty clause  $\square$ , the branch became unsatisfiable.

**Definition 33** The *implication graph* for a branch  $\beta$  is a directed graph  $G^{impl}$  such that:

- the vertices  $V(G^{impl})$  are the choice literals and implied literals of the branch, plus a special vertex  $\square_C$  for each clause  $C$  that becomes empty.
- the edges  $E(G^{impl})$  are the following:
  - when a clause  $C = \{l_1 \dots l_k, l'\} \in \Delta$  becomes unit with the implied literal  $l'$ , there will be the edges

$$\neg l_1 \rightarrow l' \tag{4.14}$$

$$\vdots \tag{4.15}$$

$$\neg l_k \rightarrow l' \tag{4.16}$$

- when a clause  $C = \{l_1 \dots l_k\}$  becomes empty there will be the edges

$$\neg l_1 \rightarrow \square_C \tag{4.17}$$

$$\vdots \tag{4.18}$$

$$\neg l_k \rightarrow \square_C \tag{4.19}$$

#### Example

Consider

$$\Delta = \{\{P, Q, \neg R\}, \{\neg P, \neg Q\}, \{R\}, \{P, \neg Q\}\}$$

1. First step: Apply the UP rule on  $R$  with the clause  $\{P, Q, \neg R\}$ , and get the set:

$$\{\{P, Q, \neg R\}, \{\neg P, \neg Q\}, \{R\}, \{P, \neg Q\}\} \text{ becomes } \{\{P, Q\}, \{\neg P, \neg Q\}, \{P, \neg Q\}\} \tag{4.20}$$

the literal  $R$  is an implied literal and becomes a vertex in the graph.

2. Second step: We apply the splitting rule on  $P$ , creating two branches, one with  $P = 0$  and the other with  $P = 1$ , we continue with the branch with  $P = 0$ . The literal  $\neg P$  becomes a choice literal so there are the vertex  $\neg P$  in the graph.

$$\{\{P, Q\}, \{\neg P, \neg Q\}, \{P, \neg Q\}\} \text{ becomes } \{\{Q\}, \{\neg Q\}\} \tag{4.21}$$

3. We apply the UP rule on  $Q$  and  $Q$  becomes an implied literal, the edges

$$R \rightarrow Q \quad (4.22)$$

$$\neg P \rightarrow Q \quad (4.23)$$

are added to the graph. After the UP rule it remains the empty clause

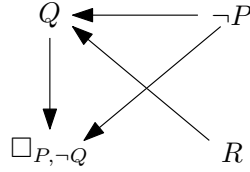
$$\{\square\}$$

So we add the conflict literal  $\square_{P,\neg Q}$  and the edges

$$Q \rightarrow \square_{P,\neg Q} \quad (4.24)$$

$$\neg P \rightarrow \square_{P,\neg Q} \quad (4.25)$$

the final graph of this branch is:



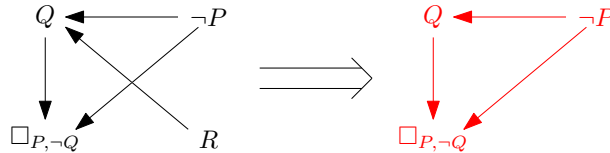
An implication graph cannot have cycle because it keeps track of chronological behavior along the current DPLL search branch  $\beta$ . Unit propagation cannot derive  $l'$  whose value was already set beforehand. The implication graph it is used to visualize and trace the logical path that led the algorithm to assign certain truth values to the variables.

There is another tool used to capture "what went wrong" in a failed node (when a branch return "unsatisfiable").

**Definition 34** Let  $\Delta$  to be a set of clauses and  $G^{impl}$  the implication graph relative to a branch of the DPLL algorithm, a **conflict graph**  $G^{conf}$  is a sub-graph of  $G^{impl}$  such that:

- $G^{conf}$  contains exactly one conflict vertex  $\square_C$
- if  $l'$  is a vertex in  $G^{conf}$ , then, all the parents of  $l'$  that are vertices  $\neg l_i$  with the arc  $(\neg l_i, l') \in E(G^{impl})$  are also vertices in  $G^{conf}$ .
- all the vertices in  $G^{conf}$  have a path to  $\square_C$ ,

We can construct the conflict graph by starting from a conflict vertex and going back through the implication graph until reaching choice literals.



Let  $choiceList(G^{conf})$  to be the set of all the choice literals in the conflict graph, given  $\Delta$ , if we set as true<sup>3</sup> all the literals in  $choiceList(G^{conf})$  we will have a false statement:

$$\Delta \models \left( \bigwedge_{l \in choiceList(G^{conf})} l \right) \rightarrow \perp \quad (4.26)$$

This can be rewritten as

$$\Delta \models \bigwedge_{l \in choiceList(G^{conf}) - l} l \quad (4.27)$$

<sup>3</sup>We recall that a literal  $l$  can be an atom or a negated atom. If  $l = P$  where  $P$  is an atom, setting  $l$  to true means  $l = P = 1$ , if  $l = \neg P$ , setting  $l$  to true means  $l = 1 = \neg P \implies P = 0$ .

**Proposition 3 (Clause Learning)** Let  $\Delta$  to be a set of clauses and let  $G^{conf}$  be a conflict graph at some time point during a run of DPLL on  $\Delta$ . Let  $choiceList(G^{conf})$  to be the set of all the choice literals in the conflict graph, then

$$\Delta \models \{\neg l \mid l \in choiceList(G^{conf})\} \quad (4.28)$$

The negation of the choice literals in a conflict graph is a valid clause.

So during the execution of the DPLL we can analyze a conflict graph and derive a new clause

$$C = \{\neg l \mid l \in choiceList(G^{conf})\} \quad (4.29)$$

we can add  $C$  into  $\Delta$  and retract the last choice literal  $l'$ .

### Example

Let

$$\Delta = \{\{\neg A, \neg B\}, \{\neg A, B\}\}$$

we apply the DPLL and set in a branch  $A = 1$  so

$$\{\{\neg B\}, \{B\}\}$$

we have a conflict and the new clause derived by negating the choice literals is

$$C = \{\neg A\} \quad (4.30)$$

Now we add this to the set of clauses and we have

$$\Delta = \{\{\neg A, \neg B\}, \{\neg A, B\}, \{\neg A\}\}$$

Now the algorithm instantly knows that  $A$  must be False and will no longer waste time proving  $A = 1$ .

After adding the new clause  $C$ , the process follows three fundamental logical phases to avoid repeating the same mistake:

1. The algorithm retracts the last choice made, denoted as  $l'$  (the truth value assigned to  $l'$  is removed and it becomes an unassigned literal again). This is necessary because the combination of previous choices  $(l_1, \dots, l_k)$  together with  $l'$  has just been shown to be inconsistent.
2. once  $C$  is added and the  $l'$  choice is cancelled, the  $C$  clause becomes a unit clause (only one unassigned literal remains). In this case,  $\neg l'$  is no longer a "free choice" of the algorithm, but becomes a literal implied by Unit Propagation (UP).
3. The algorithm reruns Unit Propagation and continues the analysis. If a new conflict were to occur, the learned clause will be even more "powerful" because it will only contain the even older choices  $(l_1, \dots, l_k)$ , allowing you to move up the decision hierarchy.

We saw how the original DPLL algorithm generates proof exponentially longer than their shortest (general) resolution proof. This is not true with clause learning, that renders DPLL equivalent to full resolution.

## 4.2 Predicate Logic (FOL)

A logic is a family of formal languages, which has the purpose of *representing* information and *manipulate* knowledge. Each logic is provided with a **syntax** and a **semantics**, the syntax defines a series of symbols and the structure that the formulas must take on to be considered valid, the semantics defines the meaning of these formulas.

In classical logic, each formula, based on the "world" to which it is applied, can be true or false. To define the syntax I must establish:

- What symbols can I use (the alphabet)
- Which finite sequences of symbols can compose a formula

Semantics, on the other hand, establishes the *truth* of formulas in the so-called possible "worlds", i.e the **interpretations**. In first-order logic, also called **FOL**, one establishes

- Syntax
  - Interpretation
  - Variable assignment
  - Model
  - Evaluating a formula
- Satisfaction
- Unsatisfiability
- Validity

### 4.2.1 Syntax

In a FOL formula there are:

- Variables:  $x, x_1, x_2, x', x'' \dots$
- True ( $\top$ ) or False ( $\perp$ ) symbols
- Operators:  $\neg, \wedge, \vee, \rightarrow, \leftrightarrow$
- Quantifiers:  $\forall, \exists$

We can have constant symbols that represents an object

$$\text{BlockA}, \text{BlockB}, a, b \quad (4.31)$$

predicate symbols that returns true or false taking an object as an input:

$$\text{Block}(.), \text{Above}(.,.)) \quad (4.32)$$

The arity (number of argument) can be  $\geq 1$ . For example:

$$\text{Chase}(\text{Cat}, \text{Mouse}) = \top \quad (4.33)$$

Means that the cat chase the mouse. Then we have the function symbols:

$$\text{WeightOf}(.), \text{Succ}(.), \text{Sum}(.,.) \quad (4.34)$$

these functions returns an object and not a truth value

$$\text{Sum}(3, 4) = 7 \quad (4.35)$$

where 3, 4 and 7 are constant symbols. Constant symbols are just function symbols of arity zero.

**Definition 35 (Signature)** A signature  $\Sigma$  in predicate logic is a finite set of constant symbols, predicate symbols, and function symbols.

Another example of formula is

$$\forall x[\text{Dog}(x) \rightarrow \exists y\text{Chase}(x, y)] \quad (4.36)$$

Means: For every  $x$  such that  $x$  is a dog, there exists something ( $y$ ) such that  $x$  chase  $y$ , in a few words: Every dog chases something.

**Definition 36 (Term)** Let  $\Sigma$  to be a signature:

- every variable and every constant symbol is a term.
- if  $t_1 \dots, t_n$  are terms and  $f$  is a  $n$ -ary function symbol, then  $f(t_1 \dots, t_n)$  is a term.

**Definition 37 (Atoms)** Let  $\Sigma$  to be a signature:

- $\top$  and  $\perp$  are atoms
- if  $t_1, \dots, t_n$  are terms and  $P$  is a  $n$ -ary predicate symbol, then  $P(t_1, \dots, t_n)$  is an atom.

**Definition 38 (Formula)** Let  $\Sigma$  to be a signature:

- each atom is a formula.
- if  $\varphi$  is a formula,  $\neg\varphi$  is a formula
- if  $\varphi$  and  $\psi$  are formulas then
  - $\varphi \wedge \psi$  is a formula (conjunction)
  - $\varphi \vee \psi$  is a formula (disjunction)
  - $\varphi \rightarrow \psi$  is a formula (implication)
  - $\varphi \leftrightarrow \psi$  is a formula (equivalence)
- if  $\varphi$  is a formula and  $x$  a variable then
  - $\forall x\varphi$  is a formula
  - $\exists x\varphi$  is a formula

Generally:

- terms represents object
- predicates represents relations on the universe

**Definition 39 (Interpretation)** Let  $\Sigma$  be a signature. A  $\Sigma$ -interpretation is a pair  $(U, I)$  where  $U$ , the universe, is an arbitrary non-empty set

$$U = \{o_1, o_2, \dots\}$$

and  $I$  is a function, notated as superscript, so that:

1.  $I$  maps constant symbols to elements of  $U$ :  $c^I \in U$

$$\text{Lassie}^I = o_1$$

2.  $I$  maps  $n$ -ary predicate symbols to  $n$ -ary relations over  $U$ :  $P^I \subseteq U^n$

$$\text{Dog}^I = \{o_1, o_3\}$$

3.  $I$  maps  $n$ -ary function symbols to  $n$ -ary functions over  $U$ :  $f^I \in [U^n \mapsto U]$

$$\text{FoodOf}^I = \{(o_1 \mapsto o_4), (o_2 \mapsto o_5), \dots\}$$

We will often refer to  $I$  as the interpretation, omitting  $U$ . Note that  $U$  may be infinite.

**Definition 40 (Ground Term Interpretation)** The interpretation of a ground term under  $I$  is

$$(f(t_1, \dots, t_n))^I = f^I(t_1^I, \dots, t_n^I)$$

**Definition 41 (Ground Atom Satisfaction)** Let  $\Sigma$  be a signature and  $I$  a  $\Sigma$ -interpretation. We say that  $I$  satisfies a ground atom  $P(t_1, \dots, t_n)$ , written  $I \models P(t_1, \dots, t_n)$ , iff  $(t_1^I, \dots, t_n^I) \in P^I$ . We also call  $I$  a model of  $P(t_1, \dots, t_n)$ .

$$I \models \text{Dog}(\text{Lassie}) \text{ because } \text{Lassie}^I = o_1 \in \text{Dog}^I$$

**Example "Integers":**

$$\begin{aligned} U &= \{1, 2, 3, \dots\} \\ 1^I &= 1, 2^I = 2, 3^I = 3, \dots \\ Even^I &= \{2, 3, 4, 6, \dots\} \\ Equals^I &= \{\langle 1, 1 \rangle, \langle 2, 2 \rangle, \dots\} \\ Succ^I &= \{(1 \mapsto 2), (2 \mapsto 3), \dots\} \end{aligned}$$

**Definition 42 (Variable Assignment)** Let  $\Sigma$  be a signature and  $(U, I)$  an interpretation. Let  $X$  be the set of all variables. A variable assignment  $\alpha$  is a function  $\alpha : X \mapsto U$ .

**Definition 43 (Term Interpretation)** The interpretation of a term under  $I$  and  $\alpha$  is:

1.  $x^{I,\alpha} = \alpha(x) \quad [x^{I,\alpha} = o_1]$
2.  $c^{I,\alpha} = c^I \quad [Lassie^{I,\alpha} = Lassie^I]$
3.  $(f(t_1, \dots, t_n))^{I,\alpha} = f^I(t_1^{I,\alpha}, \dots, t_n^{I,\alpha})$

**Definition 44 (Atom Satisfaction)** Let  $\Sigma$  be a signature,  $I$  a  $\Sigma$ -interpretation, and  $\alpha$  a variable assignment. We say that  $I$  and  $\alpha$  satisfy an atom  $P(t_1, \dots, t_n)$ , written  $I, \alpha \models P(t_1, \dots, t_n)$ , if and only if  $(t_1^{I,\alpha}, \dots, t_n^{I,\alpha}) \in P^I$ . We also call  $I$  and  $\alpha$  a model of  $P(t_1, \dots, t_n)$ .

From now, we write  $\alpha \frac{x}{o}$  to overwrite  $x$  with  $o$  in  $\alpha$ .

**Definition 45 (Formula Satisfaction)** Let  $\Sigma$  be a signature,  $I$  a  $\Sigma$ -interpretation, and  $\alpha$  a variable assignment. We set:

$$\begin{aligned} I, \alpha \models \top &\quad \text{and} \quad I, \alpha \not\models \perp \\ I, \alpha \models \neg\varphi &\quad \text{iff} \quad I, \alpha \not\models \varphi \\ I, \alpha \models \varphi \wedge \psi &\quad \text{iff} \quad I, \alpha \models \varphi \text{ and } I, \alpha \models \psi \\ I, \alpha \models \varphi \vee \psi &\quad \text{iff} \quad I, \alpha \models \varphi \text{ or } I, \alpha \models \psi \\ I, \alpha \models \varphi \rightarrow \psi &\quad \text{iff} \quad \text{if } I, \alpha \models \varphi, \text{ then } I, \alpha \models \psi \\ I, \alpha \models \varphi \leftrightarrow \psi &\quad \text{iff} \quad I, \alpha \models \varphi \text{ if and only if } I, \alpha \models \psi \\ I, \alpha \models \forall x \varphi &\quad \text{iff} \quad \text{for all } o \in U \text{ we have } I, \alpha \frac{x}{o} \models \varphi \\ I, \alpha \models \exists x \varphi &\quad \text{iff} \quad \text{there exists } o \in U \text{ so that } I, \alpha \frac{x}{o} \models \varphi \end{aligned}$$

If  $I, \alpha \models \varphi$ , we say that  $I$  and  $\alpha$  satisfy  $\varphi$  (are a model of  $\varphi$ ).

