# 3.5 INFORMED (HEURISTIC) SEARCH STRATEGIES

INFORMED SEARCH This section shows how an **informed search** strategy—one that uses problem-specific knowledge beyond the definition of the problem itself—can find solutions more efficiently than can an uninformed strategy.

FUNCTION

**HEURISTIC** FUNCTION

BEST-FIRST SEARCH The general approach we consider is called **best-first search**. Best-first search is an instance of the general TREE-SEARCH or GRAPH-SEARCH algorithm in which a node is  $\frac{E[X_0][X]}{E[X_0][X]}$  selected for expansion based on an **evaluation function**,  $f(n)$ . The evaluation function is construed as a cost estimate, so the node with the *lowest* evaluation is expanded first. The implementation of best-first graph search is identical to that for uniform-cost search (Figure 3.14), except for the use of f instead of q to order the priority queue.

> The choice of f determines the search strategy. (For example, as Exercise 3.22 shows, best-first tree search includes depth-first search as a special case.) Most best-first algorithms include as a component of f a **heuristic function**, denoted  $h(n)$ :

 $h(n)$  = estimated cost of the cheapest path from the state at node *n* to a goal state.

(Notice that  $h(n)$  takes a *node* as input, but, unlike  $g(n)$ , it depends only on the *state* at that node.) For example, in Romania, one might estimate the cost of the cheapest path from Arad to Bucharest via the straight-line distance from Arad to Bucharest.

Heuristic functions are the most common form in which additional knowledge of the problem is imparted to the search algorithm. We study heuristics in more depth in Section 3.6. For now, we consider them to be arbitrary, nonnegative, problem-specific functions, with one constraint: if n is a goal node, then  $h(n) = 0$ . The remainder of this section covers two ways to use heuristic information to guide search.

#### **3.5.1 Greedy best-first search**

GREEDY BEST-FIRST **Greedy best-first search<sup>8</sup>** tries to expand the node that is closest to the goal, on the grounds that this is likely to lead to a solution quickly. Thus, it evaluates nodes by using just the heuristic function; that is,  $f(n) = h(n)$ .

Let us see how this works for route-finding problems in Romania; we use the **straight-**STRAIGHT-LINE **line distance** heuristic, which we will call  $h_{SLD}$ . If the goal is Bucharest, we need to know the straight-line distances to Bucharest, which are shown in Figure 3.22. For example,  $h_{SLD}(In(Arad)) = 366$ . Notice that the values of  $h_{SLD}$  cannot be computed from the problem description itself. Moreover, it takes a certain amount of experience to know that  $h_{SLD}$  is correlated with actual road distances and is, therefore, a useful heuristic.

> Figure 3.23 shows the progress of a greedy best-first search using  $h_{SLD}$  to find a path from Arad to Bucharest. The first node to be expanded from Arad will be Sibiu because it is closer to Bucharest than either Zerind or Timisoara. The next node to be expanded will be Fagaras because it is closest. Fagaras in turn generates Bucharest, which is the goal. For this particular problem, greedy best-first search using  $h_{SLD}$  finds a solution without ever

SEARCH

**DISTANCE** 

<sup>8</sup> Our first edition called this **greedy search**; other authors have called it **best-first search**. Our more general usage of the latter term follows Pearl (1984).



expanding a node that is not on the solution path; hence, its search cost is minimal. It is not optimal, however: the path via Sibiu and Fagaras to Bucharest is 32 kilometers longer than the path through Rimnicu Vilcea and Pitesti. This shows why the algorithm is called "greedy"—at each step it tries to get as close to the goal as it can.

Greedy best-first tree search is also incomplete even in a finite state space, much like depth-first search. Consider the problem of getting from Iasi to Fagaras. The heuristic suggests that Neamt be expanded first because it is closest to Fagaras, but it is a dead end. The solution is to go first to Vaslui—a step that is actually farther from the goal according to the heuristic—and then to continue to Urziceni, Bucharest, and Fagaras. The algorithm will never find this solution, however, because expanding Neamt puts Iasi back into the frontier, Iasi is closer to Fagaras than Vaslui is, and so Iasi will be expanded again, leading to an infinite loop. (The graph search version *is* complete in finite spaces, but not in infinite ones.) The worst-case time and space complexity for the tree version is  $O(b^m)$ , where m is the maximum depth of the search space. With a good heuristic function, however, the complexity can be reduced substantially. The amount of the reduction depends on the particular problem and on the quality of the heuristic.

# **3.5.2 A\* search: Minimizing the total estimated solution cost**

 $A^*$  SFARCH

The most widely known form of best-first search is called **A** ∗ <sup>A</sup> **search** (pronounced "A-star <sup>∗</sup> search"). It evaluates nodes by combining  $g(n)$ , the cost to reach the node, and  $h(n)$ , the cost to get from the node to the goal:

 $f(n) = q(n) + h(n)$ .

Since  $g(n)$  gives the path cost from the start node to node n, and  $h(n)$  is the estimated cost of the cheapest path from  $n$  to the goal, we have

 $f(n) =$  estimated cost of the cheapest solution through n.

Thus, if we are trying to find the cheapest solution, a reasonable thing to try first is the node with the lowest value of  $q(n) + h(n)$ . It turns out that this strategy is more than just reasonable: provided that the heuristic function  $h(n)$  satisfies certain conditions, A<sup>\*</sup> search is both complete and optimal. The algorithm is identical to UNIFORM-COST-SEARCH except that A<sup>\*</sup> uses  $g + h$  instead of g.



#### **Conditions for optimality: Admissibility and consistency**

**HEURISTIC** 

ADMISSIBLE<br>HEIRISTIC The first condition we require for optimality is that  $h(n)$  be an **admissible heuristic**. An admissible heuristic is one that *never overestimates* the cost to reach the goal. Because  $g(n)$ is the actual cost to reach n along the current path, and  $f(n) = g(n) + h(n)$ , we have as an immediate consequence that  $f(n)$  never overestimates the true cost of a solution along the current path through  $n$ .

> Admissible heuristics are by nature optimistic because they think the cost of solving the problem is less than it actually is. An obvious example of an admissible heuristic is the straight-line distance  $h_{SLD}$  that we used in getting to Bucharest. Straight-line distance is admissible because the shortest path between any two points is a straight line, so the straight

line cannot be an overestimate. In Figure 3.24, we show the progress of an A<sup>∗</sup> tree search for Bucharest. The values of  $q$  are computed from the step costs in Figure 3.2, and the values of  $h<sub>SLD</sub>$  are given in Figure 3.22. Notice in particular that Bucharest first appears on the frontier at step (e), but it is not selected for expansion because its f-cost (450) is higher than that of Pitesti (417). Another way to say this is that there *might* be a solution through Pitesti whose cost is as low as 417, so the algorithm will not settle for a solution that costs 450.

CONSISTENCY A second, slightly stronger condition called **consistency** (or sometimes **monotonicity**) MONOTONICITY is required only for applications of A<sup>\*</sup> to graph search.<sup>9</sup> A heuristic  $h(n)$  is consistent if, for every node *n* and every successor *n'* of *n* generated by any action *a*, the estimated cost of reaching the goal from  $n$  is no greater than the step cost of getting to  $n'$  plus the estimated cost of reaching the goal from  $n'$ :

 $h(n) \le c(n, a, n') + h(n')$ .

INEQUALITY



It is fairly easy to show (Exercise 3.32) that every consistent heuristic is also admissible. Consistency is therefore a stricter requirement than admissibility, but one has to work quite hard to concoct heuristics that are admissible but not consistent. All the admissible heuristics we discuss in this chapter are also consistent. Consider, for example,  $h_{SLD}$ . We know that the general triangle inequality is satisfied when each side is measured by the straight-line distance and that the straight-line distance between n and n' is no greater than  $c(n, a, n')$ . Hence,  $h_{SLD}$  is a consistent heuristic.

#### **Optimality of A\***

As we mentioned earlier, A<sup>∗</sup> has the following properties: *the tree-search version of* A<sup>∗</sup> *is optimal if*  $h(n)$  *is admissible, while the graph-search version is optimal if*  $h(n)$  *is consistent.* 

We show the second of these two claims since it is more useful. The argument essentially mirrors the argument for the optimality of uniform-cost search, with  $g$  replaced by  $f$ —just as in the A<sup>\*</sup> algorithm itself.

The first step is to establish the following: *if*  $h(n)$  *is consistent, then the values of* f(n) *along any path are nondecreasing.* The proof follows directly from the definition of consistency. Suppose n' is a successor of n; then  $g(n') = g(n) + c(n, a, n')$  for some action a, and we have

$$
f(n') = g(n') + h(n') = g(n) + c(n, a, n') + h(n') \ge g(n) + h(n) = f(n) .
$$

The next step is to prove that *whenever* A<sup>∗</sup> *selects a node* n *for expansion, the optimal path to that node has been found.* Were this not the case, there would have to be another frontier node  $n'$  on the optimal path from the start node to n, by the graph separation property of





<sup>9</sup> With an admissible but inconsistent heuristic, A<sup>\*</sup> requires some extra bookkeeping to ensure optimality.







Figure 3.9; because f is nondecreasing along any path,  $n'$  would have lower f-cost than n and would have been selected first.

From the two preceding observations, it follows that the sequence of nodes expanded by A<sup>\*</sup> using GRAPH-SEARCH is in nondecreasing order of  $f(n)$ . Hence, the first goal node selected for expansion must be an optimal solution because  $f$  is the true cost for goal nodes (which have  $h = 0$ ) and all later goal nodes will be at least as expensive.

The fact that  $f$ -costs are nondecreasing along any path also means that we can draw CONTOUR **contours** in the state space, just like the contours in a topographic map. Figure 3.25 shows an example. Inside the contour labeled 400, all nodes have  $f(n)$  less than or equal to 400, and so on. Then, because  $A^*$  expands the frontier node of lowest  $f$ -cost, we can see that an A<sup>\*</sup> search fans out from the start node, adding nodes in concentric bands of increasing f-cost.

With uniform-cost search (A<sup>\*</sup> search using  $h(n) = 0$ ), the bands will be "circular" around the start state. With more accurate heuristics, the bands will stretch toward the goal state and become more narrowly focused around the optimal path. If  $C^*$  is the cost of the optimal solution path, then we can say the following:

- A<sup>\*</sup> expands all nodes with  $f(n) < C^*$ .
- A<sup>\*</sup> might then expand some of the nodes right on the "goal contour" (where  $f(n) = C^*$ ) before selecting a goal node.

Completeness requires that there be only finitely many nodes with cost less than or equal to  $C^*$ , a condition that is true if all step costs exceed some finite  $\epsilon$  and if b is finite.

Notice that A<sup>\*</sup> expands no nodes with  $f(n) > C^*$ —for example, Timisoara is not expanded in Figure 3.24 even though it is a child of the root. We say that the subtree below

PRUNING Timisoara is **pruned**; because  $h_{SLD}$  is admissible, the algorithm can safely ignore this subtree while still guaranteeing optimality. The concept of pruning—eliminating possibilities from consideration without having to examine them—is important for many areas of AI.

One final observation is that among optimal algorithms of this type—algorithms that extend search paths from the root and use the same heuristic information—A<sup>∗</sup> is **optimally EFFICIENT efficient** for any given consistent heuristic. That is, no other optimal algorithm is guaranteed to expand fewer nodes than A<sup>∗</sup> (except possibly through tie-breaking among nodes with  $f(n) = C^*$ ). This is because any algorithm that *does not* expand all nodes with  $f(n) < C^*$ runs the risk of missing the optimal solution.

That A<sup>∗</sup> search is complete, optimal, and optimally efficient among all such algorithms is rather satisfying. Unfortunately, it does not mean that A<sup>∗</sup> is the answer to all our searching needs. The catch is that, for most problems, the number of states within the goal contour search space is still exponential in the length of the solution. The details of the analysis are beyond the scope of this book, but the basic results are as follows. For problems with constant step costs, the growth in run time as a function of the optimal solution depth  $d$  is analyzed in ABSOLUTE ERROR terms of the the **absolute error** or the **relative error** of the heuristic. The absolute error is RELATIVE ERROR defined as  $\Delta \equiv h^* - h$ , where  $h^*$  is the actual cost of getting from the root to the goal, and the relative error is defined as  $\epsilon \equiv (h^* - h)/h^*$ .