Like the propositional logic, a formula can be

- satisfiable if there exist  $I, \alpha$  that satisfy  $\varphi$ .
- unsatisfiable if  $\varphi$  is not satisfiable.
- falsifiable if there exist  $I, \alpha$  that do not satisfy  $\varphi$ .
- valid if  $I, \alpha \models \varphi$  holds for all  $I$  and  $\alpha$ . We also call  $\varphi$  a tautology.

**Definition 46** Given two formulas  $\varphi, \psi$ , we say that  $\varphi$  **entails**  $\psi$

$$\varphi \models \psi$$

if every model of  $\varphi$  is also a model of  $\psi$ . If

$$\varphi \models \psi \wedge \psi \models \varphi$$

then the two formulas are **equivalent**:

$$\varphi \equiv \psi$$

Let's consider the formula:

$$\varphi = \forall x[R(x, y)] \tag{4.37}$$

the variable  $y$  is called *free*, so the formula is not closed and cannot be evaluated in a truth value. We define the **Knowledge Base** as a set  $KB$  of closed formulas.

### 4.2.2 Normal Forms

It is convenient to express the FOL formulas in normal forms to describe the problem.

The **Prenex normal form** move all the quantifiers at the beginning of the formula, the following rules must be applied (STEP 1):

1. Eliminate  $\leftrightarrow$ :  $(\varphi \leftrightarrow \psi) \equiv [(\varphi \rightarrow \psi) \wedge (\psi \rightarrow \varphi)]$
2. Eliminate  $\rightarrow$ :  $(\varphi \rightarrow \psi) \equiv (\neg\varphi \vee \psi)$
3. Apply De Morgan:
  - $\neg(\varphi \wedge \psi) \equiv (\neg\varphi \vee \neg\psi)$
  - $\neg(\varphi \vee \psi) \equiv (\neg\varphi \wedge \neg\psi)$
4. Move  $\neg$  inwards:
  - $\neg\forall x\varphi \equiv \exists x\neg\varphi$
  - $\neg\exists x\varphi \equiv \forall x\neg\varphi$

Let's see an example of application:

$$\begin{aligned} \neg\forall x[(\forall xP(x)) \rightarrow R(x)] &\equiv \\ \neg\forall x[\neg(\forall xP(x)) \vee R(x)] &\equiv \\ \exists x\neg[\neg(\forall xP(x)) \vee R(x)] &\equiv \\ \exists x[(\forall xP(x)) \wedge \neg R(x)] \end{aligned}$$

The general form of a Prenex normal form formula is

$$Qx_1Qx_2 \dots Qx_n\varphi \quad (4.38)$$

where  $Q \in \{\forall, \exists\}$  is a generic quantifier. After applying the first set of rules, the following must be applied (STEP 2):

- if  $x$  is not free in  $\psi$ :
 
$$\begin{aligned} (\forall x\varphi) \wedge \psi &\equiv \forall x(\varphi \wedge \psi) \\ (\forall x\varphi) \vee \psi &\equiv \forall x(\varphi \vee \psi) \\ (\exists x\varphi) \wedge \psi &\equiv \exists x(\varphi \wedge \psi) \\ (\exists x\varphi) \vee \psi &\equiv \exists x(\varphi \vee \psi) \end{aligned}$$

**Definition 47** If  $x$  is a variable,  $t$  a term and  $\varphi$  a formula, then the **instantiation** of  $x$  with  $t$  in  $\varphi$ , written  $\varphi_t^x$ , replace all free appearances of  $x$  in  $\varphi$  by  $t$ . If  $t = y$  is a variable, then  $\varphi_y^x$  just renames the variable  $x$  with  $y$ .

**Lemma 3** if  $y$  is a variable that doesn't appears in  $\varphi$ , then

$$\begin{aligned} \forall x\varphi &\equiv \forall y\varphi_y^x \\ \exists x\varphi &\equiv \exists y\varphi_y^x \end{aligned}$$

This Lemma is useful to move outwards the variable like in the following example:

$$\begin{aligned} \exists x[(\forall xP(x)) \wedge \neg R(x)] &\equiv \\ \exists x[(\forall yP(y)) \wedge \neg R(x)] &\equiv \\ \exists x\forall y[P(y) \wedge \neg R(x)] \end{aligned}$$

**Theorem 11** There exists an algorithm that, for any FOL formula  $\varphi$ , calculates in polynomial time an equivalent formula in Prenex normal form.

The **Skolem normal form** is a formula (that is also in Prenex normal form) where each quantifier is universal:

$$\forall x_1 \forall x_2 \dots \forall x_n \varphi \quad (4.39)$$

Each formula in Prenex normal form can be easily transformed, consider the following theorem.

**Theorem 12 (Skolem)** *Let  $\varphi = \forall x_1 \forall x_2 \dots \forall x_k \exists y \psi$  to be a closed FOL formula in Prenex normal form such that, all the variables are pairwise distinct*

$$x_i \neq x_j, \forall i \neq j$$

let  $f$  to be a  $k$ -ary function symbol ( $k$  can also be zero i.e.  $f$  is constant) that doesn't appear in  $\varphi$ .  $\varphi$  is satisfiable if and only if the formula

$$\forall x_1 \forall x_2 \dots \forall x_k \psi \frac{y}{f(x_1 \dots x_k)} \quad (4.40)$$

Let's see an example of application of the theorem. We consider

$$\exists x \forall y \exists z R(x, y, z)$$

we remove  $\exists x$  using the theorem:

$$\exists x \forall y \exists z R(x, y, z) \equiv \exists x [\forall y \exists z R(x, y, z)] \equiv$$

Now the formula have the form  $\exists x \varphi$  so we can substitute  $x$  with a 0-arity function (a constant).

$$\forall y \exists z R(f, y, z)$$

By applying the theorem again we can substitute  $z$  with a function  $g$  of arity 1 in the variable  $y$ :

$$\forall y R(f, y, g(y))$$

In this way a Prenex formula can be transformed in a Skolem formula.

**Theorem 13** *There exists an algorithm that, for any closed FOL formula  $\varphi$ , efficiently calculates a Skolem normal form formula that is satisfiable if and only if  $\varphi$  is satisfiable.*

If  $\varphi$  is a FOL formula, their relative Skolem formula is denoted  $\varphi^*$ . The theorem says that:

*Satisfying existential quantifier for universally quantified variables  $x_1 \dots x_k$  is equivalent to choose a value for a function of  $x_1 \dots x_k$ .*

Like the Skolem normal form it is a specialization of the Prenex normal form, the **Clausal normal form** is a specialization of the Skolem normal form. A formula is in this normal form if has the following structure:

$$\forall x_1 \forall x_2 \dots \forall x_n (l_1 \vee \dots \vee l_n) \quad (4.41)$$

where  $l_i$  are literals that (may) depends on  $x_1 \dots x_n$  (the formula must be closed). Any Skolem formula can be transformed in a Clausal formula

1. consider  $\forall x_1 \forall x_2 \dots \forall x_n \varphi$
2. transform  $\varphi$  to a SAT-equivalent CNF formula  $\psi$
3. write a set of clauses, one for each disjunction in  $\psi$
4. renames the variables so that each occurs in at most one clause.

**Theorem 14** *There exists an algorithm that, for any closed FOL formula  $\varphi$ , efficiently calculates a formula in clausal normal form that is satisfiable if and only if  $\varphi$  is satisfiable.*

### 4.2.3 FOL Via Propositional Logic

In this section we consider a way to reason about FOL using propositional logic, in particular, we bring a FOL formula in SNF (Skolem normal form) and then we generate a *Herbrand expansion*<sup>4</sup> to use propositional reasoning.

For any finite set of FOL formulas  $\theta^*$ , we denote  $CF(\theta^*)$  the set of constant symbols and function occurring in  $\theta^*$ . If no constant symbols occurs in  $\theta^*$  we add a new "virtual" symbol  $c$  in  $CF(\theta^*)$ .

**Definition 48 (Herbrand Universe)** *Given a set of FOL formulas  $\theta^*$  in SNF, the Herbrand Universe  $HU(\theta^*)$  over  $\theta^*$  is the set of all the **ground terms** that can be formed from  $CF(\theta^*)$ .*

We recall what is a ground term:

- if  $a$  is a constant term (function of arity 0), then  $a$  is a ground term
- if  $a_1, \dots, a_k$  are ground terms and  $f$  is a function of arity  $k$ , then  $f(a_1, \dots, a_k)$  is a ground term.

For example let

$$\theta^* = \{\forall x[\neg Dog(x) \vee Chases(x, f(x))]\} \quad (4.42)$$

then  $CF(\theta^*) = \{c, f\}$  and

$$HU(\theta^*) = \{c, f(c), f(f(c)), f(f(f(c))) \dots\} \quad (4.43)$$

Notice how  $HU(\theta^*)$  is infinite in general.

**Definition 49 (Herbrand Expansion)** *Given a set of FOL formulas  $\theta^*$  in SNF, the Herbrand expansion  $HE(\theta^*)$  is the following set of formulas:*

$$HE(\theta^*) = \left\{ \varphi \frac{x_1}{t_1} \dots \frac{x_n}{t_n} \mid (\forall x_1 \dots \forall x_n \varphi) \in \theta^* \wedge t_i \in HU(\theta^*) \right\} \quad (4.44)$$

We are instantiating each possible formula  $\varphi$  from the terms in  $HU(\theta^*)$  that contains ground terms, so it can be interpreted as propositional logic.

**Theorem 15 (Herbrand)** *A set of FOL formulas in SNF  $\theta^*$  is satisfiable<sup>5</sup> if and only if the entire set  $HE(\theta^*)$  is satisfiable (all the propositional logic formulas in  $HE(\theta^*)$  can be satisfied simultaneously by a model.)*

Notice how, without function symbols of arity greater than 0, the Herbrand expansion is finite, and the FOL reasoning is *equivalent* to propositional reasoning.

**Theorem 16 (Compactness of Propositional Logic)** *Any set  $\theta$  of propositional logic formulas is unsatisfiable if and only if at least one finite subset of  $\theta$  is unsatisfiable.*

We can enumerate all finite subsets  $\theta_1$  of the Herbrand expansion  $HE(\theta^*)$  and test propositional satisfiability of  $\theta_1$ ,  $\theta$  is unsatisfiable if and only if one of the subsets  $\theta_1$  is unsatisfiable.

*If the Herbrand expansion is infinite, to show unsatisfiability, we must somehow choose a "relevant" finite subset thereof.*

**Theorem 17** *The set of unsatisfiable FOL formulas is recursively enumerable.*

*Proof:* Enumerate all FOL formulas  $\varphi$ . Incrementally for all of these in parallel, enumerate all finite subsets  $\theta_1$  of the Herbrand expansion  $HE(\theta^*)$ . Test propositional satisfiability of each  $\theta_1$ . By compactness of propositional logic, if  $HE(\theta^*)$  is unsatisfiable then one of the  $\theta_1$  is. ■

**Theorem 18** *The problem of establishing whether a FOL formula is satisfiable or not is undecidable.*

**Corollary 1** *The set of satisfiable FOL formulas is not recursively enumerable.*

*If a FOL formula is unsatisfiable, then we can confirm this. Otherwise, we might end up in an infinite loop.*

<sup>4</sup>We will define it later

<sup>5</sup>A set of formulas is said to be satisfiable if there exists at least one interpretation (called model) which simultaneously makes all the formulas in the set true.

## 4.3 FOL Resolution

### 4.3.1 Substitution and Unification

Consider the Clausal normal form

$$\forall x_1 \forall x_2 \dots \forall x_n (l_1 \vee \dots \vee l_n) \quad (4.45)$$

we can write a form of that structure by omitting the quantifiers in the following notation:

$$\{l_1, \dots, l_n\} \quad (4.46)$$

The idea is to "adapt" the propositional resolution rule

$$\frac{C_1 \dot{\cup} \{l\}, C_2 \dot{\cup} \{\neg l\}}{C_1 \cup C_2} \quad (4.47)$$

to the first order logic.

**Definition 50 (Substitution)** A **Substitution**  $s = \{\frac{x_1}{t_1} \dots \frac{x_n}{t_n}\}$  is a function that substitutes the variables  $x_i$  for terms  $t_i$  where  $x_i \neq t_i$  for all  $i$ . Given a formula  $\varphi$ , we denote  $\varphi s$  the application of  $s$  on  $\varphi$ , which is  $\varphi$  with all occurrences of  $x_i$  simultaneously replaced by  $t_i$ .

Variable instantiation and renaming, as used in the prenex and Skolem transformations as well as in the Herbrand expansion, is a special case of substitution.

for example, given  $s = \{\frac{x}{y}, \frac{y}{h(a,b)}\}$ :

$$P(x, y)s = P(y, h(a, b)) \quad (4.48)$$

**Definition 51 (Composition)** Given two substitutions  $s_1, s_2$  we denote  $s_1 s_2$  the **composed substitution** that is the single subsection identical to  $s_2 \circ s_1$  (apply  $s_1$  then  $s_2$ ).

For example, given  $s_1 = \{\frac{z}{g(x,y)}, \frac{v}{w}\}$  and  $s_2 = \{\frac{x}{a}, \frac{y}{b}, \frac{w}{v}, \frac{z}{d}\}$  we have

$$P(x, y, z, v)s_1 s_2 = P(x, y, g(x, y), w)s_2 = P(a, b, g(a, b), v) \quad (4.49)$$

Given  $s_1$  and  $s_2$  to get  $s_1 s_2$  we can

1. replace all the terms  $t_i$  in  $s_1$  by applying  $s_2$
2. then for any variable  $x_i$  replaced by  $s_2$  but not by  $s_1$ , apply the respective variable/term pair  $\frac{x_i}{t_i}$  of  $s_2$
3. remove any pairs of variable  $x$  and term  $t$  where  $x = t$ .

For any formula  $\varphi$  and triplets of substitutions  $s_1, s_2, s_3$ , the following properties holds:

1.  $(\varphi s_1)s_2 = \varphi(s_1 s_2)$
2.  $(s_1 s_2)s_3 = s_1(s_2 s_3)$
3.  $s_1 s_2 \neq s_2 s_1$  in general.

A substitution  $s = \{\frac{x_1}{t_1} \dots \frac{x_n}{t_n}\}$  is idempotent

$$\varphi s s = \varphi s$$

if  $t_i$  does not contain  $x_j$  for all  $i, j$ .

**Definition 52 (Unifier)** A substitution  $s$  is a **unifier** for a set of expressions  $\{E_i\} = \{E_1 \dots E_k\}$  if  $E_i s = E_j s$  for all  $i, j$ .

**Definition 53** A substitution  $g$  that is a unifier for  $\{E_i\}$  is the **most general unifier (mgu)** if, for any  $s$  unifier for  $\{E_i\}$ , there exists an other  $s'$  unifier for  $\{E_i\}$  such that

$$\{E_i\}s = \{E_i\}gs'. \quad (4.50)$$

**Proposition 4** If any unifier exists, then an idempotent mgu exists.

We want to define an algorithm that finds the mgu for a set of expressions  $\{E_i\}$ .

**Proposition 5** If the only way to unify  $\{E_i\}$  is to unify a variable  $x$  with a term  $t$  that contains  $x$ , then,  $\{E_i\}$  cannot be unified.

We call a **disagreement set** of a set of expressions  $\{E_i\}$ , the set of sub-expressions  $\{t_i\} \subseteq \{E_i\}$  at the first position of  $\{E_i\}$  for which some of the  $\{E_i\}$  disagree. For example with

$$P(x, c, f(y)), P(x, z, z) \text{ have the disagreement set: } \{c, z\}$$

since  $c$  and  $z$  are the first terms on which the expressions in  $\{E_i\}$  disagree. Another example is

$$\{P(x, a, f(y)), P(y, a, f(y))\}$$

the expressions disagree on the first term

$$\begin{aligned} & P(\textcolor{blue}{x}, a, f(y)) \\ & P(\textcolor{blue}{y}, a, f(y)) \implies \\ & \text{disagreement set: } \{x, y\} \end{aligned}$$

We use this definition to build an algorithm that returns a unifier for a set of expressions (if any).

**Theorem 19** The Algorithm 27 succeeds if and only if there is a unifier for  $\{E_i\}$ . In the positive case, the algorithm returns an idempotent mgu of  $\{E_i\}$ .