> The complexity results depend very strongly on the assumptions made about the state space. The simplest model studied is a state space that has a single goal and is essentially a tree with reversible actions. (The 8-puzzle satisfies the first and third of these assumptions.) In this case, the time complexity of  $A^*$  is exponential in the maximum absolute error, that is,  $O(b^{\Delta})$ . For constant step costs, we can write this as  $O(b^{\epsilon d})$ , where d is the solution depth. For almost all heuristics in practical use, the absolute error is at least proportional to the path cost  $h^*$ , so  $\epsilon$  is constant or growing and the time complexity is exponential in d. We can also see the effect of a more accurate heuristic:  $O(b^{\epsilon d}) = O((b^{\epsilon})^d)$ , so the effective branching factor (defined more formally in the next section) is  $b^{\epsilon}$ .

> When the state space has many goal states—particularly *near-optimal* goal states—the search process can be led astray from the optimal path and there is an extra cost proportional to the number of goals whose cost is within a factor  $\epsilon$  of the optimal cost. Finally, in the general case of a graph, the situation is even worse. There can be exponentially many states with  $f(n) < C^*$  even if the absolute error is bounded by a constant. For example, consider a version of the vacuum world where the agent can clean up any square for unit cost without even having to visit it: in that case, squares can be cleaned in any order. With  $N$  initially dirty squares, there are  $2^N$  states where some subset has been cleaned and all of them are on an optimal solution path—and hence satisfy  $f(n) < C^*$ —even if the heuristic has an error of 1.

> The complexity of A<sup>★</sup> often makes it impractical to insist on finding an optimal solution. One can use variants of A<sup>∗</sup> that find suboptimal solutions quickly, or one can sometimes design heuristics that are more accurate but not strictly admissible. In any case, the use of a good heuristic still provides enormous savings compared to the use of an uninformed search. In Section 3.6, we look at the question of designing good heuristics.

> Computation time is not, however, A<sup>∗</sup> 's main drawback. Because it keeps all generated nodes in memory (as do all GRAPH-SEARCH algorithms), A<sup>∗</sup> usually runs out of space long

EFFICIENT

**function** RECURSIVE-BEST-FIRST-SEARCH(problem) **returns** a solution, or failure **return** RBFS(problem, MAKE-NODE(problem.INITIAL-STATE),∞) **function** RBFS(*problem, node, f<sub>-</sub>limit*) **returns** a solution, or failure and a new f-cost limit **if** problem.GOAL-TEST(node.STATE) **then return** SOLUTION(node)  $successors \leftarrow [ ]$ **for each** action **in** problem.ACTIONS(node.STATE) **do** add CHILD-NODE(problem, node, action) into successors **if** successors is empty **then return** failure,  $\infty$ **for each** s **in** successors **do** /\* update f with value from previous search, if any \*/  $s.f \leftarrow \max(s.g + s.h, node.f)$ **loop do**  $best \leftarrow$  the lowest f-value node in successors **if** best.f  $> f$  limit **then return** failure, best.f  $\mathit{alternative} \leftarrow$  the second-lowest f-value among successors  $result, best.f \leftarrow RBFS(problem, best, min(f\_limit, alternative))$ **if** result  $\neq$  failure **then return** result Figure 3.26 The algorithm for recursive best-first search.

before it runs out of time. For this reason, A<sup>\*</sup> is not practical for many large-scale problems. There are, however, algorithms that overcome the space problem without sacrificing optimality or completeness, at a small cost in execution time. We discuss these next.

# **3.5.3 Memory-bounded heuristic search**

The simplest way to reduce memory requirements for  $A^*$  is to adapt the idea of iterative deepening to the heuristic search context, resulting in the **iterative-deepening** A<sup>∗</sup> (IDA<sup>∗</sup>) algorithm. The main difference between IDA<sup>\*</sup> and standard iterative deepening is that the cutoff used is the f-cost  $(g+h)$  rather than the depth; at each iteration, the cutoff value is the smallest f-cost of any node that exceeded the cutoff on the previous iteration. IDA<sup>\*</sup> is practical for many problems with unit step costs and avoids the substantial overhead associated with keeping a sorted queue of nodes. Unfortunately, it suffers from the same difficulties with realvalued costs as does the iterative version of uniform-cost search described in Exercise 3.18. This section briefly examines two other memory-bounded algorithms, called RBFS and MA<sup>∗</sup>.

BEST-FIRST SEARCH

RECURSIVE<br>RECURSIVE SEARCH **RECURSIVE BEST-first search** (RBFS) is a simple recursive algorithm that attempts to mimic the operation of standard best-first search, but using only linear space. The algorithm is shown in Figure 3.26. Its structure is similar to that of a recursive depth-first search, but rather than continuing indefinitely down the current path, it uses the  $f$ -limit variable to keep track of the f-value of the best *alternative* path available from any ancestor of the current node. If the current node exceeds this limit, the recursion unwinds back to the alternative path. As the recursion unwinds, RBFS replaces the  $f$ -value of each node along the path BACKED-UP VALUE with a **backed-up value**—the best  $f$ -value of its children. In this way, RBFS remembers the f-value of the best leaf in the forgotten subtree and can therefore decide whether it's worth

ITERATIVE-DEEPENING<br>∗  $\mathbf{A}^*$ 



**Figure 3.27** Stages in an RBFS search for the shortest route to Bucharest. The f-limit value for each recursive call is shown on top of each current node, and every node is labeled with its f-cost. (a) The path via Rimnicu Vilcea is followed until the current best leaf (Pitesti) has a value that is worse than the best alternative path (Fagaras). (b) The recursion unwinds and the best leaf value of the forgotten subtree (417) is backed up to Rimnicu Vilcea; then Fagaras is expanded, revealing a best leaf value of 450. (c) The recursion unwinds and the best leaf value of the forgotten subtree (450) is backed up to Fagaras; then Rimnicu Vilcea is expanded. This time, because the best alternative path (through Timisoara) costs at least 447, the expansion continues to Bucharest.

reexpanding the subtree at some later time. Figure 3.27 shows how RBFS reaches Bucharest.

RBFS is somewhat more efficient than IDA<sup>\*</sup>, but still suffers from excessive node regeneration. In the example in Figure 3.27, RBFS follows the path via Rimnicu Vilcea, then "changes its mind" and tries Fagaras, and then changes its mind back again. These mind changes occur because every time the current best path is extended, its f-value is likely to increase— $h$  is usually less optimistic for nodes closer to the goal. When this happens, the second-best path might become the best path, so the search has to backtrack to follow it. Each mind change corresponds to an iteration of IDA<sup>∗</sup> and could require many reexpansions of forgotten nodes to recreate the best path and extend it one more node.

Like A<sup>\*</sup> tree search, RBFS is an optimal algorithm if the heuristic function  $h(n)$  is admissible. Its space complexity is linear in the depth of the deepest optimal solution, but its time complexity is rather difficult to characterize: it depends both on the accuracy of the heuristic function and on how often the best path changes as nodes are expanded.

IDA<sup>∗</sup> and RBFS suffer from using *too little* memory. Between iterations, IDA<sup>∗</sup> retains only a single number: the current f-cost limit. RBFS retains more information in memory, but it uses only linear space: even if more memory were available, RBFS has no way to make use of it. Because they forget most of what they have done, both algorithms may end up reexpanding the same states many times over. Furthermore, they suffer the potentially exponential increase in complexity associated with redundant paths in graphs (see Section 3.3).

It seems sensible, therefore, to use all available memory. Two algorithms that do this MA<sup>∗</sup> are MA<sup>\*</sup> (memory-bounded A<sup>\*</sup>) and **SMA<sup>\*</sup>** (simplified MA<sup>\*</sup>). SMA<sup>\*</sup> is—well—simpler, so SMA\* we will describe it. SMA<sup>∗</sup> proceeds just like A<sup>∗</sup> , expanding the best leaf until memory is full. At this point, it cannot add a new node to the search tree without dropping an old one. SMA<sup>\*</sup> always drops the *worst* leaf node—the one with the highest f-value. Like RBFS, SMA<sup>∗</sup> then backs up the value of the forgotten node to its parent. In this way, the ancestor of a forgotten subtree knows the quality of the best path in that subtree. With this information, SMA<sup>\*</sup> regenerates the subtree only when all other paths have been shown to look worse than the path it has forgotten. Another way of saying this is that, if all the descendants of a node  $n$ are forgotten, then we will not know which way to go from  $n$ , but we will still have an idea of how worthwhile it is to go anywhere from  $n$ .

> The complete algorithm is too complicated to reproduce here,  $10$  but there is one subtlety worth mentioning. We said that SMA<sup>\*</sup> expands the best leaf and deletes the worst leaf. What if *all* the leaf nodes have the same f-value? To avoid selecting the same node for deletion and expansion, SMA<sup>∗</sup> expands the *newest* best leaf and deletes the *oldest* worst leaf. These coincide when there is only one leaf, but in that case, the current search tree must be a single path from root to leaf that fills all of memory. If the leaf is not a goal node, then *even if it is on an optimal solution path*, that solution is not reachable with the available memory. Therefore, the node can be discarded exactly as if it had no successors.

> SMA<sup>\*</sup> is complete if there is any reachable solution—that is, if  $d$ , the depth of the shallowest goal node, is less than the memory size (expressed in nodes). It is optimal if any optimal solution is reachable; otherwise, it returns the best reachable solution. In practical terms, SMA<sup>∗</sup> is a fairly robust choice for finding optimal solutions, particularly when the state space is a graph, step costs are not uniform, and node generation is expensive compared to the overhead of maintaining the frontier and the explored set.

<sup>10</sup> A rough sketch appeared in the first edition of this book.

On very hard problems, however, it will often be the case that SMA<sup>∗</sup> is forced to switch back and forth continually among many candidate solution paths, only a small subset of which THRASHING can fit in memory. (This resembles the problem of **thrashing** in disk paging systems.) Then the extra time required for repeated regeneration of the same nodes means that problems that would be practically solvable by A<sup>∗</sup> , given unlimited memory, become intractable for SMA<sup>∗</sup> . That is to say, *memory limitations can make a problem intractable from the point of view of computation time.* Although no current theory explains the tradeoff between time and memory, it seems that this is an inescapable problem. The only way out is to drop the optimality requirement.

#### **3.5.4 Learning to search better**

We have presented several fixed strategies—breadth-first, greedy best-first, and so on—that have been designed by computer scientists. Could an agent *learn* how to search better? The answer is yes, and the method rests on an important concept called the **metalevel state space**. Each state in a metalevel state space captures the internal (computational) state of a program OBJECT-LEVEL STATE that is searching in an **object-level state space** such as Romania. For example, the internal state of the A<sup>∗</sup> algorithm consists of the current search tree. Each action in the metalevel state space is a computation step that alters the internal state; for example, each computation step in A<sup>∗</sup> expands a leaf node and adds its successors to the tree. Thus, Figure 3.24, which shows a sequence of larger and larger search trees, can be seen as depicting a path in the metalevel state space where each state on the path is an object-level search tree.

Now, the path in Figure 3.24 has five steps, including one step, the expansion of Fagaras, that is not especially helpful. For harder problems, there will be many such missteps, and a METALEVEL **metalevel learning** algorithm can learn from these experiences to avoid exploring unpromising subtrees. The techniques used for this kind of learning are described in Chapter 21. The goal of learning is to minimize the **total cost** of problem solving, trading off computational expense and path cost.

# 3.6 HEURISTIC FUNCTIONS

In this section, we look at heuristics for the 8-puzzle, in order to shed light on the nature of heuristics in general.

The 8-puzzle was one of the earliest heuristic search problems. As mentioned in Section 3.2, the object of the puzzle is to slide the tiles horizontally or vertically into the empty space until the configuration matches the goal configuration (Figure 3.28).

The average solution cost for a randomly generated 8-puzzle instance is about 22 steps. The branching factor is about 3. (When the empty tile is in the middle, four moves are possible; when it is in a corner, two; and when it is along an edge, three.) This means that an exhaustive tree search to depth 22 would look at about  $3^{22} \approx 3.1 \times 10^{10}$  states. A graph search would cut this down by a factor of about 170,000 because only  $9!/2$  = 181, 440 distinct states are reachable. (See Exercise 3.5.) This is a manageable number, but

METALEVEL STATE SPACE

**SPACE** 

LEARNING



the corresponding number for the 15-puzzle is roughly  $10^{13}$ , so the next order of business is to find a good heuristic function. If we want to find the shortest solutions by using  $A^*$ , we need a heuristic function that never overestimates the number of steps to the goal. There is a long history of such heuristics for the 15-puzzle; here are two commonly used candidates:

- $h_1$  = the number of misplaced tiles. For Figure 3.28, all of the eight tiles are out of position, so the start state would have  $h_1 = 8$ .  $h_1$  is an admissible heuristic because it is clear that any tile that is out of place must be moved at least once.
- $h_2$  = the sum of the distances of the tiles from their goal positions. Because tiles cannot move along diagonals, the distance we will count is the sum of the horizontal and vertical distances. This is sometimes called the **city block distance** or **Manhattan** MANHATTAN<br>DISTANCE **distance**.  $h_2$  is also admissible because all any move can do is move one tile one step closer to the goal. Tiles 1 to 8 in the start state give a Manhattan distance of

 $h_2 = 3 + 1 + 2 + 2 + 2 + 3 + 3 + 2 = 18$ .