---

#### Algorithm 27 Find-MGU

---

```

1: given  $\{E_i\}$ 
2:  $k \leftarrow 0$ ,  $T_k \leftarrow \{E_i\}$ ,  $s_k \leftarrow \{\}$ ,
3: while  $T_k$  is not a singleton do
4:   Let  $D_k$  be the disagreement set of  $T_k$ 
5:   /*if  $t_k$  contains  $x_k$  the unification is impossible*/
6:   Let  $x_k, t_k \in D_k$  be a variable and term such that  $t_k$  does not contain  $x_k$ 
7:   if such  $x_k, t_k$  does not exists then
8:     return failure
9:   end if
10:   $s_{k+1} \leftarrow s_k \{ \frac{x_k}{t_k} \}$  /* $t_k$  does not contain any of  $x_1 \dots x_k$ */
11:   $T_{k+1} \leftarrow T_k \{ \frac{x_k}{t_k} \}$  /* $x_k$  does not occur in  $T_{k+1}$ */
12:   $k \leftarrow k + 1$ 
13: end while
14: return  $s_k$ 

```

---

Let's see an example of application, the input set is

$$\{P(x, f(y), y), P(z, f(b), b)\}$$

At the first iteration with  $k = 0$  we have

$$\begin{aligned} D_0 &= \{x, z\} \\ \text{select } x \text{ and } z \text{ in } D_0 \text{ since } z \text{ does not contain } x \\ s_1 &= \{\frac{x}{z}\} \\ \text{we substitute } x \text{ with } z \text{ in the set of expressions} \\ \{P(\textcolor{blue}{x}, f(y), y), P(z, f(b), b)\} &\implies \{P(\textcolor{blue}{z}, f(y), y), P(z, f(b), b)\} \\ T_1 &= \{P(z, f(y), y), P(z, f(b), b)\} \end{aligned}$$

at the second iteration with  $k = 1$  we have

$$\begin{aligned} D_1 &= \{y, b\} \\ s_2 &= s_1 \{ \frac{y}{b} \} = \{\frac{x}{z}, \frac{y}{b}\} \\ T_2 &= \{P(z, f(b), b), P(z, f(b), b)\} = \{P(z, f(b), b)\} \end{aligned}$$

at this point  $T_2$  contains a single element so the output will be  $s_2 = \{\frac{x}{z}, \frac{y}{b}\}$ .

Now we explain in the details the FOL resolution, we assume that the formulas are in Clausal normal form:

$$\forall x_1 \dots \forall x_n (l_1 \vee \dots \vee l_n) \quad (4.51)$$

written

$$\{l_1, \dots, l_n\} \quad (4.52)$$

we recall that a clause is a set of disjunction of literal, a set of clause is denoted  $\Delta$ , the empty clause is denoted  $\square$  and a *calculus* is a set of inference rules.

**Definition 54 (Derivation)** Given a set of clause  $\Delta$  we say that a clause  $C$  can be **derived** from  $\Delta$  using a calculus  $\mathcal{R}$ , written

$$\Delta \models_{\mathcal{R}} C$$

if, starting from  $\Delta$  we can apply a sequence of rule from  $\mathcal{R}$  ending in  $C$ .

We recall the following properties

- A calculus  $\mathcal{R}$  is **sound** if  $\Delta \models_{\mathcal{R}} C \implies \Delta \models C$ . Soundness assures you that the derivation system  $\mathcal{R}$  is "reliable", and that the clause it derives are logically correct with respect to  $\Delta$ .
- A calculus  $\mathcal{R}$  is **resolution complete** if  $\Delta \models \perp$  implies  $\Delta \models_{\mathcal{R}} \square$ , i.e. if  $\Delta$  is unsatisfiable is possible to derive the empty clause from it with  $\mathcal{R}$ .

We remind that to decide whenever  $KB \models \varphi$  we can check the satisfiability of  $\psi = KB \cup \{\varphi\}$ . If this is unsatisfiable then  $KB \models \varphi$ . So the idea is

- to check if  $KB \models \varphi$  we consider  $KB \cup \{\varphi\}$
- we try to derive the empty clause  $\square$  from  $KB \cup \{\varphi\}$ .

Now let's see our resolution rule.

**Definition 55 (Propositional Resolution)** The resolution uses the following resolution rule:

$$\frac{C_1 \dot{\cup} \{l\}, C_2 \dot{\cup} \{\neg l\}}{C_1 \cup C_2} \quad (4.53)$$

So, if  $\Delta$  contains two clauses of the form

$$C_1 \dot{\cup} \{l\}, C_2 \dot{\cup} \{\neg l\}$$

the rule derive the new clause  $C_1 \cup C_2$ ,  $l$  and  $\neg l$  are called resolution literals.

**Definition 56 (Binary FOL Resolution)** The resolution uses the following resolution rule:

$$\frac{C_1 \dot{\cup} \{P_1\}, C_2 \dot{\cup} \{\neg P_2\}}{[C_1 \cup C_2]g} \quad (4.54)$$

where  $g$  is the mgu.

If  $\Delta$  contains two clauses of the form

$$\{C_1 \dot{\cup} \{P_1\}, C_2 \dot{\cup} \{\neg P_2\}\}$$

where  $P_1$  and  $P_2$  can be unified and  $g$  is an mgu for them, the rule allows to derive the clause  $[C_1 \cup C_2]g$  (the union of  $C_1$  and  $C_2$  where we apply the unification).  $P$  and  $\neg P$  are called resolution literals. The two rules presented satisfies the properties of soundness.

**Theorem 20** The calculus with only the binary FOL resolution is not complete.

As a proof consider the following example:

$$\Delta = \{\{P(x_1, x_2), P(x_2, x_1)\}, \{\neg P(y_1, y_2), \neg P(y_2, y_1)\}\} \quad (4.55)$$

this  $\Delta$  is not satisfiable, the variables in FOL clauses are universally quantified, to satisfy  $\Delta$  we would have to satisfy  $P(o, o)$  and  $\neg P(o, o)$  for all the objects  $o$  in the universe. We notice that we can't derive the empty clause  $\square$  from  $\Delta$  using the binary FOL resolution.

Now we present two rules that forms a sound and complete calculus.

**Definition 57 (Full FOL Resolution)** *The resolution uses the following resolution rule:*

$$\frac{C_1 \dot{\cup} \{P_1^1 \dots P_1^n\}, C_2 \dot{\cup} \{\neg P_2^1 \dots \neg P_2^n\}}{[C_1 \cup C_2]g} \quad (4.56)$$

where  $\{P_1^1, \dots, P_1^n, P_2^1, \dots, \neg P_2^n\}$  can be unified and  $g$  is an mgu for them.

**Definition 58 (Factoring)** *The resolution uses the following resolution rule:*

$$\frac{C_1 \dot{\cup} \{l_1\} \dot{\cup} \{l_2\}}{[C_1 \cup \{l_1\}]g} \quad (4.57)$$

where  $l_1$  and  $l_2$  can be unified and  $g$  is an mgu for them.  $[C_1 \cup \{l_1\}]g$  is a **factor** of the parent clause.

We denote  $\mathcal{R}_{FactBin}$  the calculus with the factoring rule and  $\mathcal{R}_{Full}$  the calculus with the full FOL resolution.

**Lemma 4 (Lifting Lemma)** *Let  $C_1$  and  $C_2$  be two clauses with no shared variables, and let  $C_1^g$  and  $C_2^g$  be ground instances of  $C_1$  and  $C_2$ . Say that  $C^g$  is a resolvent of  $C_1^g$  and  $C_2^g$ . Then there exists a clause  $C$  such that  $C^g$  is a ground instance of  $C$ , and:*

1.  $C$  can be derived from  $C_1$  and  $C_2$  using  $\mathcal{R}_{Full}$ .
2.  $C$  can be derived from  $C_1$  and  $C_2$  using  $\mathcal{R}_{FactBin}$ .

**Theorem 21** *The calculus with the full FOL resolution and with the factoring resolution is sound and complete.*

A sketch of the proof is presented through previous intermediate results:

- Any set  $\theta$  of FOL formulae  $s$  can be represented in clausal form  $\Delta$
- we assume that  $\Delta$  is unsatisfiable
- we use the *Herbrand* Theorem 15 and the propositional compactness Proposition 16, so the (assumed) unsatisfiability of  $\Delta$  translates to: some finite set  $\Delta'$  of ground instances is unsatisfiable.
- by compactness of propositional resolution, we can derive  $\square$  from  $\Delta'$
- $\square$  can also be derived directly from  $\Delta$  with the full FOL resolution and the factoring resolution by the lifting Lemma 4.

## CHAPTER

# 5

## PLANNING

Planning (or automatic planning) is a branch of artificial intelligence that deals with generating a strategy or sequence of actions to allow an intelligent agent (such as a robot or software) to achieve an objective starting from an initial state. It's a general way to define a problem solving procedure.

### 5.1 STRIPS

STRIPS (acronym for STanford Research Institute Problem Solver) is one of the most important automatic planning languages and systems in the history of artificial intelligence.

Developed in 1971 by Richard Fikes and Nils Nilsson, it was used to control the actions of Shakey, one of the first mobile robots with logical reasoning. In modern terms, STRIPS represents the standard for so-called classic planning.

STRIPS has only boolean variables, the preconditions and the goals are conjunctions of positive atoms, the effects are conjunctions of literals (positive or negated atoms).

**Definition 59 (STRIPS Planning Task)** A STRIPS planning task, short planning task, is a 4-tuple  $\Pi = (P, A, I, G)$  where:

- $P$  is a finite set of **facts** (aka **propositions**).
- $A$  is a finite set of **actions**; each  $a \in A$  is a triple  $a = (pre_a, add_a, del_a)$  of subsets of  $P$  referred to as the action's **precondition**, **add list**, and **delete list** respectively; we require that  $add_a \cap del_a = \emptyset$ .
- $I \subseteq P$  is the **initial state**.
- $G \subseteq P$  is the **goal**.

We will often give each action  $a \in A$  a **name** (a string), and identify  $a$  with that name.

#### Example: Australian Logistics Planning Task

- **Facts**  $P$ :  $\{at(x), visited(x) \mid x \in \{Sydney, Adelaide, Brisbane, Perth, Darwin\}\}$ .
- **Initial state**  $I$ :  $\{at(Sydney), visited(Sydney)\}$ .
- **Goal**  $G$ :  $\{at(Sydney)\} \cup \{visited(x) \mid x \in \{Sydney, Adelaide, Brisbane, Perth, Darwin\}\}$ .
- **Actions**  $a \in A$ :  $drive(x, y)$  where  $x, y$  have a road.
  - **Precondition**  $pre_a$ :  $\{at(x)\}$ .

- **Add list**  $add_a: \{at(y), visited(y)\}$ .
- **Delete list**  $del_a: \{at(x)\}$ .
- **Plan:**  $\langle drive(Sydney, Brisbane), drive(Brisbane, Sydney), drive(Sydney, Adelaide), drive(Adelaide, Perth), drive(Darwin, Adelaide), drive(Adelaide, Sydney) \rangle$ .



**Definition 60 (STRIPS State Space)** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task. The state space of  $\Pi$  is  $\Theta_\Pi = (S, A, T, I, S^G)$  where:

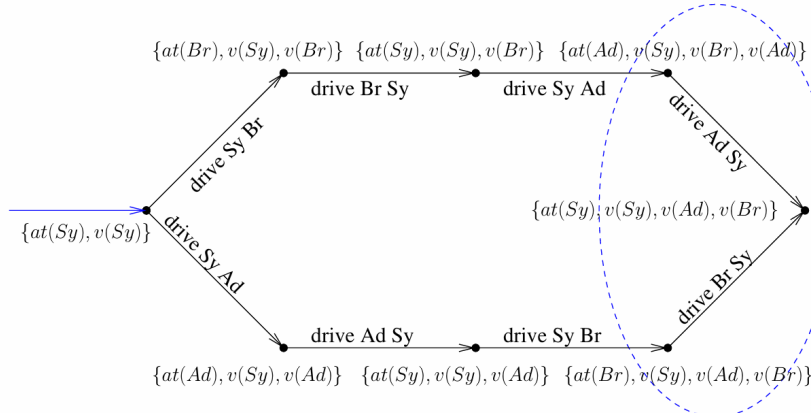
- The states (also **world states**)  $S = 2^P$  are the subsets of  $P$ .
- $A$  is  $\Pi$ 's action set.
- The transitions are  $T = \{s \xrightarrow{a} s' \mid pre_a \subseteq s, s' = (\llbracket s \rrbracket, a)\}$ .  
If  $pre_a \subseteq s$ , then  $a$  is **applicable** in  $s$  and

$$(\llbracket s \rrbracket, a) := (s \cup add_a) \setminus del_a$$

If  $pre_a \not\subseteq s$ , then  $(\llbracket s \rrbracket, a)$  is undefined.

- $I$  is  $\Pi$ 's initial state.
- The goal states  $S^G = \{s \in S \mid G \subseteq s\}$  are those that satisfy  $\Pi$ 's goal.

A **plan** for  $s \in S$  is a path from  $s$  to  $s' \in S^G$ , a plan for  $I$  is a **solution** and is called *plan for  $\Pi$* , if such plan exists,  $\Pi$  is **solvable**.



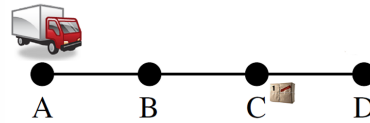
These planning examples draw heavily on the research problems presented in previous chapters, as in this case, it is possible to define a heuristic function that estimates the distance (or cost) between a current state  $s$  and the goal state  $G$ .

It is important to perform a complexity analysis of the problems that we face to identify special cases that can be solved in polynomial time, relax the input into the special case to obtain a heuristic function.

### 5.1.1 Only-Adds STRIPS Tasks

Let's consider the following example:

- **Facts  $P$ :**  $\{truck(x) \mid x \in \{A, B, C, D\}\} \cup \{pack(x) \mid x \in \{A, B, C, D, T\}\}$ .
- **Initial state  $I$ :**  $\{truck(A), pack(C)\}$ .
- **Goal  $G$ :**  $\{truck(A), pack(D)\}$ .
- **Actions  $A$ :** (Notated as "precondition  $\Rightarrow$  adds,  $\neg$  deletes")
  - $drive(x, y)$ , where  $x, y$  have a road:  
 $"truck(x) \Rightarrow truck(y), \neg truck(x)"$ .
  - $load(x)$ : " $truck(x), pack(x) \Rightarrow pack(T), \neg pack(x)$ ".
  - $unload(x)$ : " $truck(x), pack(T) \Rightarrow pack(x), \neg pack(T)$ ".



It is clear that this STRIPS planning task models a truck that must collect a package and deposit it in a designated place. One simple relaxation is the following:

**Only-Adds Relaxation:** Drop the preconditions and deletes on the actions.

We want to use this to generate a heuristic function. The general problem is

- Given a STRIPS task  $\Pi = (P, A, I, G)$ , we want to find an action sequence  $\vec{a}$  leading from  $I$  to a state that contains  $G$  when pretending that preconditions and deletes are empty.

The simplest possible approach is the following: This algorithm is applied for each state  $s$  and calculates

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**Algorithm 28** Solution 1

---

```

1:  $\vec{a} = \langle \rangle$ 
2: while  $G \neq \emptyset$  do
3:   select  $a \in A$ 
4:    $G = G \setminus add_a$  #Subtract from the remaining objectives anything that the chosen action adds
5:    $\vec{a} = \vec{a} \circ \langle a \rangle$ 
6:    $A = A \setminus \{a\}$ 
7: end while
8: return  $h = |\vec{a}|$ 

```

---

the value of  $h$ . This heuristic is not admissible, admissibility is only guaranteed if we find a shortest possible  $\vec{a}$ ; else,  $\vec{a}$  might be longer than a plan for  $\Pi$  itself. Selecting an arbitrary action each time,  $\vec{a}$  may be longer than needed. Another solution might be:

---

**Algorithm 29** Solution 2

---

```

1:  $\vec{a} = \langle \rangle$ 
2: while  $G \neq \emptyset$  do
3:   select  $a \in A$  such that  $|add_a|$  is maximal.
4:    $G = G \setminus add_a$ 
5:    $\vec{a} = \vec{a} \circ \langle a \rangle$ 
6:    $A = A \setminus \{a\}$ 
7: end while
8: return  $h = |\vec{a}|$ 

```

---

$h$  is not admissible yet, large  $add_a$  doesn't help if the intersection with  $G$  is small.

**Algorithm 30** Solution 3

---

```

1:  $\vec{a} = \langle \rangle$ 
2: while  $G \neq \emptyset$  do
3:   select  $a \in A$  such that  $|add_a \cap G|$  is maximal.
4:    $G = G \setminus add_a$ 
5:    $\vec{a} = \vec{a} \circ \langle a \rangle$ 
6:    $A = A \setminus \{a\}$ 
7: end while
8: return  $h = |\vec{a}|$ 
```

---

Also this is not admissible. To have an admissible (and optimal) heuristic in the relaxed world, we should find the shortest possible sequence of actions (optimal  $\vec{a}$ ). But finding this sequence is equivalent to solving the Minimum Cover problem, since Minimum Cover is an NP-complete problem, finding the optimal heuristic even for a relaxed problem is computationally expensive.

We call **PlanEx** the problem of deciding, given a STRIPS planning task  $\Pi$ , whether or not there exists a plan for  $\Pi$ .

**Lemma 5** *PlanEx is PSPACE-hard.*

*Proof Sketch:* Given a Turing machine with **space bounded by polynomial**  $p(|w|)$ , we can in polynomial time (in the size of the machine) generate an equivalent STRIPS planning task. Say the possible symbols in tape cells are  $x_1, \dots, x_m$  and the internal states are  $s_1, \dots, s_n$ , accepting state  $s_{acc}$ .

- The contents of the tape cells:  $in(1, x_1), \dots, in(p(|w|), x_1), \dots, in(1, x_m), \dots, in(p(|w|), x_m)$ .
- The position of the R/W head:  $at(1), \dots, at(p(|w|))$ .
- The internal state of the machine:  $state(s_1), \dots, state(s_n)$ .
- Transitions rules  $\mapsto$  STRIPS actions; accepting state  $\mapsto$  STRIPS goal  $\{state(s_{acc})\}$ ; initial state obvious.
- This reduction to STRIPS runs in polynomial-time because we need only polynomially many facts.

**Lemma 6** *PlanEx is in PSPACE.*

It is easy and sufficient to prove that PlanEx is in NPSPACE, since PSPACE = NPSPACE. From the two lemmas, the following theorems follows.

**Theorem 22** *PlanEx is PSPACE-complete.*

### 5.1.2 Transition System

The state space of a planning task, like the one described in Definition 60, is generalized in the following definition.

**Definition 61 (Transition Systems)** *A transition system is a 6-tuple  $\Theta = (S, L, c, T, I, S^G)$  where:*

- $S$  is a finite set of states
- $L$  is a finite set of transition labels (a generalization of an action)
- $c : L \mapsto \mathbb{R}^+ \cup \{0\}$  is the cost function
- $T \subseteq S \times L \times S$  is the transition relation
- $S^G \subseteq S$  is the set of the goal states.
- $I \in S$  is the initial state.

The size of  $\Theta$  is  $|S|$ . If  $\Theta$  has the transition  $(s, l, s') \in T$  we write

$$s \xrightarrow{l} s'$$

or

$$s \rightarrow s'$$

if we are not interested in the label  $l$ .  $\Theta$  has unit cost if  $c$  is the constant function  $c(l) = 1, \forall l$ . The classic definitions already addressed follow:

- $s'$  is a successor of  $s$  if  $s \rightarrow s' \in T$
- $s'$  is a predecessor of  $s$  if  $s' \rightarrow s \in T$
- a path from  $s$  to  $s'$  is a sequence of transitions:

$$s \xrightarrow{l_1} s_1 \xrightarrow{l_2} s_2 \dots \xrightarrow{l_n} s_n = s' \quad (5.1)$$

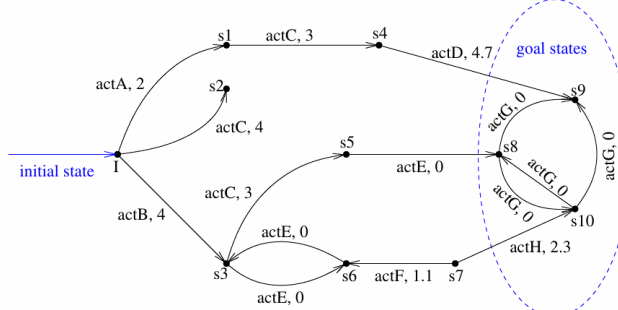
and the cost of the path is

$$\sum_{i=1}^n c(l_i) \quad (5.2)$$

we say that  $s$  is reachable (without specifying the origin) meaning that is reachable from  $I$ .

A solution for  $\Theta$  is a path from  $I$  to a state  $s \in S^G$ . If such path exists,  $\Theta$  is solvable.

**Directed labeled graphs** + mark-up for initial state and goal states:



We are interested in solving **huge** transition systems (unfeasible to apply Dijkstra), represented in a compact way as planning tasks.

## 5.2 FDR

STRIPS and FDR (Finite Domain Representation) are two different formalisms for describing planning tasks, both aimed at defining how an agent should move from an initial state to a goal. FDR is a more modern and often more efficient evolution for automatic solvers (like Fast Downward):

- Finite Domain Variables: Instead of having many Boolean facts (e.g. *truck\_in\_A*, *truck\_in\_B*), use a single multivariate variable (e.g. *TruckPosition*) that can take a value in a finite set  $\{A, B, C, D\}$
- State: A function that assigns a value to each variable.
- Advantage: Reduces the number of "mutually exclusive" facts the planner must explicitly handle, making the problem structure clearer.

**Definition 62 (FDR Planning Task)** A finite-domain representation planning task, short FDR planning task, is a 5-tuple  $\Pi = (V, A, c, I, G)$  where:

- $V$  is a finite set of state variables, each  $v \in V$  with a finite domain  $D_v$ .  
We refer to (partial) functions on  $V$ , mapping each  $v \in V$  into a member of  $D_v$ , as (partial) variable assignments.

- $A$  is a finite set of actions; each  $a \in A$  is a pair  $(\text{pre}_a, \text{eff}_a)$  of partial variable assignments referred to as the action's precondition and effects.
- $c : A \mapsto \mathbb{R}^+ \cup \{0\}$  is the cost function.
- $I$  is a complete variable assignment called the initial state.
- $G$  is a partial variable assignment called the goal.

We say that  $\Pi$  has *unit costs* if, for all  $a \in A$ ,  $c(a) = 1$ . In the FDR context we call **fact** an assignment

$$(v, d) \text{ or } v = d$$

we mean that the variable  $v$  assumes the value  $d$ . If  $p$  is a partial variable assignment, we write

$$V[p] = \{v \in V \mid p(v) \text{ is defined}\}$$

to refers to all variables where  $p$  is defined. For the map example in Australia:



- **Variables**  $V$ :  $at : \{Sydney, Adelaide, Brisbane, Perth, Darwin\}$ ;  $\text{visited}(x) : \{T, F\}$  per ogni città  $x$ .
- **Initial state**  $I$ :  $at = Sydney, \text{visited}(Sydney) = T, \text{visited}(x) = F$  per  $x \neq Sydney$ .
- **Goal**  $G$ :  $at = Sydney, \text{visited}(x) = T$  per ogni  $x$ .
- **Actions**  $a \in A$ :  $drive(x, y)$  dove  $x, y$  hanno una strada.
  - Precondizione  $\text{pre}_a$ :  $at = x$ .
  - Effetto  $\text{eff}_a$ :  $at = y, \text{visited}(y) = T$ .

- **Cost function**  $c$ :

$$c(\text{drive}(x, y)) = \begin{cases} 1 & \{x, y\} = \{Sydney, Brisbane\} \\ 1.5 & \{x, y\} = \{Sydney, Adelaide\} \\ 3.5 & \{x, y\} = \{Adelaide, Perth\} \\ 4 & \{x, y\} = \{Adelaide, Darwin\} \end{cases}$$

**Definition 63 (FDR State Space)** Let  $\Pi = (V, A, c, I, G)$  to be an FDR planning task, the **task space** of  $\Pi$  is the labeled transition system  $\Theta_\Pi = (S, L, c, T, I, S^G)$  where

- the states  $S$  are all the possible complete variable assignments.
- the labels  $L = A$  are the actions of  $\Pi$ , hence, the cost  $c$  of  $\Pi$  defined on  $A$  is equal of the cost  $c$  of  $\Theta_\Pi$  defined on  $L$ .
- The transitions are

$$T = \{s \xrightarrow{a} s' \mid a \in A[s], s' = s[a]\} \quad (5.3)$$

where

- $A[s] = \{a \in A \mid \text{pre}_a \subseteq s\}$  are the actions that can be applied in  $s$ .  $s[a]$  represents the resulting state after applying the  $a$  action.

$$s[a] = \begin{cases} \text{undefined if } a \notin A[s] \\ \text{eff}_a(v) \text{ if } v \in V[\text{eff}_a] \\ s(v) \text{ if } v \notin V[\text{eff}_a] \end{cases}$$

Although the STRIPS language is still used for historical convenience, modern planning systems (such as Fast Downward) prefer to translate everything internally into FDR for reasons of practical efficiency. The FDR format is superior because it reduces unnecessary states, allows you to better map dependencies via causal graphs, and facilitates the creation of more powerful heuristics, offering a much more natural and compact way of modeling reality than the rigid Boolean variables of STRIPS.

### 5.2.1 Conversion between FDR and STRIPS

An easy way to convert an FDR planning task in a STRIPS planning task is the following

- for each variable  $v$  with domain  $\{d_1 \dots d_k\}$ , we make  $k$  STRIPS facts

$$v = d_1$$

$$\vdots$$

$$v = d_k$$

**Definition 64** Let  $\Theta = (S, L, c, T, I, S^G)$  and  $\Theta' = (S', L', c', T', I', S'^G)$  to be two transition systems. We say that  $\Theta$  is **isomorphic** to  $\Theta'$

$$\Theta \sim \Theta'$$

if there exists two bijective functions  $\varphi : S \mapsto S'$  and  $\psi : L \mapsto L'$  such that

- $\varphi(I) = I'$
- $s \in S^G$  if and only if  $\varphi(s) \in S'^G$
- $(s, l, t) \in T$  if and only if  $(\varphi(s), \psi(l), \varphi(t)) \in T'$
- for all  $l \in L$ ,  $c(l) = c'(\psi(l))$

Isomorphisms typically result from compilations between different formalisms (see later this chapter); we will also sometimes use them as a technical device.

Now we consider a systematic way to convert an FDR planning task in a STRIPS planning task.

**Definition 65 (FDR to STRIPS)** Let  $\Pi = (V, A, c, I, G)$  to be an FDR planning task, the **STRIPS conversion** is the STRIPS planning task  $\Pi^{STR} = (P_V, A^{STR}, c, I, G)$  where

- $P_V = \{v = d \mid v \in V, d \in D_v\}$  is the set of STRIPS facts.
- $A^{STR} = \{a^{STR} \mid a \in A\}$  where  $pre_{a^{STR}} = pre_a$ ,  $add_{a^{STR}} = eff_a$  and

$$del_{a^{STR}} = \bigcup_{(v=d) \in eff_a} \begin{cases} \{v = pre_a(v)\} & \text{if } pre_a(v) \text{ is defined} \\ \{v = d' \mid d' \in D_v \setminus \{d\}\} & \text{otherwise} \end{cases} \quad (5.4)$$

- the cost function  $c$  is defined by

$$c(a^{STR}) = c(a), \forall a^{STR} \in A^{STR}$$

- $I$  and  $G$  are identical to those in  $\Pi$ .

Note how a STRIPS fact is a pair variable/value in FDR:

- If a variable in *FDR* can be

$$Weather = \{sunny, rainy\}$$

- then there will be two STRIPS facts

$$\begin{aligned} Weather &= sunny \\ Weather &= rainy \end{aligned}$$

**Proposition 6** Let  $\Pi = (V, A, c, I, G)$  to be an FDR planning task and  $\Pi^{STR} = (P_V, A^{STR}, c, I, G)$  it's STRIPS conversion as in Definition 65. Let  $\Theta_\Pi$  to be the transition system (state space) of  $\Pi$ , and  $\Theta_{\Pi^{STR}}$  the transition system with the states  $s \subseteq P_v$  where, for each  $v \in V$ ,  $s$  contains exactly one fact of the form  $v = d$ , with all other states in  $\Theta_{\Pi^{STR}}$  unreachable. Then,  $\Theta_\Pi$  is isomorphic to  $\Theta_{\Pi^{STR}}$ .

In the proposition it is important to note that, in each state, for each FDR variable  $v$ , there must be exactly one fact  $v = d$  for some  $d \in D_v$ , therefore, if there are  $n$  variables in the FDR planning task

$$v_1, v_2 \dots v_n$$

, a state of the state space of the STRIPS conversion is of the type

$$\begin{aligned} v_1 &= d_1 \\ v_2 &= d_2 \\ &\vdots \\ v_n &= d_n \end{aligned}$$

That is, each variable must have a single assignment.

- FDR  $V$ :  $at : \{Sydney, Adelaide, Brisbane\}$ ;  $visited(x) : \{T, F\}$  for  $x \in \{Sydney, Adelaide, Brisbane\}$ .
- STRIPS  $P$ :  $at(x), visited(x, T), visited(x, F)$  for  $x \in \{Sydney, Adelaide, Brisbane\}$ .
- FDR  $dr(x, y)$ :  $pre = \{at = x\}$ ,  $eff = \{at = y, v(y) = T\}$ .
- STRIPS  $dr(x, y)$ :  $pre = \{at(x)\}$ ,  $add = \{at(y), v(y, T)\}$ ,  $del = \{at(x), v(y, F)\}$ .

**Definition 66 (STRIPS to FDR)** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task, the **FDR conversion** is the FDR planning task  $\Pi^{FDR} = (V_P, A^{FDR}, c, I^{FDR}, G^{FDR})$  where

- $V_P = \{v_p \mid p \in P\}$  is the set of variables, all boolean
- $A^{FDR} = \{a^{FDR} \mid a \in A\}$  where
  - $pre_{a^{FDR}} = \{v_p = True \mid p \in pre_a\}$
  - $eff_{a^{FDR}} = \{v_p = True \mid p \in add_a\} \cup \{v_p = False \mid p \in del_a\}$
- the cost function  $c$  is defined by

$$c(a^{FDR}) = c(a), \forall a^{FDR} \in A^{FDR}$$

- $I = \{v_p = True \mid p \in I\}$
- $G = \{v_p = True \mid p \in G\}$

This conversion is simple, hence is called *Naïve translation*, all the variables of the conversion is boolean, so this does not benefit at all from the added expressivity of FDR.

**Proposition 7** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task and  $\Pi^{FDR} = (V_P, A^{FDR}, c, I^{FDR}, G^{FDR})$  it's FDR conversion as in Definition 66. Let  $\Theta_\Pi$  to be the transition system (state space) of  $\Pi$ , and  $\Theta_{\Pi^{FDR}}$  the transition system (state space) of  $\Pi^{FDR}$ , then  $\Theta_{\Pi^{FDR}}$  are isomorphic to  $\Theta_\Pi$ .

- STRIPS  $P$ :  $at(x), visited(x)$  for  $x \in \{Sydney, Adelaide, Brisbane\}$ .
- FDR  $V$ :  $at(x), visited(x) : \{T, F\}$  for  $x \in \{Sydney, Adelaide, Brisbane\}$ .
- STRIPS  $dr(x, y)$ :  $pre = \{at(x)\}$ ,  $add = \{at(y), v(y)\}$ ,  $del = \{at(x)\}$ .
- FDR  $dr(x, y)$ :  $pre = \{at(x) = T\}$ ,  $eff = \{at(y) = T, v(y) = T, at(x) = F\}$ .

There is other conversion method that are more effective and exploit the true expressivity of FDR, but we don't cover these methods here.

## 5.3 Planning Domain Definition Language

The Planning Domain Definition Language (PDDL) is an input language used in the planning area as a standard, like FOL, PDDL describes the world in a schematic way relative to a set of objects. This makes the encoding much smaller and easier to write.