As expected, neither of these overestimates the true solution cost, which is 26.

#### **3.6.1 The effect of heuristic accuracy on performance**

EFFECTIVE<br>BRANCHING FACTOR

EFFECTIVE **One** way to characterize the quality of a heuristic is the **effective branching factor**  $b^*$ . If the total number of nodes generated by  $A^*$  for a particular problem is N and the solution depth is d, then  $b^*$  is the branching factor that a uniform tree of depth d would have to have in order to contain  $N + 1$  nodes. Thus,

$$
N + 1 = 1 + b^* + (b^*)^2 + \dots + (b^*)^d.
$$

For example, if A<sup>∗</sup> finds a solution at depth 5 using 52 nodes, then the effective branching factor is 1.92. The effective branching factor can vary across problem instances, but usually it is fairly constant for sufficiently hard problems. (The existence of an effective branching factor follows from the result, mentioned earlier, that the number of nodes expanded by  $A^*$ grows exponentially with solution depth.) Therefore, experimental measurements of  $b^*$  on a small set of problems can provide a good guide to the heuristic's overall usefulness. A welldesigned heuristic would have a value of  $b^*$  close to 1, allowing fairly large problems to be solved at reasonable computational cost.

DISTANCE

To test the heuristic functions  $h_1$  and  $h_2$ , we generated 1200 random problems with solution lengths from 2 to 24 (100 for each even number) and solved them with iterative deepening search and with A<sup>\*</sup> tree search using both  $h_1$  and  $h_2$ . Figure 3.29 gives the average number of nodes generated by each strategy and the effective branching factor. The results suggest that  $h_2$  is better than  $h_1$ , and is far better than using iterative deepening search. Even for small problems with  $d = 12$ , A<sup>\*</sup> with  $h<sub>2</sub>$  is 50,000 times more efficient than uninformed iterative deepening search.



**Figure 3.29** Comparison of the search costs and effective branching factors for the ITERATIVE-DEEPENING-SEARCH and A<sup>\*</sup> algorithms with  $h_1$ ,  $h_2$ . Data are averaged over 100 instances of the 8-puzzle for each of various solution lengths d.

One might ask whether  $h_2$  is *always* better than  $h_1$ . The answer is "Essentially, yes." It is easy to see from the definitions of the two heuristics that, for any node  $n, h_2(n) \ge h_1(n)$ . DOMINATION We thus say that  $h_2$  **dominates**  $h_1$ . Domination translates directly into efficiency: A<sup>\*</sup> using  $h_2$  will never expand more nodes than A<sup>\*</sup> using  $h_1$  (except possibly for some nodes with  $f(n) = C^*$ ). The argument is simple. Recall the observation on page 97 that every node with  $f(n) < C^*$  will surely be expanded. This is the same as saying that every node with  $h(n) < C^* - g(n)$  will surely be expanded. But because  $h_2$  is at least as big as  $h_1$  for all nodes, every node that is surely expanded by A<sup>\*</sup> search with  $h_2$  will also surely be expanded with  $h_1$ , and  $h_1$  might cause other nodes to be expanded as well. Hence, it is generally better to use a heuristic function with higher values, provided it is consistent and that the computation time for the heuristic is not too long.

# **3.6.2 Generating admissible heuristics from relaxed problems**

We have seen that both  $h_1$  (misplaced tiles) and  $h_2$  (Manhattan distance) are fairly good heuristics for the 8-puzzle and that  $h_2$  is better. How might one have come up with  $h_2$ ? Is it possible for a computer to invent such a heuristic mechanically?

 $h_1$  and  $h_2$  are estimates of the remaining path length for the 8-puzzle, but they are also perfectly accurate path lengths for *simplified* versions of the puzzle. If the rules of the puzzle

were changed so that a tile could move anywhere instead of just to the adjacent empty square, then  $h_1$  would give the exact number of steps in the shortest solution. Similarly, if a tile could move one square in any direction, even onto an occupied square, then  $h_2$  would give the exact number of steps in the shortest solution. A problem with fewer restrictions on the actions is RELAXED PROBLEM called a **relaxed problem**. The state-space graph of the relaxed problem is a *supergraph* of the original state space because the removal of restrictions creates added edges in the graph.



Because the relaxed problem adds edges to the state space, any optimal solution in the original problem is, by definition, also a solution in the relaxed problem; but the relaxed problem may have *better* solutions if the added edges provide short cuts. Hence, *the cost of an optimal solution to a relaxed problem is an admissible heuristic for the original problem.* Furthermore, because the derived heuristic is an exact cost for the relaxed problem, it must obey the triangle inequality and is therefore **consistent** (see page 95).

If a problem definition is written down in a formal language, it is possible to construct relaxed problems automatically.<sup>11</sup> For example, if the 8-puzzle actions are described as

A tile can move from square A to square B if

A is horizontally or vertically adjacent to B **and** B is blank,

we can generate three relaxed problems by removing one or both of the conditions:

- (a) A tile can move from square A to square B if A is adjacent to B.
- (b) A tile can move from square A to square B if B is blank.
- (c) A tile can move from square A to square B.

From (a), we can derive  $h_2$  (Manhattan distance). The reasoning is that  $h_2$  would be the proper score if we moved each tile in turn to its destination. The heuristic derived from (b) is discussed in Exercise 3.34. From (c), we can derive  $h_1$  (misplaced tiles) because it would be the proper score if tiles could move to their intended destination in one step. Notice that it is crucial that the relaxed problems generated by this technique can be solved essentially *without search*, because the relaxed rules allow the problem to be decomposed into eight independent subproblems. If the relaxed problem is hard to solve, then the values of the corresponding heuristic will be expensive to obtain.<sup>12</sup>

A program called ABSOLVER can generate heuristics automatically from problem definitions, using the "relaxed problem" method and various other techniques (Prieditis, 1993). ABSOLVER generated a new heuristic for the 8-puzzle that was better than any preexisting heuristic and found the first useful heuristic for the famous Rubik's Cube puzzle.

One problem with generating new heuristic functions is that one often fails to get a single "clearly best" heuristic. If a collection of admissible heuristics  $h_1 \ldots h_m$  is available for a problem and none of them dominates any of the others, which should we choose? As it turns out, we need not make a choice. We can have the best of all worlds, by defining

 $h(n) = \max\{h_1(n), \ldots, h_m(n)\}\.$ 

 $11$  In Chapters 8 and 10, we describe formal languages suitable for this task; with formal descriptions that can be manipulated, the construction of relaxed problems can be automated. For now, we use English.

<sup>&</sup>lt;sup>12</sup> Note that a perfect heuristic can be obtained simply by allowing h to run a full breadth-first search "on the sly." Thus, there is a tradeoff between accuracy and computation time for heuristic functions.



get tiles 1, 2, 3, and 4 into their correct positions, without worrying about what happens to the other tiles.

This composite heuristic uses whichever function is most accurate on the node in question. Because the component heuristics are admissible,  $h$  is admissible; it is also easy to prove that  $h$  is consistent. Furthermore,  $h$  dominates all of its component heuristics.

#### **3.6.3 Generating admissible heuristics from subproblems: Pattern databases**

SUBPROBLEM Admissible heuristics can also be derived from the solution cost of a **subproblem** of a given problem. For example, Figure 3.30 shows a subproblem of the 8-puzzle instance in Figure 3.28. The subproblem involves getting tiles 1, 2, 3, 4 into their correct positions. Clearly, the cost of the optimal solution of this subproblem is a lower bound on the cost of the complete problem. It turns out to be more accurate than Manhattan distance in some cases.

PATTERN DATABASE The idea behind **pattern databases** is to store these exact solution costs for every possible subproblem instance—in our example, every possible configuration of the four tiles and the blank. (The locations of the other four tiles are irrelevant for the purposes of solving the subproblem, but moves of those tiles do count toward the cost.) Then we compute an admissible heuristic  $h_{DB}$  for each complete state encountered during a search simply by looking up the corresponding subproblem configuration in the database. The database itself is constructed by searching back<sup>13</sup> from the goal and recording the cost of each new pattern encountered; the expense of this search is amortized over many subsequent problem instances.

> The choice of 1-2-3-4 is fairly arbitrary; we could also construct databases for 5-6-7-8, for 2-4-6-8, and so on. Each database yields an admissible heuristic, and these heuristics can be combined, as explained earlier, by taking the maximum value. A combined heuristic of this kind is much more accurate than the Manhattan distance; the number of nodes generated when solving random 15-puzzles can be reduced by a factor of 1000.

> One might wonder whether the heuristics obtained from the 1-2-3-4 database and the 5-6-7-8 could be *added*, since the two subproblems seem not to overlap. Would this still give an admissible heuristic? The answer is no, because the solutions of the 1-2-3-4 subproblem and the 5-6-7-8 subproblem for a given state will almost certainly share some moves—it is

<sup>&</sup>lt;sup>13</sup> By working backward from the goal, the exact solution cost of every instance encountered is immediately available. This is an example of **dynamic programming**, which we discuss further in Chapter 17.

unlikely that 1-2-3-4 can be moved into place without touching 5-6-7-8, and vice versa. But what if we don't count those moves? That is, we record not the total cost of solving the 1-2- 3-4 subproblem, but just the number of moves involving 1-2-3-4. Then it is easy to see that the sum of the two costs is still a lower bound on the cost of solving the entire problem. This DISJOINT PATTERN is the idea behind **disjoint pattern databases**. With such databases, it is possible to solve random 15-puzzles in a few milliseconds—the number of nodes generated is reduced by a factor of 10,000 compared with the use of Manhattan distance. For 24-puzzles, a speedup of roughly a factor of a million can be obtained.

> Disjoint pattern databases work for sliding-tile puzzles because the problem can be divided up in such a way that each move affects only one subproblem—because only one tile is moved at a time. For a problem such as Rubik's Cube, this kind of subdivision is difficult because each move affects 8 or 9 of the 26 cubies. More general ways of defining additive, admissible heuristics have been proposed that do apply to Rubik's cube (Yang *et al.*, 2008), but they have not yielded a heuristic better than the best nonadditive heuristic for the problem.

#### **3.6.4 Learning heuristics from experience**

A heuristic function  $h(n)$  is supposed to estimate the cost of a solution beginning from the state at node  $n$ . How could an agent construct such a function? One solution was given in the preceding sections—namely, to devise relaxed problems for which an optimal solution can be found easily. Another solution is to learn from experience. "Experience" here means solving lots of 8-puzzles, for instance. Each optimal solution to an 8-puzzle problem provides examples from which  $h(n)$  can be learned. Each example consists of a state from the solution path and the actual cost of the solution from that point. From these examples, a learning algorithm can be used to construct a function  $h(n)$  that can (with luck) predict solution costs for other states that arise during search. Techniques for doing just this using neural nets, decision trees, and other methods are demonstrated in Chapter 18. (The reinforcement learning methods described in Chapter 21 are also applicable.)

DATABASES

FEATURE Inductive learning methods work best when supplied with **features** of a state that are relevant to predicting the state's value, rather than with just the raw state description. For example, the feature "number of misplaced tiles" might be helpful in predicting the actual distance of a state from the goal. Let's call this feature  $x_1(n)$ . We could take 100 randomly generated 8-puzzle configurations and gather statistics on their actual solution costs. We might find that when  $x_1(n)$  is 5, the average solution cost is around 14, and so on. Given these data, the value of  $x_1$  can be used to predict  $h(n)$ . Of course, we can use several features. A second feature  $x_2(n)$  might be "number of pairs of adjacent tiles that are not adjacent in the goal state." How should  $x_1(n)$  and  $x_2(n)$  be combined to predict  $h(n)$ ? A common approach is to use a linear combination:

 $h(n) = c_1x_1(n) + c_2x_2(n)$ .

The constants  $c_1$  and  $c_2$  are adjusted to give the best fit to the actual data on solution costs. One expects both  $c_1$  and  $c_2$  to be positive because misplaced tiles and incorrect adjacent pairs make the problem harder to solve. Notice that this heuristic does satisfy the condition that  $h(n) = 0$  for goal states, but it is not necessarily admissible or consistent.