In a few words, the PDDL is used to explain to a computer:

1. What are the "rules of the world" (what can be done).
2. How is the current situation.
3. What is the objective to be achieved.

The schematic encoding of the PDDL use *predicates* instead of the STRIPS propositions, and *action schemas* instead of the STRIPS actions.

A PDDL planner takes as an input two different files:

- **domain.pddl**: It defines the "universe" of the problem. It is the constant part that does not change between different exercises of the same type. Contains predicates, types, action schemas.
- **problem.pddl**: Defines the specific instance you want to resolve, contains the objects, the initial state and the goal.

### 5.3.1 The Domain File

This file contains a requirements definition (by default simply write at the beginning of the file `:adl :typing`), the definition of the types for the variables, the definition of the predicates and the action schemas. An example is the following:

```
(define (domain BlocksWorld)
  (:requirements :adl :typing)
  (:types block - object
         blueblock smallblock - block)
  (:predicates (on ?x - smallblock ?y - block)
               (ontable ?x - block)
               (clear ?x - block))
  )
```

An action schema require a list of parameter

```
(:action <name>
  (?x- type1 ?y- type2 ?z- type3))
```

a precondition written as a formula

```
<predicate>
  (&lt;formula> ... <formula>)
  (&or <formula> ... <formula>)
  (&not <formula>)
  (&forall (?x1- type1 ... ?xn- typen) <formula>)
  (&exists (?x1- type1 ... ?xn- typen) <formula>)
```

and the effect of the action as a combination of literals, conjunction, conditional effects, and quantification over effects:

```
<predicate>
  (&not <predicate>)
  (&and <effect> ... <effect>)
  (&when <formula> <effect>)
  (&forall (?x1- type1 ... ?xn- typen) <effect>)
```

### 5.3.2 The Problem File

A PDDL problem file consists of:

- the definition of the problem name and the relative domain:

```
(define (problem <name>)
(:domain <name>)
```

- the definitions of objects belonging to each type
- definition of the initial state (list of ground predicates initially true)
- definition of the goal (a formula like action preconditions).

An example of a problem file is the following

```
(define (problem example)
(:domain BlocksWorld)
(:objects a b c - smallblock)
d e - block
f - blueblock)
(:init (clear a) (clear b) (clear c)
(clear d) (clear e) (clear f)
(ontable a) (ontable b) (ontable c)
(ontable d) (ontable e) (ontable f))
(:goal (and (on a d) (on b e) (on c f)))
)
```

So a PDDL program is a "translation" for the computer of a STRIPS planning task.

### 5.3.3 Syntax

We give a brief and intuitive explanation of the PDDL syntax. In the domain file, the first thing to do is to define the types:

```
(:types
    vehicle - object
    truck car - vehicle
    pack
)
```

When you write on a single line

```
obj1 obj2 ... objn - root_obj
```

you are defining the objects `obj1 obj2 ... objn` and declaring that they are of the type `root_obj`. If not specified, each object inheritance the root type `object`.

When you define actions that require a certain object type `x`, you can also pass an object of a type that inherit that type `x`.

After defining the types, it is necessary to define the predicates, these describe properties and relations of the world that can be true or false. The PDDL adopts the Closed World Assumption:

- If a predicate is listed in its current state, it is TRUE.
- If a predicate is NOT listed, it is automatically considered FALSE.

There is no such thing as "unknown" status. A predicate can only go from true to false (or vice versa) through the effects of actions. A predicates is written as follows:

```
(:predicates
    (pred_name ?var1 - typ1 ?var2 - typ2 ... ?varn - typn)
)
```

The strings with `?` as the prefix are the variables, in the problem file we will use the predicates to assign relations between the objects. An example is

```
(:predicates
  (has-key ?r - robot)
  (door-open ?p - door)
  (is_close_to ?r - robot ?p - door)
)
```

In the problem file, you can define several robot-type objects and a door-type object, and then declare that some robots are near the door using the appropriate predicate. An example is the following:

```
(:objects
  r2d2 c3po - robot ;; Lets define two specific robots
)
(:init
  (has-key r2d2) ;; r2d2 has the key, c3po doesn't
)
```

An action scheme use the predicates for the precondition and the effect as follows:

```
(:action key_pass
  :parameters (?A - robot ?B - robot)
  :precondition (has-key ?A)
  :effect (and (not (has-key ?A)) (has-key ?B))
)
```

In the action it is specified that, if a robot `?A` has the key (as specified by the pre-conditions) it can pass it to a robot `?B`, and in the effects it is specified that, after the action, `has-key ?A` will be false (the object that passed the key will no longer have the key) while `has-key ?B` will be true.

In a problem file it is necessary to define a series of objects and assign them the relative types, to then define the initial conditions through the predicates, and the goal state to always be reached through the use of the predicates. The solver will take care of using the actions to verify whether the goal is reachable and possibly provide a resolution strategy.

```
(define (problem key_mission)
  (:domain robot-world)
  (:objects
    r2d2 c3po - robot
    golden_key - key
    room_a room_b - place
  )
  (:init
    (at r2d2 room_a)
    (at c3po room_b)
    (at golden_key room_a)
    (has-key r2d2 golden_key)
  )
  (:goal
    (and
      (at c3po room_b)
      (has-key c3po golden_key)
    )
  )
)
```

### 5.3.4 Example

A classic and intuitive example for PDDL is the Logistics Problem. In this scenario, we have a robot that needs to move packages between different locations.

#### The Scenario

- Locations: A Warehouse and a Store.
- Objects: One Package and one Robot.
- Goal: Move the package from the Warehouse to the Store.
- Actions: The robot can move between locations, pick-up a package, and drop-off a package.

The domain file is the following:

```
(define (domain logistics-study)

(:requirements :strips :typing)

(:types
  location item robot
)

(:predicates
  (at ?obj - object ?loc - location)
  (carrying ?robot - robot ?item - item)
  (empty ?robot - robot)
)

(:action move
  :parameters (?r - robot ?from - location ?to - location)
  :precondition (at ?r ?from)
  :effect (and
    (at ?r ?to)
    (not (at ?r ?from))
  )
)
)
```

the predicate `at` is needed to model the fact that a generic object can be in relation with an object of type `location`, obviously the meaning is that that object is in that location. The meaning of the predicate `carrying` is also obvious, as for `empty`, which serves to identify when a robot is not carrying anything.

```
(:action pick-up
  :parameters (?r - robot ?i - item ?l - location)
  :precondition (and (at ?r ?l) (at ?i ?l) (empty ?r))
  :effect (and
    (carrying ?r ?i)
    (not (at ?i ?l))
    (not (empty ?r))
  )
)
)

(:action drop-off
  :parameters (?r - robot ?i - item ?l - location)
  :precondition (and (at ?r ?l) (carrying ?r ?i))
  :effect (and
    (at ?i ?l)
    (empty ?r)
    (not (carrying ?r ?i))
  )
)
)
```

The problem file is the following:

```
(define (problem move-package-01)
  (:domain logistics-study)

  (:objects
    warehouse store - location
    package1 - item
    bot1 - robot
  )

  (:init
    (at bot1 warehouse)
    (at package1 warehouse)
    (empty bot1)
  )

  (:goal
    (at package1 store)
  )
)
```

The solver will take care of finding the right sequence of actions (navigating in the state space) to achieve the goal starting from the defined initial state.

## 5.4 Causal Graph

**Definition 67** Let  $\Pi = (V, A, c, I, G)$  to be an FDR planning task, the **causal graph** of  $\Pi$  is the directed graph  $CG(\Pi)$  where

- the vertices of  $CG(\Pi)$  are the variables  $V$  of  $\Pi$ .
- there exists an arc  $(u, v)$  with  $u \neq v$  if there exists an action  $a \in A$  such that
  - $pre_a(u)$  and  $eff_a(v)$  are both defined, OR
  - $eff_a(u)$  and  $eff_a(v)$  are both defined.

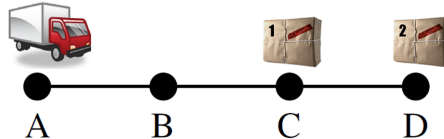
The causal graph capture the dependencies from the variables, the first type of arc have the following meaning

we may have to change  $u$  to be able to change  $v$

the second type of arc have the following meaning

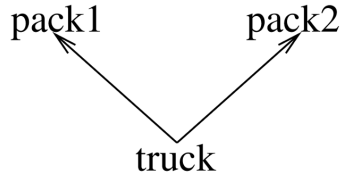
changing  $u$  may (as a side effect) change  $v$  as well (and viceversa)

this second type of arc create a cycle between the two variables. Let's consider the truck example.



- State variables  $V$ :  $truck : \{A, B, C, D\}$ ;  $pack1, pack2 : \{A, B, C, D, T\}$ .
- Initial state  $I$ :  $truck = A, pack1 = C, pack2 = D$ .
- Goal  $G$ :  $truck = A, pack1 = D$ .
- Actions  $A$   
(unit costs):  $drive(x, y), load(p, x), unload(p, x)$ .

we notice how the action *unload* have in the preconditions the truck (it must have a pack) and in its side effect, one of the packs (since their location will change) so the causal graph is the following:



### Second Example (Australian Cities/Visiting)

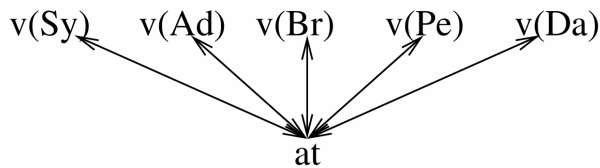
- Variables  $V$ :  $at : \{Sy, Ad, Br, Pe, Da\}; v(x) : \{T, F\}$  for  $x \in \{Sy, Ad, Br, Pe, Da\}$ .
- Initial state  $I$ :  $at = Sy, v(Sy) = T, v(x) = F$  for  $x \neq Sy$ .
- Goal  $G$ :  $at = Sy, v(x) = T$  for all  $x$ .
- Actions  $A$ :  $drive(x, y)$  where  $x, y$  have a road; pre  $at = x$ , eff  $at = y, v(y)$ .
- Cost function  $c$ :  $Sy \leftrightarrow Br : 1, Sy \leftrightarrow Ad : 1.5, Ad \leftrightarrow Pe : 3.5, Ad \leftrightarrow Da : 4$ .



In the action  $drive(x, y)$  we notice how

- the variable  $at$  is in the precondition and the variable  $v(x)$  is in the effects, so there will be an arc from  $at$  to  $v(x)$ .
- either  $at$  and  $v(x)$  are in the effect of the same action, so there will be a bi-directed arc between these two variables.

The causal graph is:



So, we defined two types of arc, the class 1 (precondition-effect) and the class 2 (effect-effect). In the causal graph, an "arc" of class two is in fact a couple of directed arc forming a cycle between two variables, occur whenever an action has more than one effect.

A cycle of class 1 (precondition-effect) occur when there are "cyclic support dependencies", where moving variable  $x$  requires a precondition on  $y$  which (transitively) requires a precondition on  $x$ .

A causal graph don't depend on the initial state  $I$  or the goal state  $G$ , this is a main weakness of causal graphs, they capture only the structure of the variables and actions, and can by design not account for the influence of different initial states and goals.

### Why FDR and not STRIPS?

- **Richness of Variables:** FDR uses multi-valued variables instead of binary propositions, leading to a more compact and readable graph.
- **Structural Clarity:** Causal Graphs in FDR explicitly show dependencies between state variables, which are often hidden or fragmented in STRIPS.
- **Heuristic Efficiency:** Modern planners use FDR Causal Graphs to identify cycles and hierarchical dependencies, facilitating more effective state-space search.

### 5.4.1 Domain Transition Graphs

**Definition 68** Let  $\Pi = (V, A, c, I, G)$  be an FDR planning task, and let  $v \in V$ . The domain transition graph (DTG) of  $v$  is the labeled directed graph  $DTG(v, \Pi)$  with vertices  $D_v$  and an arc  $(d, d')$  labeled with action  $a \in A$  whenever either:

- (i)  $pre_a(v) = d$  and  $eff_a(v) = d'$ , or
- (ii)  $pre_a(v)$  is not defined and  $eff_a(v) = d'$ .

We refer to  $(d, d')$  as a value transition of  $v$ . We write  $d \xrightarrow{a} \varphi d'$  where  $\varphi = pre_a \setminus \{v = d\}$  is the (outside) condition. Where not relevant, we omit “ $a$ ” and/or “ $\varphi$ ”.

A DTG capture “where” a variable can go and how, and which values can assume.

**Definition 69 (Invertible Value Transition)** Let  $\Pi = (V, A, c, I, G)$  be an FDR planning task, and let  $v \in V$ , and let  $d \rightarrow_\varphi d'$  to be a value transition of  $v$ . We say that  $d \rightarrow_\varphi d'$  is **invertible** if there exists a value transition  $d' \rightarrow_{\varphi'} d$  where  $\varphi' \subseteq \varphi$ .

With this definition we emphasize the fact that the DTG captures whether a variable can “go back”.

### 5.4.2 Task Decomposition and Task Simplification

The following fact:

Unconnected parts of the task can be solved separately

is true and is modeled by the following lemma.

**Lemma 7** Let  $\Pi = (V, A, c, I, G)$  be an FDR planning task, and let  $V_1, V_2$  to be a partition of  $V$  such that  $CG(\Pi)$  contains no arc between the two sets. Let

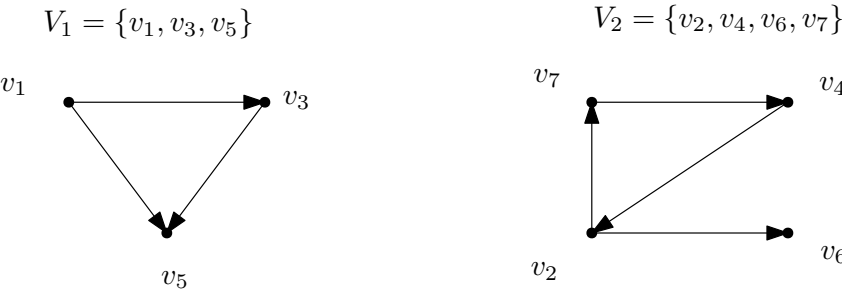
- $\Pi_1$  to be an FDR identical to  $\Pi$  except that includes only the variables  $V_1$ ,  $I$  and  $G$  are restricted on  $V_1$  and for without all the actions  $a$  where either  $pre_a$  or  $eff_a$  are defined on a variable in  $V_2$ .
- $\Pi_2$  to be an FDR identical to  $\Pi$  except that includes only the variables  $V_2$ ,  $I$  and  $G$  are restricted on  $V_2$  and for without all the actions  $a$  where either  $pre_a$  or  $eff_a$  are defined on a variable in  $V_1$ .

then, if

- $\vec{a}_1$  is an optimal plan for  $\Pi_1$
- $\vec{a}_2$  is an optimal plan for  $\Pi_2$

then  $\vec{a}_1 \circ \vec{a}_2$  is an optimal plan for  $\Pi$ .

Any plan for  $\Pi$  can be separated into independent sub-sequences touching  $V_1$  respectively  $V_2$ , corresponding to plans for  $\Pi_1$  respectively  $\Pi_2$ . The Lemma also holds for non-optimal plan.



Let's now consider an FDR task planning  $\Pi$  where the causal graph have a vertex  $v$  such that

- $v$  is a leaf in the causal graph
- $G(v)$  is not defined, so the value of  $v$  is independent from the goal

Since  $v$  is a leaf in  $CG(\Pi)$ , the actions that do affect  $v$  affect no other variables, and the actions that do not affect  $v$  do not have preconditions on  $v$ . So  $v$  is a "client": it moves only for its own purpose. But if  $v$  has no own goal, then it has no "purpose". Thus, denoting by  $\Pi'$  the modified task where  $v$  has been removed, any (optimal) plan for  $\Pi'$  is an (optimal) plan for  $\Pi$ .

Let's now consider a variable with a domain transition graph that is invertible and connected (so the variable  $v$  can "move" around and assume any value), and is a root vertex in the causal graph, we can remove  $v$  from  $\Pi$  and obtain a smaller problem  $\Pi'$  to solve, with an optimal plan  $\vec{a}$  for  $\Pi'$ , then extend  $\vec{a}$  with a move sequence for  $v$  that achieves all preconditions on  $v$  needed to moves the variable to their goal value (if any).