# 3.7 SUMMARY

This chapter has introduced methods that an agent can use to select actions in environments that are deterministic, observable, static, and completely known. In such cases, the agent can construct sequences of actions that achieve its goals; this process is called **search**.

- Before an agent can start searching for solutions, a **goal** must be identified and a welldefined **problem** must be formulated.
- A problem consists of five parts: the **initial state**, a set of **actions**, a **transition model** describing the results of those actions, a **goal test** function, and a **path cost** function. The environment of the problem is represented by a **state space**. A **path** through the state space from the initial state to a goal state is a **solution**.
- Search algorithms treat states and actions as **atomic**: they do not consider any internal structure they might possess.
- A general TREE-SEARCH algorithm considers all possible paths to find a solution, whereas a GRAPH-SEARCH algorithm avoids consideration of redundant paths.
- Search algorithms are judged on the basis of **completeness**, **optimality**, **time complexity**, and **space complexity**. Complexity depends on b, the branching factor in the state space, and d, the depth of the shallowest solution.
- **Uninformed search** methods have access only to the problem definition. The basic algorithms are as follows:
	- **Breadth-first search** expands the shallowest nodes first; it is complete, optimal for unit step costs, but has exponential space complexity.
	- **Uniform-cost search** expands the node with lowest path cost,  $q(n)$ , and is optimal for general step costs.
	- **Depth-first search** expands the deepest unexpanded node first. It is neither complete nor optimal, but has linear space complexity. **Depth-limited search** adds a depth bound.
	- **Iterative deepening search** calls depth-first search with increasing depth limits until a goal is found. It is complete, optimal for unit step costs, has time complexity comparable to breadth-first search, and has linear space complexity.
	- **Bidirectional search** can enormously reduce time complexity, but it is not always applicable and may require too much space.
- **Informed search** methods may have access to a **heuristic** function  $h(n)$  that estimates the cost of a solution from  $n$ .
	- **–** The generic **best-first search** algorithm selects a node for expansion according to an **evaluation function**.
	- $-$  **Greedy best-first search** expands nodes with minimal  $h(n)$ . It is not optimal but is often efficient.
- **A<sup>\*</sup> search** expands nodes with minimal  $f(n) = g(n) + h(n)$ . A<sup>\*</sup> is complete and optimal, provided that  $h(n)$  is admissible (for TREE-SEARCH) or consistent (for GRAPH-SEARCH). The space complexity of A<sup>∗</sup> is still prohibitive.
- **RBFS** (recursive best-first search) and **SMA**<sup>∗</sup> (simplified memory-bounded A<sup>∗</sup> ) are robust, optimal search algorithms that use limited amounts of memory; given enough time, they can solve problems that A<sup>∗</sup> cannot solve because it runs out of memory.
- The performance of heuristic search algorithms depends on the quality of the heuristic function. One can sometimes construct good heuristics by relaxing the problem definition, by storing precomputed solution costs for subproblems in a pattern database, or by learning from experience with the problem class.

# BIBLIOGRAPHICAL AND HISTORICAL NOTES

The topic of state-space search originated in more or less its current form in the early years of AI. Newell and Simon's work on the Logic Theorist (1957) and GPS (1961) led to the establishment of search algorithms as the primary weapons in the armory of 1960s AI researchers and to the establishment of problem solving as the canonical AI task. Work in operations research by Richard Bellman (1957) showed the importance of additive path costs in simplifying optimization algorithms. The text on *Automated Problem Solving* by Nils Nilsson (1971) established the area on a solid theoretical footing.

Most of the state-space search problems analyzed in this chapter have a long history in the literature and are less trivial than they might seem. The missionaries and cannibals problem used in Exercise 3.9 was analyzed in detail by Amarel (1968). It had been considered earlier—in AI by Simon and Newell (1961) and in operations research by Bellman and Dreyfus (1962).

The 8-puzzle is a smaller cousin of the 15-puzzle, whose history is recounted at length by Slocum and Sonneveld (2006). It was widely believed to have been invented by the famous American game designer Sam Loyd, based on his claims to that effect from 1891 onward (Loyd, 1959). Actually it was invented by Noyes Chapman, a postmaster in Canastota, New York, in the mid-1870s. (Chapman was unable to patent his invention, as a generic patent covering sliding blocks with letters, numbers, or pictures was granted to Ernest Kinsey in 1878.) It quickly attracted the attention of the public and of mathematicians (Johnson and Story, 1879; Tait, 1880). The editors of the *American Journal of Mathematics* stated, "The '15' puzzle for the last few weeks has been prominently before the American public, and may safely be said to have engaged the attention of nine out of ten persons of both sexes and all ages and conditions of the community." Ratner and Warmuth (1986) showed that the general  $n \times n$  version of the 15-puzzle belongs to the class of NP-complete problems.

The 8-queens problem was first published anonymously in the German chess magazine *Schach* in 1848; it was later attributed to one Max Bezzel. It was republished in 1850 and at that time drew the attention of the eminent mathematician Carl Friedrich Gauss, who

attempted to enumerate all possible solutions; initially he found only 72, but eventually he found the correct answer of 92, although Nauck published all 92 solutions first, in 1850. Netto (1901) generalized the problem to  $n$  queens, and Abramson and Yung (1989) found an  $O(n)$  algorithm.

Each of the real-world search problems listed in the chapter has been the subject of a good deal of research effort. Methods for selecting optimal airline flights remain proprietary for the most part, but Carl de Marcken (personal communication) has shown that airline ticket pricing and restrictions have become so convoluted that the problem of selecting an optimal flight is formally *undecidable*. The traveling-salesperson problem is a standard combinatorial problem in theoretical computer science (Lawler *et al.*, 1992). Karp (1972) proved the TSP to be NP-hard, but effective heuristic approximation methods were developed (Lin and Kernighan, 1973). Arora (1998) devised a fully polynomial approximation scheme for Euclidean TSPs. VLSI layout methods are surveyed by Shahookar and Mazumder (1991), and many layout optimization papers appear in VLSI journals. Robotic navigation and assembly problems are discussed in Chapter 25.

Uninformed search algorithms for problem solving are a central topic of classical computer science (Horowitz and Sahni, 1978) and operations research (Dreyfus, 1969). Breadthfirst search was formulated for solving mazes by Moore (1959). The method of **dynamic programming** (Bellman, 1957; Bellman and Dreyfus, 1962), which systematically records solutions for all subproblems of increasing lengths, can be seen as a form of breadth-first search on graphs. The two-point shortest-path algorithm of Dijkstra (1959) is the origin of uniform-cost search. These works also introduced the idea of explored and frontier sets (closed and open lists).

A version of iterative deepening designed to make efficient use of the chess clock was first used by Slate and Atkin (1977) in the CHESS 4.5 game-playing program. Martelli's algorithm B (1977) includes an iterative deepening aspect and also dominates A<sup>∗</sup> 's worst-case performance with admissible but inconsistent heuristics. The iterative deepening technique came to the fore in work by Korf (1985a). Bidirectional search, which was introduced by Pohl (1971), can also be effective in some cases.

The use of heuristic information in problem solving appears in an early paper by Simon and Newell (1958), but the phrase "heuristic search" and the use of heuristic functions that estimate the distance to the goal came somewhat later (Newell and Ernst, 1965; Lin, 1965). Doran and Michie (1966) conducted extensive experimental studies of heuristic search. Although they analyzed path length and "penetrance" (the ratio of path length to the total number of nodes examined so far), they appear to have ignored the information provided by the path cost  $g(n)$ . The A<sup>\*</sup> algorithm, incorporating the current path cost into heuristic search, was developed by Hart, Nilsson, and Raphael (1968), with some later corrections (Hart *et al.*, 1972). Dechter and Pearl (1985) demonstrated the optimal efficiency of A<sup>∗</sup> .

The original A<sup>∗</sup> paper introduced the consistency condition on heuristic functions. The monotone condition was introduced by Pohl (1977) as a simpler replacement, but Pearl (1984) showed that the two were equivalent.

Pohl (1977) pioneered the study of the relationship between the error in heuristic functions and the time complexity of A<sup>∗</sup> . Basic results were obtained for tree search with unit step

costs and a single goal node (Pohl, 1977; Gaschnig, 1979; Huyn *et al.*, 1980; Pearl, 1984) and with multiple goal nodes (Dinh *et al.*, 2007). The "effective branching factor" was proposed by Nilsson (1971) as an empirical measure of the efficiency; it is equivalent to assuming a time cost of  $O((b^*)^d)$ . For tree search applied to a graph, Korf *et al.* (2001) argue that the time cost is better modeled as  $O(b^{d-k})$ , where k depends on the heuristic accuracy; this analysis has elicited some controversy, however. For graph search, Helmert and Röger (2008) noted that several well-known problems contained exponentially many nodes on optimal solution paths, implying exponential time complexity for  $A^*$  even with constant absolute error in h.

There are many variations on the A<sup>∗</sup> algorithm. Pohl (1973) proposed the use of *dynamic weighting*, which uses a weighted sum  $f_w(n) = w_g g(n) + w_h h(n)$  of the current path length and the heuristic function as an evaluation function, rather than the simple sum  $f(n) = g(n) +$  $h(n)$  used in A<sup>\*</sup>. The weights  $w_g$  and  $w_h$  are adjusted dynamically as the search progresses. Pohl's algorithm can be shown to be  $\epsilon$ -admissible—that is, guaranteed to find solutions within a factor  $1 + \epsilon$  of the optimal solution, where  $\epsilon$  is a parameter supplied to the algorithm. The same property is exhibited by the  $A_{\epsilon}^{*}$  algorithm (Pearl, 1984), which can select any node from the frontier provided its f-cost is within a factor  $1 + \epsilon$  of the lowest-f-cost frontier node. The selection can be done so as to minimize search cost.

Bidirectional versions of A<sup>\*</sup> have been investigated; a combination of bidirectional A<sup>\*</sup> and known landmarks was used to efficiently find driving routes for Microsoft's online map service (Goldberg *et al.*, 2006). After caching a set of paths between landmarks, the algorithm can find an optimal path between any pair of points in a 24 million point graph of the United States, searching less than 0.1% of the graph. Others approaches to bidirectional search include a breadth-first search backward from the goal up to a fixed depth, followed by a forward IDA<sup>∗</sup> search (Dillenburg and Nelson, 1994; Manzini, 1995).

A ∗ and other state-space search algorithms are closely related to the *branch-and-bound* techniques that are widely used in operations research (Lawler and Wood, 1966). The relationships between state-space search and branch-and-bound have been investigated in depth (Kumar and Kanal, 1983; Nau *et al.*, 1984; Kumar *et al.*, 1988). Martelli and Montanari (1978) demonstrate a connection between dynamic programming (see Chapter 17) and certain types of state-space search. Kumar and Kanal (1988) attempt a "grand unification" of heuristic search, dynamic programming, and branch-and-bound techniques under the name of CDP—the "composite decision process."

Because computers in the late 1950s and early 1960s had at most a few thousand words of main memory, memory-bounded heuristic search was an early research topic. The Graph Traverser (Doran and Michie, 1966), one of the earliest search programs, commits to an operator after searching best-first up to the memory limit. IDA<sup>∗</sup> (Korf, 1985a, 1985b) was the first widely used optimal, memory-bounded heuristic search algorithm, and a large number of variants have been developed. An analysis of the efficiency of IDA<sup>∗</sup> and of its difficulties with real-valued heuristics appears in Patrick *et al.* (1992).

RBFS (Korf, 1993) is actually somewhat more complicated than the algorithm shown in Figure 3.26, which is closer to an independently developed algorithm called **iterative ex-PERATTLE FEAT (RUSSELLEED) pansion** (Russell, 1992). RBFS uses a lower bound as well as the upper bound; the two algorithms behave identically with admissible heuristics, but RBFS expands nodes in best-first

**EXPANSION** 

order even with an inadmissible heuristic. The idea of keeping track of the best alternative path appeared earlier in Bratko's (1986) elegant Prolog implementation of A<sup>\*</sup> and in the DTA<sup>\*</sup> algorithm (Russell and Wefald, 1991). The latter work also discusses metalevel state spaces and metalevel learning.

The MA<sup>∗</sup> algorithm appeared in Chakrabarti *et al.* (1989). SMA<sup>∗</sup>, or Simplified MA<sup>∗</sup>, emerged from an attempt to implement MA<sup>∗</sup> as a comparison algorithm for IE (Russell, 1992). Kaindl and Khorsand (1994) have applied SMA<sup>∗</sup> to produce a bidirectional search algorithm that is substantially faster than previous algorithms. Korf and Zhang (2000) describe a divideand-conquer approach, and Zhou and Hansen (2002) introduce memory-bounded  $A^*$  graph search and a strategy for switching to breadth-first search to increase memory-efficiency (Zhou and Hansen, 2006). Korf (1995) surveys memory-bounded search techniques.