Such variable  $v$  is called "servant", thanks to its connected and invertible DTG, it can always go wherever it is needed. The optimal plan for  $\Pi'$  ignores the cost of moving  $v$  so may incur unnecessarily high costs on  $v$ . This idea is expressed in the following Lemma.

**Lemma 8** *Let  $\Pi = (V, A, c, I, G)$  be an FDR planning task, and let  $v \in V$  to be a root vertex in  $CG(\Pi)$  such that  $DTG(v, \Pi)$  is connected and all value transitions of  $v$  is invertible. Let  $\Pi'$  to be identical to  $\Pi$  except that*

- *we remove  $v$*
- *we restrict  $I$  and  $G$  to  $V \setminus \{v\}$*
- *remove any assignment to  $v$  from all action preconditions*
- *remove all actions  $a$  where  $eff_a(v)$  is defined*

*then, for any plan  $\vec{a}$  for  $\Pi'$ , a plan  $\Pi$  can be obtained in polynomial time in  $|\Pi|$  and  $|\vec{a}|$ .*

**Theorem 23** *If an FDR have an acyclic causal graph and all the variables have all value transitions invertible, the task of finding a path can be decided in polynomial time.*

## CHAPTER

# 6

## ADVANCED SEARCH TECHNIQUES AND HEURISTICS

We treat the planning as a classical search problem, there are three independent choice to make, the first one is the search Space:

- Progression: search forward from initial state to goal. Search states = world states.
- Regression: Search backward from goal to initial state. Search states = sub-goals we would need to achieve.

Then, we have to choose the search algorithm:

- Blind search: DFS, BFS ecc...
- Heuristic search (systematic): A\*
- Heuristic search (local): hill-climbing

The last choice is on the search control:

- Heuristic function (for heuristic cases): Critical-path heuristics, delete-relaxation heuristics, abstraction heuristics, landmarks heuristics, ...
- Pruning techniques: Helpful actions pruning, symmetry elimination, dominance pruning, partial-order reduction.

### 6.1 Progression and Regression

A classical search space is defined by the initial state, the goal state, and the transitions between the states generated by an applicable action.

**Definition 70** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task, the **progression search space** of  $\Pi$  is given by

- $InitialState() = I$
- $GoalState(s) = \begin{cases} true & if G \subseteq s \\ false & otherwise \end{cases}$
- $ChildState(s, a) = \{s' \mid \Theta_\Pi \text{ has the transition } s \xrightarrow{a} s'\}$

The same definition applies to FDR tasks. Start from initial state, and apply actions until a goal state is reached.

**Definition 71** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task, the **regression search space** of  $\Pi$  is given by

- $InitialState() = G$
- $GoalState(g) = \begin{cases} \text{true if } g \subseteq I \\ \text{false otherwise} \end{cases}$
- $ChildState(g, a) = \{g' \mid g' = \text{regr}(g, a)\}$

The definition of the regression  $\text{regr}(g, a)$  for an action depends on whether you consider FDR or STRIPS and will be given later. The following condition on  $g' = \text{regr}(g, a)$  is required

$$g' = \text{regr}(g, a) \implies \forall s' \text{ s.t. } s' \models g' \text{ we have } s'[a] = s \text{ where } s \models g. \quad (6.1)$$

we recall the previous definition:

$$s[a] = \begin{cases} \text{undefined if } a \notin A[s] \\ \text{eff}_a(v) \text{ if } v \in V[\text{eff}_a] \\ s(v) \text{ if } v \notin V[\text{eff}_a] \end{cases}$$

**Definition 72 (FDR Regression)** Let  $(V, A, c, I, G)$  be an FDR planning task, let  $g$  to be a partial variable assignment and  $a \in A$  an action, we say that  $g$  is **regressable** over  $a$  if

- (i)  $\text{eff}_a \cap g \neq \emptyset$
- (ii) there is no  $v \in V$  such that  $v \in V[\text{eff}_a] \cap V[g]$  and  $\text{eff}_a(v) \neq g(v)$
- (iii) there is no  $v \in V$  such that  $v \notin V[\text{eff}_a]$ ,  $v \in V[\text{pre}_a]$  and  $\text{pre}_a(v) \neq g(v)$ .

if these conditions are met, then the **regression** of  $g$  over  $a$  is

$$\text{regr}(g, a) = \{g \setminus \text{eff}_a\} \cup \text{pre}_a \quad (6.2)$$

otherwise the regression is undefined:

$$\text{regr}(g, a) = \perp. \quad (6.3)$$

The three conditions of Definition 6.1 affirms that

- (i) The action must serve some purpose. There must be at least one variable in common between the effects of the action ( $\text{eff}_a$ ) and the goal  $g$  ( $\text{eff}_a \cap g \neq \emptyset$ ). If the action doesn't accomplish any part of the goal, there's no point in regressing on it.
- (ii) The action must not destroy the goal. If the action assigns a value to a variable that is also present in the target  $g$ , that value must be identical. You can't use an action that makes a variable "red" if the goal is for it to be "blue".
- (iii) There must be no internal conflicts. If a variable appears in both the action's preconditions ( $\text{pre}_a$ ) and the goal  $g$ , but the action does not change that variable, then the two values must match. Otherwise, you would ask the variable to have two different values at the same time at the same instant in time before the action.

**Proposition 8** The regression of Definition 6.1 satisfies the condition on Equation (6.1).

**Definition 73 (STRIPS Regression)** Let  $(P, A, c, I, G)$  be a STRIPS planning task, let  $g \subseteq P$  and  $a \in A$ , we say that  $g$  is **regressable** over  $a$  if

- (i)  $\text{add}_a \cap g \neq \emptyset$ : The action must add at least one facts/proposition that is part of  $g$
- (ii)  $\text{del}_a \cap g = \emptyset$ . The action must not delete anything required in  $g$ .

In that case the regression of  $g$  over  $a$  is

$$\text{regr}(g, a) = \{g \setminus \text{add}_a\} \cup \text{pre}_a \quad (6.4)$$

otherwise the regression is undefined:

$$\text{regr}(g, a) = \perp. \quad (6.5)$$

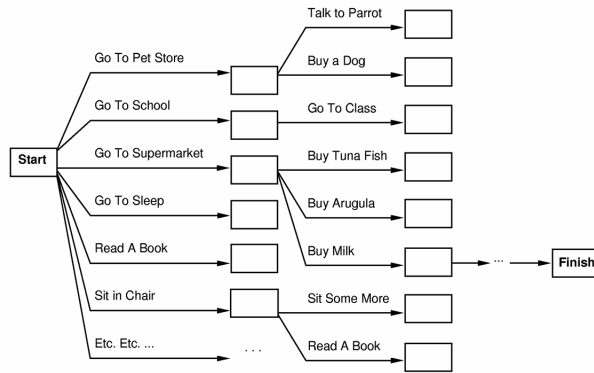
**Proposition 9** *The regression of Definition 73 satisfies the condition on Equation (6.1).*

We want to emphasize the difference between the STRIPS regression and the FDR regression

- the condition (ii) of the FDR regression prevents there from being assignments of variables that contradict those of the objective, whereas for STRIPS regression, only immediate cancellations of the action are considered.
- the condition (iii) in the STRIPS regression is missing because in STRIPS there is no possibility for a subgoal to be "self-contradictory".

STRIPS regression is "weaker" because it cannot see certain logical conflicts that FDR instead captures naturally thanks to the finite domains of the variables.

Observe how the progression doesn't know what's "relevant" to what contributes to reaching the goal.



So a heuristic function is needed in such case. Differently, regression explores only solvable states, but may explore unreachable ones. Regression doesn't know what's "reachable", i.e. what contributes to reaching the initial state. Also in this case a heuristic function is needed to guide the search towards the initial state.

Given that, for the rest of this course we will use *progression*, since:

- Regression has in the past often had serious trouble getting lost in a lot of solvable but unreachable states. Reachable dead end states tend to be less frequent in practice.
- Progression allows easy formulation of searches for more complex planning formalisms (numbers, durations, uncertainty).
- Basically all current heuristic search planners, including Fast Downward use progression.

## 6.2 Heuristic Search

We already know that a heuristic function  $h$  for a search space estimates the cost of an optimal path from a given state  $s$  to the goal, given such function  $h$ , we might search among the states by expanding the node with the smallest value on  $h$ .

The reader is advised to review the definitions of a heuristic function given in the 2.4 section. We will see later also inadmissible heuristics, that typically arise as approximations of admissible heuristics that are too costly to compute.

**Definition 74 (Domination)** Let  $\Pi$  be a planning task, and let  $h$  and  $h'$  to be two admissible heuristic functions, we say that  $h'$  **dominates**  $h$  if

$$h(s) \leq h'$$

for all the states  $s$ .

If  $h'$  dominates  $h$  then,  $h'$  provides a lower bound at least as good as  $h$ .

**Definition 75 (Additivity)** Let  $\Pi$  be a planning task, and let  $h_1, h_2 \dots h_n$  to be  $n$  admissible heuristic functions. The functions  $h_1, h_2 \dots h_n$  are additive if

$$h = h_1 + h_2 + \dots + h_n$$

is an admissible function, i.e. we have:

$$h_1(s) + h_2(s) + \dots + h_n(s) \leq h^*(s) \quad \forall s \quad (6.6)$$

where  $h^*$  is the perfect heuristic.

It's clearly better to consider the sum of a sequence of heuristic since it always dominates any heuristic of the sum.

There exists a modified version of the A\* algorithm 9, called **Weighted A\***.

---

### Algorithm 31 Weighted-A\*

---

**Require:** A problem  $\Pi$

```

 $n \leftarrow$  a node  $n$  such that  $n.State = \Pi.InitialState$ 
 $frontier \leftarrow$  a priority queue ordered by ascending  $g + W \cdot h$ 
 $explored \leftarrow$  empty set of states
while True do
    if  $frontier$  is empty Then Return failure
     $n \leftarrow Pop(frontier)$ 
    if  $n.State$  is the goal state Then Return Solution( $s$ )
     $explored \leftarrow explored \cup n.State$ 
    for each action  $a$  in  $\Pi.Actions(n.State)$  do
         $n' \leftarrow ChildNode(\Pi, n, a)$ 
        if  $n'.State \notin explored \cup States(frontier)$  then
            insert( $n', W \cdot h(n') + g(n')$ ,  $frontier$ )
        else if  $\exists n''$  s.t.  $n''.State = n'.State$  and  $g(n') < g(n'')$ 
            Replace  $n''$  in  $frontier$  with  $n'$ 
        end if
    end for
end while

```

---

This algorithm explores the states by increasing weighted-plan-cost estimate  $g + W \cdot h$ . The weight  $W \in \mathbb{R}^+ \cup \{0\}$  is an algorithm parameter:

- if  $W = 0$  we are not using the heuristic function
- if  $W = 1$  the algorithm is A\*
- for  $W \rightarrow \infty$  the algorithm tends to behave like the greedy best-first search algorithm 8.

We recall the hill climbing algorithm:

The Hill climbing algorithm 32 can easily get stuck in a local minima where immediate improvements of  $h(n)$  are not possible. A possible improvement is to perform a BFS to find the node on which  $h$  decreases:

This algorithm 33 is still not optimal:

- If the problem contains dead ends that are not recognized by the heuristic, the algorithm may crash without finding a solution.
- It is only complete if it is guaranteed that no such states exist and that a goal is always reachable with a lower heuristic value.

**Algorithm 32** Hill-Climbing**Require:** A problem  $\Pi$ 

```

 $n \leftarrow$  a node  $n$  with  $n.State = \Pi.InitialState$ 
while True do
    if  $n.State$  is the goal state then return  $n$ 
     $N \leftarrow$  set of all child nodes of  $n$ 
     $n \leftarrow \arg \min_{n \in N} h(n)$ 
    if stopping condition are met then return  $n$ 
end while

```

**Algorithm 33** Enforced Hill-Climbing**Require:** A problem  $\Pi$ 

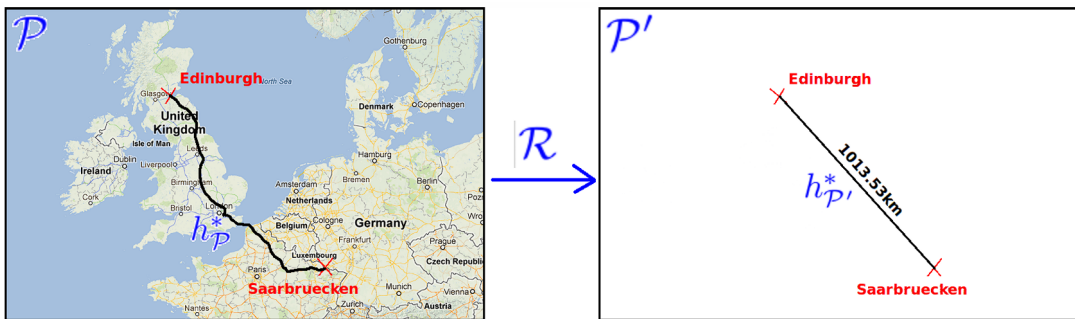
```

 $n \leftarrow$  a node  $n$  with  $n.State = \Pi.InitialState$ 
while True do
    if  $n.State$  is the goal state then return  $n$ 
    perform a BFS to find  $n'$  s.t.  $h(n') < h(n)$ 
     $n \leftarrow n'$ 
    if stopping condition are met then return  $n$ 
end while

```

**6.2.1 Computing the Heuristic Function**In general, how do we choose the function  $h$ ?

- Let  $\mathcal{P}$  to be a class of problems that we want to solve, whose perfect heuristic is denoted  $h_{\mathcal{P}}^*$ .
- It can be defined a class  $\mathcal{P}'$  of simpler problems that are a relaxation of  $\mathcal{P}$ , whose perfect heuristic is  $h_{\mathcal{P}'}^*$ .
- We define a **relaxation mapping**  $\mathcal{R}$  that maps instances of  $\Pi \in \mathcal{P}$  into instances of  $\Pi' \in \mathcal{P}'$ .
- Given a problem  $\Pi \in \mathcal{P}$ , we let  $\Pi' = \mathcal{R}(\Pi)$  and we use the function  $h_{\mathcal{P}'}^*(\Pi')$  to estimate  $h_{\mathcal{P}}^*(\Pi)$ .



- Problem class  $\mathcal{P}$ : Route finding.
- Perfect heuristic  $h_{\mathcal{P}}^*$  for  $\mathcal{P}$ : Length of a shortest route.
- Simpler problem class  $\mathcal{P}'$ : Route finding on an empty map.
- Perfect heuristic  $h_{\mathcal{P}'}^*$  for  $\mathcal{P}'$ : Straight-line distance.
- Transformation  $\mathcal{R}$ : Throw away the map.

**6.3 Critical Path Heuristics**

Critical path heuristics are a family of methods to relax planning tasks, and thus automatically compute heuristic functions  $h$ .

**Definition 76** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, the **perfect regression heuristic**  $r^*$  for  $\Pi$  is the function  $r^*(s, g)$  where

$$r^*(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \min_{a \in A^*} c(a) + r^*(s, \text{regr}(g, a)) & \text{otherwise} \end{cases} \quad (6.7)$$

where  $A^* = \{a \in A : \text{regr}(g, a) \neq \perp\}$ .

Is omitted that this is the perfect heuristic when  $g = G$  (the goal) and is evaluated by letting  $g = G$  fixed. The heuristic  $r^*$  is defined as the cost of reaching a set of goals  $g$  starting from a state  $s$ :

- Base case: If the goal  $g$  is already contained in the state  $s$ , the cost is 0.
- Recursive case: If the objective is not satisfied, the cost is given by choosing the best action  $a$  (the one that minimizes the sum) among all those that can "generate" a part of the objective through regression. The total cost is the cost of action  $c(a)$  plus the cost of achieving the new subgoals needed to apply that action.

**Proposition 10** For a STRIPS planning task  $\Pi$  the perfect heuristic  $h^*$  is  $r^*$ .

**Definition 77** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, the **critical path heuristic**  $h^1$  for  $\Pi$  is the function  $h^1(s, g)$  defined as follows

$$h^1(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \min_{a \in A^*} c(a) + h^1(s, \text{regr}(g, a)) & \text{if } |g| = 1 \\ \max_{g' \subseteq g} h^1(s, \{g'\}) & \text{if } |g| > 1 \end{cases} \quad (6.8)$$

where  $A^* = \{a \in A : \text{regr}(g, a) \neq \perp\}$ .

Is omitted that this is the perfect heuristic when  $g = G$  (the goal) and is evaluated by letting  $g = G$  fixed. This function is similar to  $h^*$ :

- For singleton subgoals  $g$ , use regression as in  $h^*$ .
- For subgoal sets  $g$ , use the cost of the most costly singleton subgoal  $g' \in g$ .

It follows the critical path, defined as the cheapest path trough the most costly sub goal  $g'$ . This heuristic can be generalized by considering the critical path as the cheapest path trough the most costly  $m$  sub goals.

**Definition 78** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, the **critical path heuristic**  $h^m$  for  $\Pi$  is the function  $h^m(s, g)$  defined as follows

$$h^m(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \min_{a \in A^*} c(a) + h^m(s, \text{regr}(g, a)) & \text{if } |g| \leq m \\ \max_{g' \subseteq g, |g'|=m} h^m(s, \{g'\}) & \text{if } |g| > m \end{cases} \quad (6.9)$$

where  $A^* = \{a \in A : \text{regr}(g, a) \neq \perp\}$ .

For a fixed  $m$ , the value of  $h^m(s, g)$  can be computed in polynomial time in the size of  $\Pi$ .

**Proposition 11**  $h^m$  is consistent and goal-aware, hence is admissible and safe.

Intuition:  $h^m$  is admissible because it is always more difficult to achieve larger subgoals (so  $m$ -subsets can only be cheaper)

**Proposition 12**  $h^{m+1}$  dominates  $h^m$ ,  $h^m$  gets more accurate as  $m$  grows.

**Proposition 13** There exists  $m$  such that  $h^m = r^*$ .

*Proof:* Let  $m = |P|$ , in that case since  $g \subseteq P$  we have  $|g| \leq m$  so  $h^m = r^*$ .

### 6.3.1 Computing $h^m$

The function  $h^m$  can be computed in polynomial time through dynamic programming, but we have to give an iterative definition. We will denote  $h_i^m$  the value of the iterative  $h^m$  at the iteration  $i$ . This function is defined as follows:

$$h_0^m(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \infty & \text{otherwise} \end{cases} \quad (6.10)$$

Given  $h_i^m(s, g)$ , the next iteration is:

$$h_{i+1}^m(s, g) = \begin{cases} \min \left\{ h_i^m(s, g), \min_{a \in A, \text{regr}(g, a) \neq \perp} c(a) + h_i^m(s, \text{regr}(g, a)) \right\} & \text{if } |g| \leq m \\ \max_{g' \subseteq g, |g'|=m} h_{i+1}^m(s, g') & \text{if } |g| > m \end{cases} \quad (6.11)$$

**Proposition 14** *the series  $\{h_i^m\}$  converges to  $h^m$ .*

We give a proof sketch, if  $h_{i+1}^m(s, g) \neq h_i^m(s, g)$  then  $h_{i+1}^m(s, g) < h_i^m(s, g)$ , since there exists  $\epsilon \in \mathbb{R}^+$  such that

$$h_i^m(s, g) - h_{i+1}^m(s, g) \geq \epsilon \quad (6.12)$$

at some point  $h_i^m$  will stop decreasing. If  $h_{i+1}^m(s, g) = h_i^m(s, g)$  then  $h_i^m$  satisfy the  $h^m$  definition, and no other function greater than  $h_i^m$  at any point can satisfy that definition.

To compute  $h_i^m$  dynamic programming is involved. With *dynamic programming*, we mean that paradigm of algorithm design, which, in order to find a final solution to a problem, it considers a series of sub-problems minors who will then be useful in deciding the aforementioned final solution.

#### Dynamic Programming Algorithm

```

new table  $T_0^m(g)$ , for all  $g \subseteq P$  with  $|g| \leq m$ 
For all  $g \subseteq P$  with  $|g| \leq m$ :  $T_0^m(g) := \begin{cases} 0 & g \subseteq s \\ \infty & \text{otherwise} \end{cases}$ 
fn  $Cost_i(g) := \begin{cases} T_i^m(g) & |g| \leq m \\ \max_{g' \subseteq g, |g'|=m} T_i^m(g') & |g| > m \end{cases}$ 
fn  $Next_i(g) := \min[Cost_i(g), \min_{a \in A, \text{regr}(g, a) \neq \perp} c(a) + Cost_i(\text{regr}(g, a))]$ 
i := 0
do forever:
  new table  $T_{i+1}^m(g)$ , for all  $g \subseteq P$  with  $|g| \leq m$ 
  For all  $g \subseteq P$  with  $|g| \leq m$ :  $T_{i+1}^m(g) := Next_i(g)$ 
  if  $T_{i+1}^m = T_i^m$  then stop endif
  i := i + 1
enddo

```

It is easy to prove that  $h_i^m(s, g) = Cost_i(g)$  for all  $i$ . This table computation always first finds the shortest path to achieve a subgoal  $g$ . Hence, with unit action costs, the value of  $g$  is fixed once it becomes less than  $\infty$ , and equals the  $i$  where that happens. With non-unit action costs, neither is true.

For any fixed  $m$ ,  $h_m$  can be computed in polynomial time.

In practice, only  $m = 1, 2$  are used, higher values of  $m$  are infeasible.

### 6.3.2 Graphplan Representation

The critical path heuristic  $h^m$  can be computed with a planning graph, we consider the case  $m = 1$  and the Algorithm 34.

This algorithm computes a sequence  $\{F_i\}$ , this induce a function  $h_{PG}^1$  called **1-planning graph heuristic** defined as follows

$$h_{PG}^1 = \min\{i \mid s \subseteq F_i\}. \quad (6.13)$$

**Algorithm 34** 1-Planning-Graph

---

```

1:  $F_0 = s$ ,  $i = 0$ 
2: while  $G \subseteq F_i$  do
3:    $A_i = \{a \in A \mid pre_a \subseteq F_i\}$ 
4:    $F_{i+1} = F_i \cup \left\{ \bigcup_{a \in A_i} add_a \right\}$ 
5:   if  $F_{i+1} = F_i$  then
6:     return
7:   end if
8:    $i = i + 1$ 
9: end while

```

---

1. **Initialization:** Set the initial fact layer  $F_0$  to the initial state  $s$ . Initialize the layer counter  $i = 0$ .
2. **Goal Check:** Determine if the goal set  $G$  is a subset of the current fact layer  $F_i$  ( $G \subseteq F_i$ ). If it is, the algorithm terminates successfully.
3. **Action Selection ( $A_i$ ):** Identify the set of all actions  $A_i$  whose preconditions are satisfied by the facts in the current layer:

$$A_i := \{a \in A \mid pre_a \subseteq F_i\}$$

4. **Fact Expansion ( $F_{i+1}$ ):** Generate the next fact layer by taking the union of the current facts and all additive effects of the actions in  $A_i$ :

$$F_{i+1} := F_i \cup \bigcup_{a \in A_i} add_a$$

5. **Fixed-Point Check:** Check for convergence. If  $F_{i+1} = F_i$ , the graph has "leveled off". If the goal has not been reached by this point, it is unreachable, and the algorithm stops.
6. **Iteration:** Increment  $i$  and repeat from Step 2.

**Proposition 15** For a STRIPS planning task with unit cost,  $h_{PG}^1 = h^1$ .

The algorithm 35 is the generalization of a generic  $m \in \mathbb{N}$ .

**Algorithm 35**  $m$ -Planning-Graph

---

```

1:  $F_0 = s$ ,  $i = 0$ ,  $M = \emptyset$ 

2: function REACHED( $i, g$ )
3:   if  $g \subseteq F_i \wedge \exists g' \in M_i$  such that  $g' \subseteq g$  then
4:     return True
5:   end if
6:   return False
7: end function

8: while not REACHED( $i, G$ ) do
9:    $A_i = \{a \in A \mid \text{REACHED}(i, pre_a)\}$ 
10:   $F_{i+1} = F_i \cup \left\{ \bigcup_{a \in A_i} add_a \right\}$ 
11:   $M_{i+1} = \{g \subseteq P : |g| \leq m \wedge \forall a \in A_i, \text{not REACHED}(i, regr(g, a))\}$ 
12:  if  $F_{i+1} = F_i$  and  $M_{i+1} = M_i$  then
13:    return
14:  end if
15:   $i = i + 1$ 
16: end while

```

---

Intuition: all  $m$ -subsets  $g$  of  $F_i$  are reachable within  $i$  steps, except for those  $g$  listed in  $M_i$ .

All the algorithms and arguments presented applies also for FDR.

## 6.4 Delete Relaxation Heuristics

We will present some formal definition and states some simple properties that immediately result in a simple "greedy" heuristic.

**Definition 79 (Delete Relaxation)** For a STRIPS action  $a$ , we denote  $a^+$  the **delete relaxed action** (or short **relaxed action**), defined as follows:

$$\begin{aligned} pre_{a^+} &= pre_a \\ add_{a^+} &= add_a \\ del_{a^+} &= \emptyset \end{aligned}$$

Given a set of action  $A$ , the relaxed actions  $A^+$  are

$$A^+ = \{a^+ \mid a \in A\} \quad (6.14)$$

and we denote  $\vec{a}^+$  a sequence of relaxed actions

$$\vec{a}^+ = \langle a_1^+ \dots a_n^+ \rangle \quad (6.15)$$

relative to the sequence of actions

$$\vec{a} = \langle a_1 \dots a_n \rangle \quad (6.16)$$

Given a STRIPS planning task  $\Pi = (P, A, c, I, G)$ , we denote  $\Pi^+ = (P, A^+, c, I, G)$  the corresponding **relaxed planning task**.

A delete relaxed planning task (denoted as  $\Pi^+$ ) is a simplified version of a standard planning problem where actions are modified to never remove facts from the current state.

**Definition 80** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, and let  $s$  to be a state. A **relaxed plan** for  $s$  is a plan<sup>1</sup> for  $\Pi_s^+$  where  $\Pi_s = (P, A, c, s, G)$  so  $\Pi_s^+ = (P, A^+, c, s, G)$ .

**Definition 81** A state  $s'$  for a STRIPS planning task **dominates**  $s$  if  $s \subseteq s'$ .

A state (set of facts) dominates another state if it has "more facts true".

**Proposition 16** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, and let  $s, s'$  be states where  $s'$  dominates  $s$ , then

1. if  $s$  is a goal then  $s'$  is a goal as well.
2. if  $\vec{a}$  is applicable in  $s$  then is applicable in  $s'$ .
3.  $s'[\vec{a}]$  dominates  $s[\vec{a}]$ .

The proof is trivial. This proposition says that "it is always better to have more facts true". The following proposition is crucial.

**Proposition 17** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, let  $s \subseteq P$  be a state and  $a \in A$  an action applicable in  $s$ , then

- $s[a^+]$  dominates  $s$
- if  $s'$  dominates  $s$ , then  $s'[a^+]$  dominates  $s[a]$

The first follows from the facts that  $a^+$  doesn't delete any fact from  $s$ , so it can only add facts. The second is more trivial, since  $s'$  already have all the facts in  $s$

$$s \subseteq s'$$

and  $s'[a^+]$  dominates  $s'$ , so

$$s \subseteq s' \subseteq s'[a^+]$$

then,  $s[a]$  may loose some facts and get added more, but all the facts that it loses are in  $s'$ , and all the facts that it obtain are in  $s'[a^+]$ , so the proposition holds. The meaning of the proposition is

---

<sup>1</sup>this plan may be optimal or not. If the plan is optimal, also the relaxed plan is optimal.

any real plan also works in the relaxed world.

**Proposition 18** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, let  $s \subseteq P$  be a state and  $\vec{a}$  a plan for  $\Pi_s$ . Then,  $\vec{a}^+$  is a relaxed plan for  $s$ .

So, if we apply a relaxed action we can only have more facts true, and that cannot render the task unsolvable, so we can keep applying relaxed actions until the goal is reached to get a relaxed plan  $\vec{a}^+$ , as done in algorithm 36.

---

**Algorithm 36** Greedy Relaxed Planning

---

```

1:  $s^+ = s, \vec{a}^+ = \langle \rangle$ 
2: while  $G \not\subseteq s^+$  do
3:   if  $\exists a \in A$  such that  $pre_a \subseteq s^+$  and  $s^+ \llbracket a^+ \rrbracket \neq s^+$  then
4:     select one such  $a$ 
5:      $s^+ = s^+ \llbracket a^+ \rrbracket$ 
6:      $\vec{a}^+ = \vec{a}^+ \circ \langle a^+ \rangle$ 
7:   else
8:     return  $\Pi_s^+$  is unsolvable
9:   end if
10: end while
```

---

This algorithm is sound, complete and terminates in polynomial time in the size of  $\Pi$ . It is easy to decide whether a relaxed plan exists.

This kind of relaxation is an over-approximation of the problem, in fact, if we use a greedy relaxed plan to generate a heuristic function  $h$ , we wouldn't have an accurate function.

- In the search state  $s$ , during forward search, we run the greedy relaxed planning for  $\Pi_s^+$
- we set  $h(s)$  to the cost of  $\vec{a}^+$ , or  $\infty$  if  $\Pi_s^+$  is unsolvable.

TO be accurate, a heuristic function needs to approximate the minimum effort needed to reach the goal, the greedy relaxed plan may select arbitrary actions that aren't relevant at all, over-estimating dramatically. The next definition describes the function we need.

**Definition 82** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task with state space  $\theta_\Pi = (S, A, c, T, I, G)$ . The **optimal delete relaxation heuristic**  $h^+$  for  $\Pi$  is the function  $h^+ : S \rightarrow \mathbb{R}^+ \cup \{0, \infty\}$  where  $h^+(s)$  is defined as the cost of an optimal relaxed plan for  $s$ .

$h^+$  defines the minimum effort to reach the goal under delete relaxation.

**Proposition 19** The optimal delete relaxation heuristic  $h^+$  for a state space is consistent, and thus admissible, safe and goal aware.

**Proposition 20** For the task to find the shortest path between two nodes in a graph, the optimal delete relaxation heuristic is the perfect heuristic.

**Definition 83** By  $PlanOpt^+$  we denote the problem of deciding, given a STRIPS planning task  $\Pi = (P, A, c, I, G)$  and  $B \in \mathbb{R}^+ \cup \{0\}$  whether there exists a relaxed plan for  $\Pi$  whose cost is at most  $B$ .

It is true that, by computing  $h^+$ , we would solve  $PlanOpt^+$ .

**Theorem 24**  $PlanOpt^+$  is NP-complete.

From the theorem follows that computing  $h^+$  is an hard task, so instead of directly computing it, we may consider an approximation.

*The delete relaxation heuristic we want is  $h^+$ . Unfortunately, this is hard to compute so the computational overhead is very likely to be prohibitive. All implemented systems using the delete relaxation approximate  $h^+$  in one or the other way.*

### 6.4.1 The Additive and Max Heuristic

**Definition 84** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task, the **additive heuristic**  $h^{\text{add}}$  for  $\Pi$  is the function  $h^{\text{add}}(s, g)$  where  $s, g$  are set of facts and

$$h^{\text{add}}(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \min_{a \in A, g' \in \text{add}_a} c(a) + h^{\text{add}}(s, \text{pre}_a) & \text{if } g = \{g'\} \\ \sum_{g' \in g} h^{\text{add}}(s, \{g'\}) & \text{if } |g| > 1 \end{cases} \quad (6.17)$$

**Definition 85** Let  $\Pi = (P, A, c, I, G)$  to be a STRIPS planning task, the **max heuristic**  $h^{\text{max}}$  for  $\Pi$  is the function  $h^{\text{max}}(s, g)$  where  $s, g$  are set of facts and

$$h^{\text{max}}(s, g) = \begin{cases} 0 & \text{if } g \subseteq s \\ \min_{a \in A, g' \in \text{add}_a} c(a) + h^{\text{max}}(s, \text{pre}_a) & \text{if } g = \{g'\} \\ \max_{g' \in g} h^{\text{max}}(s, \{g'\}) & \text{if } |g| > 1 \end{cases} \quad (6.18)$$

The provided definitions describe two common relaxation-based heuristics used in STRIPS automated planning to estimate the cost of reaching a goal state  $G$  from a current state  $s$ . Both functions decompose the planning problem into achieving individual subgoals but handle the combination of those costs differently.