The idea that admissible heuristics can be derived by problem relaxation appears in the seminal paper by Held and Karp (1970), who used the minimum-spanning-tree heuristic to solve the TSP. (See Exercise 3.33.)

The automation of the relaxation process was implemented successfully by Prieditis (1993), building on earlier work with Mostow (Mostow and Prieditis, 1989). Holte and Hernadvolgyi (2001) describe more recent steps towards automating the process. The use of pattern databases to derive admissible heuristics is due to Gasser (1995) and Culberson and Schaeffer (1996, 1998); disjoint pattern databases are described by Korf and Felner (2002); a similar method using symbolic patterns is due to Edelkamp (2009). Felner *et al.* (2007) show how to compress pattern databases to save space. The probabilistic interpretation of heuristics was investigated in depth by Pearl (1984) and Hansson and Mayer (1989).

By far the most comprehensive source on heuristics and heuristic search algorithms is Pearl's (1984) *Heuristics* text. This book provides especially good coverage of the wide variety of offshoots and variations of A<sup>∗</sup>, including rigorous proofs of their formal properties. Kanal and Kumar (1988) present an anthology of important articles on heuristic search, and Rayward-Smith *et al.* (1996) cover approaches from Operations Research. Papers about new search algorithms—which, remarkably, continue to be discovered—appear in journals such as *Artificial Intelligence* and *Journal of the ACM*.

PARALLEL SEARCH The topic of **parallel search** algorithms was not covered in the chapter, partly because it requires a lengthy discussion of parallel computer architectures. Parallel search became a popular topic in the 1990s in both AI and theoretical computer science (Mahanti and Daniels, 1993; Grama and Kumar, 1995; Crauser *et al.*, 1998) and is making a comeback in the era of new multicore and cluster architectures (Ralphs *et al.*, 2004; Korf and Schultze, 2005). Also of increasing importance are search algorithms for very large graphs that require disk storage (Korf, 2008).

#### **EXERCISES**

- **3.1** Explain why problem formulation must follow goal formulation.
- **3.2** Give a complete problem formulation for each of the following problems. Choose a

formulation that is precise enough to be implemented.

- **a**. There are six glass boxes in a row, each with a lock. Each of the first five boxes holds a key unlocking the next box in line; the last box holds a banana. You have the key to the first box, and you want the banana.
- **b**. You start with the sequence ABABAECCEC, or in general any sequence made from A, B, C, and E. You can transform this sequence using the following equalities:  $AC = E$ ,  $AB = BC$ ,  $BB = E$ , and  $Ex = x$  for any x. For example, ABBC can be transformed into AEC, and then AC, and then E. Your goal is to produce the sequence E.
- **c**. There is an  $n \times n$  grid of squares, each square initially being either unpainted floor or a bottomless pit. You start standing on an unpainted floor square, and can either paint the square under you or move onto an adjacent unpainted floor square. You want the whole floor painted.
- **d**. A container ship is in port, loaded high with containers. There 13 rows of containers, each 13 containers wide and 5 containers tall. You control a crane that can move to any location above the ship, pick up the container under it, and move it onto the dock. You want the ship unloaded.

**3.3** You have a  $9 \times 9$  grid of squares, each of which can be colored red or blue. The grid is initially colored all blue, but you can change the color of any square any number of times. Imagining the grid divided into nine  $3 \times 3$  sub-squares, you want each sub-square to be all one color but neighboring sub-squares to be different colors.

- **a**. Formulate this problem in the straightforward way. Compute the size of the state space.
- **b**. You need color a square only once. Reformulate, and compute the size of the state space. Would breadth-first graph search perform faster on this problem than on the one in (a)? How about iterative deepening tree search?
- **c**. Given the goal, we need consider only colorings where each sub-square is uniformly colored. Reformulate the problem and compute the size of the state space.
- **d**. How many solutions does this problem have?
- **e**. Parts (b) and (c) successively abstracted the original problem (a). Can you give a translation from solutions in problem (c) into solutions in problem (b), and from solutions in problem (b) into solutions for problem (a)?

**3.4** Suppose two friends live in different cities on a map, such as the Romania map shown in Figure 3.2. On every turn, we can simultaneously move each friend to a neighboring city on the map. The amount of time needed to move from city i to neighbor j is equal to the road distance  $d(i, j)$  between the cities, but on each turn the friend that arrives first must wait until the other one arrives (and calls the first on his/her cell phone) before the next turn can begin. We want the two friends to meet as quickly as possible.

- **a**. Write a detailed formulation for this search problem. (You will find it helpful to define some formal notation here.)
- **b**. Let  $D(i, j)$  be the straight-line distance between cities i and j. Which of the following heuristic functions are admissible? (i)  $D(i, j)$ ; (ii)  $2 \cdot D(i, j)$ ; (iii)  $D(i, j)/2$ .



- **c**. Are there completely connected maps for which no solution exists?
- **d**. Are there maps in which all solutions require one friend to visit the same city twice?

**3.5** Show that the 8-puzzle states are divided into two disjoint sets, such that any state is reachable from any other state in the same set, while no state is reachable from any state in the other set. (*Hint:* See Berlekamp *et al.* (1982).) Devise a procedure to decide which set a given state is in, and explain why this is useful for generating random states.

**3.6** Consider the *n*-queens problem using the "efficient" incremental formulation given on **5.0** Consider the *n*-queens problem using the entirement incremental formulation given on page 72. Explain why the state space has at least  $\sqrt[3]{n!}$  states and estimate the largest *n* for which exhaustive exploration is feasible. (*Hint*: Derive a lower bound on the branching factor by considering the maximum number of squares that a queen can attack in any column.)

**3.7** Consider the problem of finding the shortest path between two points on a plane that has convex polygonal obstacles as shown in Figure 3.31. This is an idealization of the problem that a robot has to solve to navigate in a crowded environment.

- **a**. Suppose the state space consists of all positions  $(x, y)$  in the plane. How many states are there? How many paths are there to the goal?
- **b**. Explain briefly why the shortest path from one polygon vertex to any other in the scene must consist of straight-line segments joining some of the vertices of the polygons. Define a good state space now. How large is this state space?
- **c**. Define the necessary functions to implement the search problem, including an ACTIONS function that takes a vertex as input and returns a set of vectors, each of which maps the current vertex to one of the vertices that can be reached in a straight line. (Do not forget the neighbors on the same polygon.) Use the straight-line distance for the heuristic function.
- **d**. Apply one or more of the algorithms in this chapter to solve a range of problems in the domain, and comment on their performance.



**3.8** On page 68, we said that we would not consider problems with negative path costs. In this exercise, we explore this decision in more depth.

- **a**. Suppose that actions can have arbitrarily large negative costs; explain why this possibility would force any optimal algorithm to explore the entire state space.
- **b**. Does it help if we insist that step costs must be greater than or equal to some negative constant c? Consider both trees and graphs.
- **c**. Suppose that a set of actions forms a loop in the state space such that executing the set in some order results in no net change to the state. If all of these actions have negative cost, what does this imply about the optimal behavior for an agent in such an environment?
- **d**. One can easily imagine actions with high negative cost, even in domains such as route finding. For example, some stretches of road might have such beautiful scenery as to far outweigh the normal costs in terms of time and fuel. Explain, in precise terms, within the context of state-space search, why humans do not drive around scenic loops indefinitely, and explain how to define the state space and actions for route finding so that artificial agents can also avoid looping.
- **e**. Can you think of a real domain in which step costs are such as to cause looping?

**3.9** The **missionaries and cannibals** problem is usually stated as follows. Three missionaries and three cannibals are on one side of a river, along with a boat that can hold one or two people. Find a way to get everyone to the other side without ever leaving a group of missionaries in one place outnumbered by the cannibals in that place. This problem is famous in AI because it was the subject of the first paper that approached problem formulation from an analytical viewpoint (Amarel, 1968).

- **a**. Formulate the problem precisely, making only those distinctions necessary to ensure a valid solution. Draw a diagram of the complete state space.
- **b**. Implement and solve the problem optimally using an appropriate search algorithm. Is it a good idea to check for repeated states?
- **c**. Why do you think people have a hard time solving this puzzle, given that the state space is so simple?

**3.10** Define in your own words the following terms: state, state space, search tree, search node, goal, action, transition model, and branching factor.

**3.11** What's the difference between a world state, a state description, and a search node? Why is this distinction useful?

**3.12** An action such as  $Go( Sibiu)$  really consists of a long sequence of finer-grained actions: turn on the car, release the brake, accelerate forward, etc. Having composite actions of this kind reduces the number of steps in a solution sequence, thereby reducing the search time. Suppose we take this to the logical extreme, by making super-composite actions out of every possible sequence of *Go* actions. Then every problem instance is solved by a single supercomposite action, such as *Go(Sibiu)Go(Rimnicu Vilcea)Go(Pitesti)Go(Bucharest)*. Explain how search would work in this formulation. Is this a practical approach for speeding up problem solving?





**3.13** Does a finite state space always lead to a finite search tree? How about a finite state space that is a tree? Can you be more precise about what types of state spaces always lead to finite search trees? (Adapted from Bender, 1996.)

**3.14** Prove that GRAPH-SEARCH satisfies the graph separation property illustrated in Figure 3.9. (*Hint*: Begin by showing that the property holds at the start, then show that if it holds before an iteration of the algorithm, it holds afterwards.) Describe a search algorithm that violates the property.

- **3.15** Which of the following are true and which are false? Explain your answers.
	- a. Depth-first search always expands at least as many nodes as A<sup>\*</sup> search with an admissible heuristic.
	- **b**.  $h(n) = 0$  is an admissible heuristic for the 8-puzzle.
	- **c**. A<sup>∗</sup> is of no use in robotics because percepts, states, and actions are continuous.
	- **d**. Breadth-first search is complete even if zero step costs are allowed.
	- **e**. Assume that a rook can move on a chessboard any number of squares in a straight line, vertically or horizontally, but cannot jump over other pieces. Manhattan distance is an admissible heuristic for the problem of moving the rook from square A to square B in the smallest number of moves.

**3.16** A basic wooden railway set contains the pieces shown in Figure 3.32. The task is to connect these pieces into a railway that has no overlapping tracks and no loose ends where a train could run off onto the floor.

- **a**. Suppose that the pieces fit together *exactly* with no slack. Give a precise formulation of the task as a search problem.
- **b**. Identify a suitable uninformed search algorithm for this task and explain your choice.
- **c**. Explain why removing any one of the "fork" pieces makes the problem unsolvable.
- **d**. Give an upper bound on the total size of the state space defined by your formulation. (*Hint*: think about the maximum branching factor for the construction process and the maximum depth, ignoring the problem of overlapping pieces and loose ends. Begin by pretending that every piece is unique.)



**3.17** Implement two versions of the  $RESULT(s, a)$  function for the 8-puzzle: one that copies

and edits the data structure for the parent node s and one that modifies the parent state directly (undoing the modifications as needed). Write versions of iterative deepening depth-first search that use these functions and compare their performance.



**3.18** On page 90, we mentioned **iterative lengthening search**, an iterative analog of uniform cost search. The idea is to use increasing limits on path cost. If a node is generated whose path cost exceeds the current limit, it is immediately discarded. For each new iteration, the limit is set to the lowest path cost of any node discarded in the previous iteration.

- **a**. Show that this algorithm is optimal for general path costs.
- **b**. Consider a uniform tree with branching factor  $b$ , solution depth  $d$ , and unit step costs. How many iterations will iterative lengthening require?
- **c**. Now consider step costs drawn from the continuous range  $[\epsilon, 1]$ , where  $0 < \epsilon < 1$ . How many iterations are required in the worst case?
- **d**. Implement the algorithm and apply it to instances of the 8-puzzle and traveling salesperson problems. Compare the algorithm's performance to that of uniform-cost search, and comment on your results.

**3.19** Describe a state space in which iterative deepening search performs much worse than depth-first search (for example,  $O(n^2)$  vs.  $O(n)$ ).

**3.20** Write a program that will take as input two Web page URLs and find a path of links from one to the other. What is an appropriate search strategy? Is bidirectional search a good idea? Could a search engine be used to implement a predecessor function?

**3.21** Consider the vacuum-world problem defined in Figure 2.2.

- **a**. Which of the algorithms defined in this chapter would be appropriate for this problem? Should the algorithm use tree search or graph search?
- **b**. Apply your chosen algorithm to compute an optimal sequence of actions for a  $3 \times 3$ world whose initial state has dirt in the three top squares and the agent in the center.
- **c**. Construct a search agent for the vacuum world, and evaluate its performance in a set of  $3 \times 3$  worlds with probability 0.2 of dirt in each square. Include the search cost as well as path cost in the performance measure, using a reasonable exchange rate.
- **d**. Compare your best search agent with a simple randomized reflex agent that sucks if there is dirt and otherwise moves randomly.
- **e**. Consider what would happen if the world were enlarged to  $n \times n$ . How does the performance of the search agent and of the reflex agent vary with  $n$ ?
- **3.22** Prove each of the following statements, or give a counterexample:
	- **a**. Breadth-first search is a special case of uniform-cost search.
	- **b**. Depth-first search is a special case of best-first tree search.
	- **c**. Uniform-cost search is a special case of A<sup>∗</sup> search.









**3.23** Compare the performance of A<sup>∗</sup> and RBFS on a set of randomly generated problems

in the 8-puzzle (with Manhattan distance) and TSP (with MST—see Exercise 3.33) domains. Discuss your results. What happens to the performance of RBFS when a small random number is added to the heuristic values in the 8-puzzle domain?

**3.24** Trace the operation of A<sup>∗</sup> search applied to the problem of getting to Bucharest from Lugoj using the straight-line distance heuristic. That is, show the sequence of nodes that the algorithm will consider and the  $f$ ,  $g$ , and  $h$  score for each node.

**3.25** Sometimes there is no good evaluation function for a problem but there is a good comparison method: a way to tell whether one node is better than another without assigning numerical values to either. Show that this is enough to do a best-first search. Is there an analog of A<sup>∗</sup> for this setting?

**3.26** Devise a state space in which A<sup>∗</sup> using GRAPH-SEARCH returns a suboptimal solution with an  $h(n)$  function that is admissible but inconsistent.

**3.27** Accurate heuristics don't necessarily reduce search time in the worst case. Given any depth d, define a search problem with a goal node at depth d, and write a heuristic function such that  $|h(n) - h^{*}(n)| \le O(\log h^{*}(n))$  but A<sup>\*</sup> expands all nodes of depth less than d.

**HEURISTIC PATH 3.28** The **heuristic path algorithm** (Pohl, 1977) is a best-first search in which the evaluation function is  $f(n) = (2 - w)g(n) + wh(n)$ . For what values of w is this complete? For what values is it optimal, assuming that  $h$  is admissible? What kind of search does this perform for  $w = 0$ ,  $w = 1$ , and  $w = 2$ ?

> **3.29** Consider the unbounded version of the regular 2D grid shown in Figure 3.9. The start state is at the origin,  $(0,0)$ , and the goal state is at  $(x, y)$ .

- **a**. What is the branching factor b in this state space?
- **b**. How many distinct states are there at depth  $k$  (for  $k > 0$ )?
- **c**. What is the maximum number of nodes expanded by breadth-first tree search?
- **d**. What is the maximum number of nodes expanded by breadth-first graph search?
- **e**. Is  $h = |u x| + |v y|$  an admissible heuristic for a state at  $(u, v)$ ? Explain.
- **f**. How many nodes are expanded by A<sup>∗</sup> graph search using h?
- **g**. Does h remain admissible if some links are removed?
- **h**. Does h remain admissible if some links are added between nonadjacent states?

**3.30** Consider the problem of moving k knights from k starting squares  $s_1, \ldots, s_k$  to k goal squares  $g_1, \ldots, g_k$ , on an unbounded chessboard, subject to the rule that no two knights can land on the same square at the same time. Each action consists of moving *up to* k knights simultaneously. We would like to complete the maneuver in the smallest number of actions.

- **a**. What is the maximum branching factor in this state space, expressed as a function of k?
- **b**. Suppose  $h_i$  is an admissible heuristic for the problem of moving knight i to goal  $g_i$  by itself. Which of the following heuristics are admissible for the  $k$ -knight problem? Of those, which is the best?
	- (i)  $\min\{h_1, \ldots, h_k\}.$



ALGORITHM

- (ii)  $\max\{h_1, ..., h_k\}.$
- (iii)  $\sum_{i=1}^{k} h_i$ .
- **c**. Repeat (b) for the case where you are allowed to move only one knight at a time.

**3.31** We saw on page 93 that the straight-line distance heuristic leads greedy best-first search astray on the problem of going from Iasi to Fagaras. However, the heuristic is perfect on the opposite problem: going from Fagaras to Iasi. Are there problems for which the heuristic is misleading in both directions?

**3.32** Prove that if a heuristic is consistent, it must be admissible. Construct an admissible heuristic that is not consistent.

**3.33** The traveling salesperson problem (TSP) can be solved with the minimum-spanningtree (MST) heuristic, which estimates the cost of completing a tour, given that a partial tour has already been constructed. The MST cost of a set of cities is the smallest sum of the link costs of any tree that connects all the cities.

- **a**. Show how this heuristic can be derived from a relaxed version of the TSP.
- **b**. Show that the MST heuristic dominates straight-line distance.
- **c**. Write a problem generator for instances of the TSP where cities are represented by random points in the unit square.
- **d**. Find an efficient algorithm in the literature for constructing the MST, and use it with A<sup>∗</sup> graph search to solve instances of the TSP.

**3.34** On page 105, we defined the relaxation of the 8-puzzle in which a tile can move from square A to square B if B is blank. The exact solution of this problem defines **Gaschnig's heuristic** (Gaschnig, 1979). Explain why Gaschnig's heuristic is at least as accurate as  $h_1$ (misplaced tiles), and show cases where it is more accurate than both  $h_1$  and  $h_2$  (Manhattan distance). Explain how to calculate Gaschnig's heuristic efficiently.

**3.35** We gave two simple heuristics for the 8-puzzle: Manhattan distance and misplaced tiles. Several heuristics in the literature purport to improve on this—see, for example, Nilsson (1971), Mostow and Prieditis (1989), and Hansson *et al.* (1992). Test these claims by implementing the heuristics and comparing the performance of the resulting algorithms.





# LOGICAL AGENTS

*In which we design agents that can form representations of a complex world, use a process of inference to derive new representations about the world, and use these new representations to deduce what to do.*

Humans, it seems, know things; and what they know helps them do things. These are not empty statements. They make strong claims about how the intelligence of humans is REASONING achieved—not by purely reflex mechanisms but by processes of **reasoning** that operate on REPRESENTATION internal **representations** of knowledge. In AI, this approach to intelligence is embodied in **knowledge-based agents**.

> The problem-solving agents of Chapters 3 and 4 know things, but only in a very limited, inflexible sense. For example, the transition model for the 8-puzzle—knowledge of what the actions do—is hidden inside the domain-specific code of the RESULT function. It can be used to predict the outcome of actions but not to deduce that two tiles cannot occupy the same space or that states with odd parity cannot be reached from states with even parity. The atomic representations used by problem-solving agents are also very limiting. In a partially observable environment, an agent's only choice for representing what it knows about the current state is to list all possible concrete states—a hopeless prospect in large environments.

Chapter 6 introduced the idea of representing states as assignments of values to variables; this is a step in the right direction, enabling some parts of the agent to work in a domain-independent way and allowing for more efficient algorithms. In this chapter and LOGIC those that follow, we take this step to its logical conclusion, so to speak—we develop **logic** as a general class of representations to support knowledge-based agents. Such agents can combine and recombine information to suit myriad purposes. Often, this process can be quite far removed from the needs of the moment—as when a mathematician proves a theorem or an astronomer calculates the earth's life expectancy. Knowledge-based agents can accept new tasks in the form of explicitly described goals; they can achieve competence quickly by being told or learning new knowledge about the environment; and they can adapt to changes in the environment by updating the relevant knowledge.

> We begin in Section 7.1 with the overall agent design. Section 7.2 introduces a simple new environment, the wumpus world, and illustrates the operation of a knowledge-based agent without going into any technical detail. Then we explain the general principles of **logic**

KNOWLEDGE-BASED AGENTS

in Section 7.3 and the specifics of **propositional logic** in Section 7.4. While less expressive than **first-order logic** (Chapter 8), propositional logic illustrates all the basic concepts of logic; it also comes with well-developed inference technologies, which we describe in sections 7.5 and 7.6. Finally, Section 7.7 combines the concept of knowledge-based agents with the technology of propositional logic to build some simple agents for the wumpus world.

# 7.1 KNOWLEDGE-BASED AGENTS

**REPRESENTATION** LANGUAGE

BACKGROUND KNOWLEDGE

KNOWLEDGE BASE The central component of a knowledge-based agent is its **knowledge base**, or KB. A knowl-SENTENCE edge base is a set of **sentences**. (Here "sentence" is used as a technical term. It is related but not identical to the sentences of English and other natural languages.) Each sentence is expressed in a language called a **knowledge representation language** and represents some REPRESENTATION AXIOM assertion about the world. Sometimes we dignify a sentence with the name **axiom**, when the sentence is taken as given without being derived from other sentences.

There must be a way to add new sentences to the knowledge base and a way to query what is known. The standard names for these operations are TELL and ASK, respectively. INFERENCE Both operations may involve **inference**—that is, deriving new sentences from old. Inference must obey the requirement that when one ASKs a question of the knowledge base, the answer should follow from what has been told (or TELLed) to the knowledge base previously. Later in this chapter, we will be more precise about the crucial word "follow." For now, take it to mean that the inference process should not make things up as it goes along.

> Figure 7.1 shows the outline of a knowledge-based agent program. Like all our agents, it takes a percept as input and returns an action. The agent maintains a knowledge base, KB, which may initially contain some **background knowledge**.

> Each time the agent program is called, it does three things. First, it TELLs the knowledge base what it perceives. Second, it ASKs the knowledge base what action it should perform. In the process of answering this query, extensive reasoning may be done about the current state of the world, about the outcomes of possible action sequences, and so on. Third, the agent program TELLs the knowledge base which action was chosen, and the agent executes the action.

> The details of the representation language are hidden inside three functions that implement the interface between the sensors and actuators on one side and the core representation and reasoning system on the other. MAKE-PERCEPT-SENTENCE constructs a sentence asserting that the agent perceived the given percept at the given time. MAKE-ACTION-QUERY constructs a sentence that asks what action should be done at the current time. Finally, MAKE-ACTION-SENTENCE constructs a sentence asserting that the chosen action was executed. The details of the inference mechanisms are hidden inside TELL and ASK. Later sections will reveal these details.

> The agent in Figure 7.1 appears quite similar to the agents with internal state described in Chapter 2. Because of the definitions of TELL and ASK, however, the knowledge-based agent is not an arbitrary program for calculating actions. It is amenable to a description at

**function** KB-AGENT(*percept*) **returns** an *action* **persistent**: KB, a knowledge base  $t$ , a counter, initially 0, indicating time TELL $(KB, \text{MAKE-PERCEPT-SENTENCE}(percent, t))$ 

 $action \leftarrow ASK(KB, MAKE-ACTION-QUERV(t))$ TELL(KB, MAKE-ACTION-SENTENCE(action,t))  $t \leftarrow t + 1$ **return** action

**Figure 7.1** A generic knowledge-based agent. Given a percept, the agent adds the percept to its knowledge base, asks the knowledge base for the best action, and tells the knowledge base that it has in fact taken that action.

KNOWLEDGE LEVEL the **knowledge level**, where we need specify only what the agent knows and what its goals are, in order to fix its behavior. For example, an automated taxi might have the goal of taking a passenger from San Francisco to Marin County and might know that the Golden Gate Bridge is the only link between the two locations. Then we can expect it to cross the Golden Gate Bridge *because it knows that that will achieve its goal*. Notice that this analysis IMPLEMENTATION is independent of how the taxi works at the **implementation level**. It doesn't matter whether LEVEL its geographical knowledge is implemented as linked lists or pixel maps, or whether it reasons by manipulating strings of symbols stored in registers or by propagating noisy signals in a network of neurons.

A knowledge-based agent can be built simply by TELLing it what it needs to know. Starting with an empty knowledge base, the agent designer can TELL sentences one by one DECLARATIVE until the agent knows how to operate in its environment. This is called the **declarative** approach to system building. In contrast, the **procedural** approach encodes desired behaviors directly as program code. In the 1970s and 1980s, advocates of the two approaches engaged in heated debates. We now understand that a successful agent often combines both declarative and procedural elements in its design, and that declarative knowledge can often be compiled into more efficient procedural code.

> We can also provide a knowledge-based agent with mechanisms that allow it to learn for itself. These mechanisms, which are discussed in Chapter 18, create general knowledge about the environment from a series of percepts. A learning agent can be fully autonomous.