**Additive Heuristic ( $h^{\text{add}}$ ):** This function estimates the total cost by **summing** the individual costs of achieving every atom in the goal set. It assumes subgoals are independent, which often leads to overestimation (making it non-admissible) because it ignores "positive interactions" where one action might satisfy multiple subgoals simultaneously.

**Max Heuristic ( $h^{\text{max}}$ ):** This function estimates the cost by taking the **maximum** cost among all individual subgoals. It represents the cost of the most "difficult" subgoal to achieve. Unlike  $h^{\text{add}}$ ,  $h^{\text{max}}$  is admissible (it never overestimates the true cost) because the total plan cost must be at least as great as the cost to reach its hardest component, but it is often less informative (less "informed") than the additive version.

Both heuristics share a recursive structure for single goals ( $g = \{g'\}$ ), where the cost is determined by the cheapest action  $a$  that can achieve  $g'$ , added to the cost of achieving that action's preconditions  $\text{pre}_a$ .

**Proposition 21** The function  $h^{\text{max}}$  is optimistic:

$$h^{\text{max}} \leq h^+ \implies \quad (6.19)$$

$$h^{\text{max}} \leq h^* \quad (6.20)$$

$h^{\text{max}}$  simplifies relaxed planning by assuming that, to achieve a set  $g$  of subgoals, it suffices to achieve the single most costly  $g' \in g$ . Actual relaxed planning, i.e.  $h^+$ , can only be more expensive.

**Proposition 22** The function  $h^{\text{add}}$  is pessimistic:

$$h^{\text{add}} \geq h^+ \implies \quad (6.21)$$

there exists a  $\Pi$  and  $s$  such that  $h^{\text{add}}(s) > h^*(s)$ .

$h^{\text{add}}$  simplifies relaxed planning by assuming that, to achieve a set  $g$  of subgoals, we must achieve every  $g' \in g$  separately. Actual relaxed planning, i.e.  $h^+$ , can only be less expensive.

Both  $h^{\text{max}}$  and  $h^{\text{add}}$  approximate  $h^+$  by assuming that singleton subgoal facts are achieved independently.  $h^{\text{max}}$  estimates optimistically by the most costly singleton subgoal,  $h^{\text{add}}$  estimates pessimistically by summing over all singleton subgoals.

**Proposition 23** For all STRIPS planning task  $\Pi$ , given  $h^+, h^{\text{max}}$  and  $h^{\text{add}}$ , let  $s$  to be a state, it holds that:

$$h^+(s) = \infty \iff h^{\text{max}}(s) = \infty \iff h^{\text{add}}(s) = \infty \quad (6.22)$$

*Proof:* The proof is structured into two main observations:

1. *Equivalence between  $h^{\max}$  and  $h^{\text{add}}$  regarding infinity:* The values  $h^{\max}$  and  $h^{\text{add}}$  agree on states with infinite heuristic value because their only operational difference lies in the choice of the *max* vs.  $\sum$  (summation) operators.

- If any component of a sum is  $\infty$ , the sum is  $\infty$ .
- If any component of a set is  $\infty$ , the maximum is  $\infty$ .

Therefore, the use of these different operators does not affect the property of being infinite.

2. *Equivalence with  $h^+$ :*

- *Direction ( $\implies$ ):*  $h^+(s) < \infty$  implies  $h^{\max}(s) < \infty$  because  $h^{\max}$  is an admissible heuristic for the relaxed task, meaning  $h^{\max} \leq h^+$ .
- *Direction ( $\Leftarrow$ ):* Conversely,  $h^{\max}(s) < \infty$  implies  $h^+(s) < \infty$ . If  $h^{\max}$  is finite, it can be used to generate a *closed, well-founded best-supporter function*. From this function, a valid relaxed plan can be extracted, ensuring that  $h^+(s)$  is also finite. ■

States for which no relaxed plan exists are easy to recognize, and that is done by both  $h^{\max}$  and  $h^{\text{add}}$ . Approximation is needed only for the cost of an optimal relaxed plan, if it exists.

Consider the Definition 77 of the function  $h^1$ , its easy to notice that

$$h^1 = h^{\max} \quad (6.23)$$

We already know how to compute  $h^1$  by dynamic programming (we saw the general algorithm for any  $h^m$ ), so we already know how to compute  $h^{\max}$ . To compute  $h^{\text{add}}$  we use the following (similar) algorithm:

#### Dynamic Programming algorithm computing $h^{\text{add}}$ for state $s$

```

new table  $T_0^{\text{add}}(g)$ , for  $g \in P$ 
For all  $g \in P$ :  $T_0^{\text{add}}(g) := \begin{cases} 0 & g \in s \\ \infty & \text{otherwise} \end{cases}$ 
fn  $Cost_i(g) := \begin{cases} T_i^{\text{add}}(g) & |g| = 1 \\ \sum_{g' \in g} T_i^{\text{add}}(g') & |g| > 1 \end{cases}$ 
fn  $Next_i(g) := \min[Cost_i(g), \min_{a \in A, g' \in add_a} c(a) + Cost_i(pre_a)]$ 
do forever:
  new table  $T_{i+1}^{\text{add}}(g)$ , for  $g \in P$ 
  For all  $g \in P$ :  $T_{i+1}^{\text{add}}(g) := Next_i(g)$ 
  if  $T_{i+1}^{\text{add}} = T_i^{\text{add}}$  then stop endif
   $i := i + 1$ 
enddo

```

Both  $h^{\text{add}}$  and  $h^{\max}$  can be computed quickly, but in practice, there are some issues:

- $h^{\max}$  is admissible, but is typically far too optimistic.
- $h^{\text{add}}$  is not admissible, but is typically a lot more informed than  $h^{\max}$
- $h^{\text{add}}$  is sometimes better informed than  $h^+$ , but "for the wrong reasons", rather than accounting for deletes, it overcounts by ignoring positive interactions, i.e., sub-plans shared between subgoals.

#### 6.4.2 The Relaxed Plan Heuristic

We will define the **best supporters functions** for the additive and max heuristic, that are two functions which, for every fact  $p \in P$  returns an action that is deemed to be the cheapest achiever of  $p$  (within the relaxation). Then extract a relaxed plan from that function, by applying it to singleton subgoals and collecting all the actions.

The best-supporter function can be based directly on  $h^{\max}$  or  $h^{\text{add}}$ , simply selecting an action  $a$  achieving  $p$  that minimizes  $[c(a) + \text{cost estimate for } pre_a]$ .

**Definition 86 (Best Supporters)** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, and let  $s$  be a state:

- The  $h^{\max}$  **supporter function**  $bs_s^{\max}$  is the function

$$bs_s^{\max} : \{p \in P \mid 0 < h^{\max}(s, \{p\}) < \infty\} \mapsto A \quad (6.24)$$

defined as follows:

$$bs_s^{\max}(p) = \arg \min_{a \in A, p \in add_a} c(a) + h^{\max}(s, pre_a) \quad (6.25)$$

- The  $h^{add}$  **supporter function**  $bs_s^{add}$  is the function

$$bs_s^{add} : \{p \in P \mid 0 < h^{add}(s, \{p\}) < \infty\} \mapsto A \quad (6.26)$$

defined as follows:

$$bs_s^{add}(p) = \arg \min_{a \in A, p \in add_a} c(a) + h^{add}(s, pre_a) \quad (6.27)$$

The following algorithm is used to construct a relaxed plan ( $RPlan$ ) given a best-supporter function ( $bs$ ). A best-supporter function simply identifies which action is "best" (cheapest) to achieve a specific fact.

#### Relaxed Plan Extraction for state $s$ and best-supporter function $bs$

```

Open := G \ s; Closed := ∅; RPlan := ∅
while Open ≠ ∅ do:
    select g ∈ Open
    Open := Open \ {g}; Closed := Closed ∪ {g};
    RPlan := RPlan ∪ {bs(g)}; Open := Open ∪ (prebs(g) \ (s ∪ Closed))
endwhile
return RPlan

```

Essentially, the algorithm works backward from the goals to the initial state, picking the best actions to satisfy each requirement until every precondition is accounted for. The number of iterations of this algorithm is bounded by  $|P|$ . To make the algorithm work it is important that the following three conditions are met

1. The chosen action must actually produce the desired goal. Formally, for each goal  $g$ , the action indicated as its best supporter must contain  $g$  among its added effects ( $g \in add_{bs(g)}$ ).
2. The function must always be defined for the necessary objectives. In other words,  $bs(g)$  must not be "undefined" to avoid errors in the algorithm.
3. The support must be well-founded. This means that there must be no circular dependencies where support for a goal  $g$  ultimately requires  $g$  itself as a precondition. If this condition is not satisfied, the algorithm would fail to generate a valid plan.

The general definition of a best supporter function is the following.

**Definition 87** Let  $\Pi = (P, A, c, I, G)$  be a STRIPS planning task, and let  $s$  be a state, a **best supporter function** for  $s$  is the partial function

$$bs : P \setminus \{s\} \mapsto A \quad (6.28)$$

such that  $p \in add_a$  whenever  $a = bs(p)$ .

From this definition of  $bs$  we can construct a **support graph** where the vertices are

$$\{P \setminus \{s\}\} \cup A$$

and the edges are

$$\{(a, p) \mid a = bs(p)\} \cup \{(p, a) \mid p \in pre_a\}$$

- We say that  $bs$  is **closed** if  $bs(p)$  is defined for every  $p \in P \setminus \{s\}$  that has a path to a goal  $g \in G$  in the support graph.

- We say that  $bs$  is **well-founded** if the support graph is acyclic.

A support function  $bs$  defined in that way that is closed and well founded satisfies the three conditions to make the *Relaxed Plan Extraction* algorithm works.

**Proposition 24** *The best supporters  $bs_s^{add}$  and  $bs_s^{\max}$  are closed well-founded supporter functions for  $s$ .*

**Proposition 25** *If  $bs$  is closed well-founded supporter function the RPlan returned by the *Relaxed Plan Extraction* algorithm can be sequenced into a relaxed plan  $\vec{a}^+$  for  $s$ .*

**Definition 88** *A heuristic function is called **relaxed plan heuristic**, and is denoted  $h^{FF}$  if, given a state  $s$ , it returns  $\infty$  if no relaxed plan exists, and otherwise returns  $\sum_{a \in RPlan} c(a)$  where  $RPlan$  is the action set returned by the *Relaxed Plan Extraction* algorithm on a closed well-founded best-supporter function for  $s$ .*

**Proposition 26** *For all STRIPS planning tasks  $\Pi$ :*

- $h^{FF} \geq h^+$
- For all states  $s$ ,  $h^+(s) = \infty$  if and only if  $h^{FF}(s) = \infty$ .
- There exist  $\Pi$  and  $s$  so that  $h^{FF}(s) > h^*(s)$ .

*Proof:* The proof consists of three observations:

- $h^{FF} \geq h^+$ : follows directly from the previous slide (since  $h^{FF}$  generates a valid relaxed plan, its cost must be at least that of the optimal relaxed plan  $h^+$ ).
- Agrees with  $h^+$  on  $\infty$ : Direct from definition.
- Inadmissibility: Occurs whenever the best-supporter function  $bs$  makes sub-optimal choices.

In practice,  $h^{FF}$  typically does not over-estimate  $h^*$  (or not by a large amount, anyway), may be inadmissible just like  $h^{add}$ , but for more subtle reasons.  $h^{FF}$  does not over count, i.e., count sub-plans shared between subgoals more than once. This is due to the set union in  $RPlan = RPlan \cup \{bs(g)\}$ .

**Definition 89 (Helpful Actions)** *Let  $h^{FF}$  be a relaxed plan heuristic, let  $s$  be a state, and let  $RPlan$  be the action set returned by the *Relaxed Plan Extraction* on the closed well-founded best-supporter function for  $s$  which underlies  $h^{FF}$ . Then an action  $a$  applicable to  $s$  is called **helpful** if it is contained in  $RPlan$ .*

What about the FDR language?

**Definition 90** *Let  $\Pi = (V, A, c, I, G)$  to be an FDR planning task, we call  $P_V = \{v = d \mid v \in V, d \in D_v\}$  the set of "FDR facts". The **relaxed state space** of  $\Pi$  is the labeled transition system  $\Theta_\Pi^+ = (S^+, L, c, T, I, S^{G^+})$  where*

- the (relaxed) states  $S^+$  are the subsets  $s^+ \in \mathcal{P}(P_V)$ .
- the labels  $L = A$  are the actions of  $\Pi$ , and the cost function  $c$  is the same of  $\Pi$ .
- The transitions are  

$$T = \{s^+ \xrightarrow{a} s'^+ \mid \text{pre}_a \subseteq s^+, s'^+ = s^+ \cup \text{eff}_a\}$$
- the initial state  $I$  is the same of that in  $\Pi$
- the goal states are  

$$S^{G^+} = \{s^+ \in S^+ \mid G \subseteq s^+\}$$

A solution for  $s^+$  in  $\Theta_\Pi^+$  is a **relaxed plan**<sup>2</sup> for  $s^+$ .

Let  $\Theta_\Pi = (S, A, c, T, I, G)$  be the state space of  $\Pi$ . The optimal delete relaxation heuristic  $h^+$  for  $\Pi$  is the function

$$h^+ : S \mapsto \mathbb{R}^+ \cup \{0, \infty\}$$

where  $h^+(s)$  is defined as the cost of an optimal relaxed plan for  $s$ .

FDR states contain exactly one fact for each variable  $v \in V$ . There is no such restriction on FDR relaxed states. We recall the following proposition:

---

<sup>2</sup>if the solution is optimal, the relaxed plan is optimal

**Proposition 27** Let  $\Pi = (V, A, c, I, G)$  be an FDR planning task, and let  $\Pi^{STR}$  be its STRIPS translation. Then  $\Theta_\Pi$  is isomorphic to the sub-system of  $\Theta_{\Pi^{STR}}$  induced by those  $s \subseteq P_V$  where, for each  $v \in V$ ,  $s$  contains exactly one fact of the form  $v = d$ . All other states in  $\Theta_{\Pi^{STR}}$  are unreachable.

Observe how  $\Theta_\Pi^+$  has transition  $s^+ \xrightarrow{a} s'^+$  if and only if  $s^+ \llbracket a^{STR+} \rrbracket = s'^+$  in  $\Pi^{STR}$  (because  $s^+ \llbracket a^{STR+} \rrbracket = s^+ \cup \text{eff}_a$ ).

**Proposition 28** Denote by  $h_\Pi^*$  and  $h_\Pi^+$  the perfect heuristic and the optimal delete relaxation heuristic in  $\Pi$ , and denote by  $h_{\Pi^{STR}}^*$  and  $h_{\Pi^{STR}}^+$  these heuristics in  $\Pi^{STR}$ . Then, for all states  $s$  of  $\Pi$ ,  $h_\Pi^*(s) = h_{\Pi^{STR}}^*(s)$  and  $h_\Pi^+(s) = h_{\Pi^{STR}}^+(s)$ .

Given an FDR task  $\Pi$ , everything we have done here can be done for  $\Pi$  by doing it within  $\Pi^{STR}$ . As a summary:

- The delete relaxation simplifies STRIPS by removing all delete effects of the actions.
- The cost of optimal relaxed plans yields the heuristic function  $h^+$ , which is admissible but hard to compute.
- We can approximate  $h^+$  optimistically by  $h^{\max}$ , and pessimistically by  $h^{\text{add}}$ .  $h^{\max}$  is admissible,  $h^{\text{add}}$  is not.  $h^{\text{add}}$  is typically much more informative, but can suffer from over-counting.
- Either of  $h^{\max}$  or  $h^{\text{add}}$  can be used to generate a closed well-founded best-supporter function, from which we can extract a relaxed plan.
- The resulting relaxed plan heuristic  $h^{\text{FF}}$  does not do over-counting, but otherwise has the same theoretical properties as  $h^{\text{add}}$ ; in practice, it typically does not over-estimate  $h^*$ .
- The delete relaxation can be applied to FDR simply by accumulating variable values, rather than over-writing them. This is formally equivalent to treating variable/value pairs like STRIPS facts.