# 7.2 THE WUMPUS WORLD

In this section we describe an environment in which knowledge-based agents can show their WUMPUS WORLD worth. The **wumpus world** is a cave consisting of rooms connected by passageways. Lurking somewhere in the cave is the terrible wumpus, a beast that eats anyone who enters its room. The wumpus can be shot by an agent, but the agent has only one arrow. Some rooms contain

bottomless pits that will trap anyone who wanders into these rooms (except for the wumpus, which is too big to fall in). The only mitigating feature of this bleak environment is the possibility of finding a heap of gold. Although the wumpus world is rather tame by modern computer game standards, it illustrates some important points about intelligence.

A sample wumpus world is shown in Figure 7.2. The precise definition of the task environment is given, as suggested in Section 2.3, by the PEAS description:

- **Performance measure**: +1000 for climbing out of the cave with the gold, –1000 for falling into a pit or being eaten by the wumpus,  $-1$  for each action taken and  $-10$  for using up the arrow. The game ends either when the agent dies or when the agent climbs out of the cave.
- **Environment**: A  $4 \times 4$  grid of rooms. The agent always starts in the square labeled [1,1], facing to the right. The locations of the gold and the wumpus are chosen randomly, with a uniform distribution, from the squares other than the start square. In addition, each square other than the start can be a pit, with probability 0.2.
- **Actuators**: The agent can move *Forward*, *TurnLeft* by 90◦ , or *TurnRight* by 90◦ . The agent dies a miserable death if it enters a square containing a pit or a live wumpus. (It is safe, albeit smelly, to enter a square with a dead wumpus.) If an agent tries to move forward and bumps into a wall, then the agent does not move. The action *Grab* can be used to pick up the gold if it is in the same square as the agent. The action *Shoot* can be used to fire an arrow in a straight line in the direction the agent is facing. The arrow continues until it either hits (and hence kills) the wumpus or hits a wall. The agent has only one arrow, so only the first *Shoot* action has any effect. Finally, the action *Climb* can be used to climb out of the cave, but only from square [1,1].
- **Sensors**: The agent has five sensors, each of which gives a single bit of information:
	- **–** In the square containing the wumpus and in the directly (not diagonally) adjacent squares, the agent will perceive a *Stench*.
	- **–** In the squares directly adjacent to a pit, the agent will perceive a *Breeze*.
	- **–** In the square where the gold is, the agent will perceive a *Glitter*.
	- **–** When an agent walks into a wall, it will perceive a *Bump*.
	- **–** When the wumpus is killed, it emits a woeful *Scream* that can be perceived anywhere in the cave.

The percepts will be given to the agent program in the form of a list of five symbols; for example, if there is a stench and a breeze, but no glitter, bump, or scream, the agent program will get [Stench, Breeze, None, None, None].

We can characterize the wumpus environment along the various dimensions given in Chapter 2. Clearly, it is discrete, static, and single-agent. (The wumpus doesn't move, fortunately.) It is sequential, because rewards may come only after many actions are taken. It is partially observable, because some aspects of the state are not directly perceivable: the agent's location, the wumpus's state of health, and the availability of an arrow. As for the locations of the pits and the wumpus: we could treat them as unobserved parts of the state that happen to be immutable—in which case, the transition model for the environment is completely



known; or we could say that the transition model itself is unknown because the agent doesn't know which *Forward* actions are fatal—in which case, discovering the locations of pits and wumpus completes the agent's knowledge of the transition model.

For an agent in the environment, the main challenge is its initial ignorance of the configuration of the environment; overcoming this ignorance seems to require logical reasoning. In most instances of the wumpus world, it is possible for the agent to retrieve the gold safely. Occasionally, the agent must choose between going home empty-handed and risking death to find the gold. About 21% of the environments are utterly unfair, because the gold is in a pit or surrounded by pits.

Let us watch a knowledge-based wumpus agent exploring the environment shown in Figure 7.2. We use an informal knowledge representation language consisting of writing down symbols in a grid (as in Figures 7.3 and 7.4).

The agent's initial knowledge base contains the rules of the environment, as described previously; in particular, it knows that it is in  $[1,1]$  and that  $[1,1]$  is a safe square; we denote that with an "A" and "OK," respectively, in square [1,1].

The first percept is [None, None, None, None, None], from which the agent can conclude that its neighboring squares,  $[1,2]$  and  $[2,1]$ , are free of dangers—they are OK. Figure 7.3(a) shows the agent's state of knowledge at this point.

A cautious agent will move only into a square that it knows to be OK. Let us suppose the agent decides to move forward to [2,1]. The agent perceives a breeze (denoted by "B") in [2,1], so there must be a pit in a neighboring square. The pit cannot be in [1,1], by the rules of the game, so there must be a pit in  $[2,2]$  or  $[3,1]$  or both. The notation "P?" in Figure 7.3(b) indicates a possible pit in those squares. At this point, there is only one known square that is OK and that has not yet been visited. So the prudent agent will turn around, go back to [1,1], and then proceed to [1,2].

The agent perceives a stench in [1,2], resulting in the state of knowledge shown in Figure 7.4(a). The stench in [1,2] means that there must be a wumpus nearby. But the

1,4	2,4	3,4	4,4	A  $= Agent$ B $= Breeze$ = Glitter, Gold G $OK = Safe square$	1,4	2,4	3,4	4,4
1,3	2,3	3,3	4,3	$= Pit$ P $=$ Stench S V = Visited W $=$ Wumpus	1,3	2,3	3,3	4,3
1,2 OK	2,2	3,2	4,2		1,2 OK	2,2 P?	3,2	4,2
1,1 $\vert$ A OK	2,1 OK	3,1	4,1		1,1 $\mathbf{V}$ OK	2,1 $\vert$ A $\bf{B}$ OK	3,1 P?	4,1
	(a)				(b)			

**Figure 7.3** The first step taken by the agent in the wumpus world. (a) The initial situation, after percept [None, None, None, None, None]. (b) After one move, with percept [None, Breeze, None, None, None].

1,4	2,4	3,4	4,4	$\mathbf{A}$ $= Agent$ $= Breeze$ B = Glitter, Gold G $OK = Safe square$	1,4	2,4 P?	3,4	4,4
$1^{1,3}$ W!	2,3	3,3	4,3	$= Pit$ P S $=$ Stench $= Visited$ V W $=$ Wumpus	$11,3$ W!	$\frac{12,3}{\underline{A}}$ S G B	$ 3,3 \text{ p}$ ?	4,3
$11,2$ $\boxed{A}$ ${\bf S}$ OK	2,2 OK	3,2	4,2		1,2 S V OK	2,2 $\bf{V}$ OK	3,2	4,2
1,1 V OK	2,1 $\bf{B}$ $\bf{V}$ OK	3,1 P!	4,1		1,1 V OK	2,1 $\bf{B}$ V OK	3,1 P!	4,1
(a)					(b)			

**Figure 7.4** Two later stages in the progress of the agent. (a) After the third move, with percept [Stench, None, None, None, None]. (b) After the fifth move, with percept [Stench, Breeze, Glitter , None, None].

wumpus cannot be in  $[1,1]$ , by the rules of the game, and it cannot be in  $[2,2]$  (or the agent would have detected a stench when it was in  $[2,1]$ ). Therefore, the agent can infer that the wumpus is in [1,3]. The notation W! indicates this inference. Moreover, the lack of a breeze in  $[1,2]$  implies that there is no pit in  $[2,2]$ . Yet the agent has already inferred that there must be a pit in either  $[2,2]$  or  $[3,1]$ , so this means it must be in  $[3,1]$ . This is a fairly difficult inference, because it combines knowledge gained at different times in different places and relies on the lack of a percept to make one crucial step.

The agent has now proved to itself that there is neither a pit nor a wumpus in [2,2], so it is OK to move there. We do not show the agent's state of knowledge at [2,2]; we just assume that the agent turns and moves to [2,3], giving us Figure 7.4(b). In [2,3], the agent detects a glitter, so it should grab the gold and then return home.

Note that in each case for which the agent draws a conclusion from the available information, that conclusion is *guaranteed* to be correct if the available information is correct. This is a fundamental property of logical reasoning. In the rest of this chapter, we describe how to build logical agents that can represent information and draw conclusions such as those described in the preceding paragraphs.

# 7.3 LOGIC

This section summarizes the fundamental concepts of logical representation and reasoning. These beautiful ideas are independent of any of logic's particular forms. We therefore postpone the technical details of those forms until the next section, using instead the familiar example of ordinary arithmetic.

In Section 7.1, we said that knowledge bases consist of sentences. These sentences SYNTAX are expressed according to the **syntax** of the representation language, which specifies all the sentences that are well formed. The notion of syntax is clear enough in ordinary arithmetic: " $x + y = 4$ " is a well-formed sentence, whereas " $x4y + z$ " is not.

SEMANTICS A logic must also define the **semantics** or meaning of sentences. The semantics defines TRUTH the **truth** of each sentence with respect to each **possible world**. For example, the semantics POSSIBLE WORLD for arithmetic specifies that the sentence " $x + y = 4$ " is true in a world where x is 2 and y is 2, but false in a world where  $x$  is 1 and  $y$  is 1. In standard logics, every sentence must be either true or false in each possible world—there is no "in between."<sup>1</sup>

MODEL When we need to be precise, we use the term **model** in place of "possible world." Whereas possible worlds might be thought of as (potentially) real environments that the agent might or might not be in, models are mathematical abstractions, each of which simply fixes the truth or falsehood of every relevant sentence. Informally, we may think of a possible world as, for example, having x men and y women sitting at a table playing bridge, and the sentence  $x + y = 4$  is true when there are four people in total. Formally, the possible models are just all possible assignments of real numbers to the variables x and y. Each such assignment fixes the truth of any sentence of arithmetic whose variables are x and y. If a sentence  $\alpha$  is true in SATISFACTION model m, we say that m **satisfies**  $\alpha$  or sometimes m **is a model of**  $\alpha$ . We use the notation  $M(\alpha)$  to mean the set of all models of  $\alpha$ .

Now that we have a notion of truth, we are ready to talk about logical reasoning. This ENTAILMENT involves the relation of logical **entailment** between sentences—the idea that a sentence *follows logically* from another sentence. In mathematical notation, we write

 $\alpha \models \beta$ 

<sup>1</sup> **Fuzzy logic**, discussed in Chapter 14, allows for degrees of truth.



to mean that the sentence  $\alpha$  entails the sentence  $\beta$ . The formal definition of entailment is this:  $\alpha \models \beta$  if and only if, in every model in which  $\alpha$  is true,  $\beta$  is also true. Using the notation just introduced, we can write

$$
\alpha \models \beta
$$
 if and only if  $M(\alpha) \subseteq M(\beta)$ .

(Note the direction of the  $\subseteq$  here: if  $\alpha \models \beta$ , then  $\alpha$  is a *stronger* assertion than  $\beta$ : it rules out *more* possible worlds.) The relation of entailment is familiar from arithmetic; we are happy with the idea that the sentence  $x = 0$  entails the sentence  $xy = 0$ . Obviously, in any model where x is zero, it is the case that  $xy$  is zero (regardless of the value of y).

We can apply the same kind of analysis to the wumpus-world reasoning example given in the preceding section. Consider the situation in Figure  $7.3(b)$ : the agent has detected nothing in [1,1] and a breeze in [2,1]. These percepts, combined with the agent's knowledge of the rules of the wumpus world, constitute the KB. The agent is interested (among other things) in whether the adjacent squares [1,2], [2,2], and [3,1] contain pits. Each of the three squares might or might not contain a pit, so (for the purposes of this example) there are  $2^3 = 8$ possible models. These eight models are shown in Figure 7.5.<sup>2</sup>

The KB can be thought of as a set of sentences or as a single sentence that asserts all the individual sentences. The KB is false in models that contradict what the agent knows for example, the KB is false in any model in which [1,2] contains a pit, because there is no breeze in [1,1]. There are in fact just three models in which the KB is true, and these are

<sup>2</sup> Although the figure shows the models as partial wumpus worlds, they are really nothing more than assignments of true and false to the sentences "there is a pit in [1,2]" etc. Models, in the mathematical sense, do not need to have 'orrible 'airy wumpuses in them.

shown surrounded by a solid line in Figure 7.5. Now let us consider two possible conclusions:

 $\alpha_1$  = "There is no pit in [1,2]."

 $\alpha_2$  = "There is no pit in [2,2]."

We have surrounded the models of  $\alpha_1$  and  $\alpha_2$  with dotted lines in Figures 7.5(a) and 7.5(b), respectively. By inspection, we see the following:

in every model in which KB is true,  $\alpha_1$  is also true.

Hence,  $KB \models \alpha_1$ : there is no pit in [1,2]. We can also see that

in some models in which  $KB$  is true,  $\alpha_2$  is false.

Hence,  $KB \not\models \alpha_2$ : the agent *cannot* conclude that there is no pit in [2,2]. (Nor can it conclude that there *is* a pit in  $[2,2]$ .)<sup>3</sup>

The preceding example not only illustrates entailment but also shows how the definition LOGICAL INFERENCE of entailment can be applied to derive conclusions—that is, to carry out **logical inference**. MODEL CHECKING The inference algorithm illustrated in Figure 7.5 is called **model checking**, because it enumerates all possible models to check that  $\alpha$  is true in all models in which KB is true, that is, that  $M(KB) \subseteq M(\alpha)$ .

> In understanding entailment and inference, it might help to think of the set of all consequences of KB as a haystack and of  $\alpha$  as a needle. Entailment is like the needle being in the haystack; inference is like finding it. This distinction is embodied in some formal notation: if an inference algorithm i can derive  $\alpha$  from KB, we write

 $KB \vdash_i \alpha$ ,

which is pronounced " $\alpha$  is derived from KB by i" or "i derives  $\alpha$  from KB."

SOUND An inference algorithm that derives only entailed sentences is called **sound** or **truth-**TRUTH-PRESERVING **preserving**. Soundness is a highly desirable property. An unsound inference procedure es-



sentially makes things up as it goes along—it announces the discovery of nonexistent needles.

We have described a reasoning process whose conclusions are guaranteed to be true in any world in which the premises are true; in particular, *if* KB *is true in the* real *world, then any sentence*  $\alpha$  *derived from KB by a sound inference procedure is also true in the real world.* So, while an inference process operates on "syntax"—internal physical configurations such as bits in registers or patterns of electrical blips in brains—the process *corresponds*

<sup>3</sup> The agent can calculate the *probability* that there is a pit in [2,2]; Chapter 13 shows how.

<sup>4</sup> Model checking works if the space of models is finite—for example, in wumpus worlds of fixed size. For arithmetic, on the other hand, the space of models is infinite: even if we restrict ourselves to the integers, there are infinitely many pairs of values for x and y in the sentence  $x + y = 4$ .

 $5$  Compare with the case of infinite search spaces in Chapter 3, where depth-first search is not complete.



to the real-world relationship whereby some aspect of the real world is the case $<sup>6</sup>$  by virtue</sup> of other aspects of the real world being the case. This correspondence between world and representation is illustrated in Figure 7.6.

GROUNDING The final issue to consider is **grounding**—the connection between logical reasoning processes and the real environment in which the agent exists. In particular, *how do we know that* KB *is true in the real world?* (After all, KB is just "syntax" inside the agent's head.) This is a philosophical question about which many, many books have been written. (See Chapter 26.) A simple answer is that the agent's sensors create the connection. For example, our wumpus-world agent has a smell sensor. The agent program creates a suitable sentence whenever there is a smell. Then, whenever that sentence is in the knowledge base, it is true in the real world. Thus, the meaning and truth of percept sentences are defined by the processes of sensing and sentence construction that produce them. What about the rest of the agent's knowledge, such as its belief that wumpuses cause smells in adjacent squares? This is not a direct representation of a single percept, but a general rule—derived, perhaps, from perceptual experience but not identical to a statement of that experience. General rules like this are produced by a sentence construction process called **learning**, which is the subject of Part V. Learning is fallible. It could be the case that wumpuses cause smells *except on February 29 in leap years*, which is when they take their baths. Thus, KB may not be true in the real world, but with good learning procedures, there is reason for optimism.

# 7.4 PROPOSITIONAL LOGIC: A VERY SIMPLE LOGIC

LOGIC

PROPOSITIONAL We now present a simple but powerful logic called **propositional logic**. We cover the syntax of propositional logic and its semantics—the way in which the truth of sentences is determined. Then we look at **entailment**—the relation between a sentence and another sentence that follows from it—and see how this leads to a simple algorithm for logical inference. Everything takes place, of course, in the wumpus world.

<sup>6</sup> As Wittgenstein (1922) put it in his famous *Tractatus*: "The world is everything that is the case."

# **7.4.1 Syntax**



SYMBOL

SENTENCES

CONNECTIVES

The **syntax** of propositional logic defines the allowable sentences. The **atomic sentences** consist of a single **proposition symbol**. Each such symbol stands for a proposition that can be true or false. We use symbols that start with an uppercase letter and may contain other letters or subscripts, for example:  $P$ ,  $Q$ ,  $R$ ,  $W_{1,3}$  and North. The names are arbitrary but are often chosen to have some mnemonic value—we use  $W_{1,3}$  to stand for the proposition that the wumpus is in [1,3]. (Remember that symbols such as  $W_{1,3}$  are *atomic*, i.e.,  $W$ , 1, and 3 are not meaningful parts of the symbol.) There are two proposition symbols with fixed meanings: True is the always-true proposition and False is the always-false proposition. **Complex sentences** are constructed from simpler sentences, using parentheses and **logical** COMPLEX **connectives**. There are five connectives in common use: LOGICAL

- NEGATION  $\Box$  (not). A sentence such as  $\neg W_{1,3}$  is called the **negation** of  $W_{1,3}$ . A **literal** is either an LITERAL atomic sentence (a **positive literal**) or a negated atomic sentence (a **negative literal**).
- ∧ (and). A sentence whose main connective is ∧, such as W1,<sup>3</sup> ∧ P3,1, is called a **con-**CONJUNCTION **junction**; its parts are the **conjuncts**. (The ∧ looks like an "A" for "And.")
- DISJUNCTION ∨ (or). A sentence using ∨, such as (W1,3∧P3,1)∨W2,2, is a **disjunction** of the **disjuncts**  $(W_{1,3} \wedge P_{3,1})$  and  $W_{2,2}$ . (Historically, the ∨ comes from the Latin "vel," which means "or." For most people, it is easier to remember  $\vee$  as an upside-down  $\wedge$ .)
- IMPLICATION  $\Rightarrow$  (implies). A sentence such as  $(W_{1,3} \land P_{3,1}) \Rightarrow \neg W_{2,2}$  is called an **implication** (or con-PREMISE ditional). Its **premise** or **antecedent** is (W1,<sup>3</sup> ∧P3,1), and its **conclusion** or **consequent** CONCLUSION is  $\neg W_{2,2}$ . Implications are also known as **rules** or **if–then** statements. The implication RULES symbol is sometimes written in other books as  $\supset$  or  $\rightarrow$ .

BICONDITIONAL  $\Leftrightarrow$  (if and only if). The sentence  $W_{1,3} \Leftrightarrow \neg W_{2,2}$  is a **biconditional**. Some other books write this as  $\equiv$ .



**Figure 7.7** A BNF (Backus–Naur Form) grammar of sentences in propositional logic, along with operator precedences, from highest to lowest.

Figure 7.7 gives a formal grammar of propositional logic; see page 1066 if you are not familiar with the BNF notation. The BNF grammar by itself is ambiguous; a sentence with several operators can be parsed by the grammar in multiple ways. To eliminate the ambiguity we define a precedence for each operator. The "not" operator  $(\neg)$  has the highest precedence, which means that in the sentence  $\neg A \land B$  the  $\neg$  binds most tightly, giving us the equivalent of  $(\neg A) \land B$  rather than  $\neg (A \land B)$ . (The notation for ordinary arithmetic is the same:  $-2+4$ is 2, not –6.) When in doubt, use parentheses to make sure of the right interpretation. Square brackets mean the same thing as parentheses; the choice of square brackets or parentheses is solely to make it easier for a human to read a sentence.

#### **7.4.2 Semantics**

Having specified the syntax of propositional logic, we now specify its semantics. The semantics defines the rules for determining the truth of a sentence with respect to a particular TRUTH VALUE model. In propositional logic, a model simply fixes the **truth value**—true or false—for every proposition symbol. For example, if the sentences in the knowledge base make use of the proposition symbols  $P_{1,2}$ ,  $P_{2,2}$ , and  $P_{3,1}$ , then one possible model is

$$
m_1 = \{P_{1,2} = false, P_{2,2} = false, P_{3,1} = true\}.
$$

With three proposition symbols, there are  $2^3 = 8$  possible models—exactly those depicted in Figure 7.5. Notice, however, that the models are purely mathematical objects with no necessary connection to wumpus worlds.  $P_{1,2}$  is just a symbol; it might mean "there is a pit in [1,2]" or "I'm in Paris today and tomorrow."

The semantics for propositional logic must specify how to compute the truth value of *any* sentence, given a model. This is done recursively. All sentences are constructed from atomic sentences and the five connectives; therefore, we need to specify how to compute the truth of atomic sentences and how to compute the truth of sentences formed with each of the five connectives. Atomic sentences are easy:

- True is true in every model and  $False$  is false in every model.
- The truth value of every other proposition symbol must be specified directly in the model. For example, in the model  $m_1$  given earlier,  $P_{1,2}$  is false.

For complex sentences, we have five rules, which hold for any subsentences  $P$  and  $Q$  in any model  $m$  (here "iff" means "if and only if"):

- $\neg P$  is true iff P is false in m.
- $P \wedge Q$  is true iff both P and Q are true in m.
- $P \vee Q$  is true iff either P or Q is true in m.
- $P \Rightarrow Q$  is true unless P is true and Q is false in m.
- $P \Leftrightarrow Q$  is true iff P and Q are both true or both false in m.

TRUTH TABLE The rules can also be expressed with **truth tables** that specify the truth value of a complex sentence for each possible assignment of truth values to its components. Truth tables for the five connectives are given in Figure 7.8. From these tables, the truth value of any sentence s can be computed with respect to any model  $m$  by a simple recursive evaluation. For example,

	$\mathcal{Q}$	$\neg P$	$P \wedge Q$	$P \vee Q$	$\Rightarrow$	⇔
false	false	true	false	false	true	true
false	true	true	false	true	true	false
true	false	false	false	true	false	false
true	true	false	true	<i>true</i>	true	true

**Figure 7.8** Truth tables for the five logical connectives. To use the table to compute, for example, the value of  $P \vee Q$  when P is true and Q is false, first look on the left for the row where P is true and Q is false (the third row). Then look in that row under the  $P \vee Q$  column to see the result: true.

the sentence  $\neg P_{1,2} \land (P_{2,2} \lor P_{3,1})$ , evaluated in  $m_1$ , gives true  $\land$  (false  $\lor$  true) = true  $\land$  $true = true$ . Exercise 7.3 asks you to write the algorithm PL-TRUE?(s, m), which computes the truth value of a propositional logic sentence  $s$  in a model  $m$ .

The truth tables for "and," "or," and "not" are in close accord with our intuitions about the English words. The main point of possible confusion is that  $P \vee Q$  is true when P is true or Q is true *or both*. A different connective, called "exclusive or" ("xor" for short), yields false when both disjuncts are true.<sup>7</sup> There is no consensus on the symbol for exclusive or; some choices are  $\dot{\vee}$  or  $\neq$  or  $\oplus$ .

The truth table for  $\Rightarrow$  may not quite fit one's intuitive understanding of "P implies Q" or "if P then Q." For one thing, propositional logic does not require any relation of *causation* or *relevance* between P and Q. The sentence "5 is odd implies Tokyo is the capital of Japan" is a true sentence of propositional logic (under the normal interpretation), even though it is a decidedly odd sentence of English. Another point of confusion is that any implication is true whenever its antecedent is false. For example, "5 is even implies Sam is smart" is true, regardless of whether Sam is smart. This seems bizarre, but it makes sense if you think of " $P \Rightarrow Q$ " as saying, "If P is true, then I am claiming that Q is true. Otherwise I am making no claim." The only way for this sentence to be *false* is if P is true but Q is false.

The biconditional,  $P \Leftrightarrow Q$ , is true whenever both  $P \Rightarrow Q$  and  $Q \Rightarrow P$  are true. In English, this is often written as "P if and only if  $Q$ ." Many of the rules of the wumpus world are best written using ⇔. For example, a square is breezy *if* a neighboring square has a pit, and a square is breezy *only if* a neighboring square has a pit. So we need a biconditional,

 $B_{1,1} \Leftrightarrow (P_{1,2} \vee P_{2,1}),$ 

where  $B_{1,1}$  means that there is a breeze in [1,1].

# **7.4.3 A simple knowledge base**

Now that we have defined the semantics for propositional logic, we can construct a knowledge base for the wumpus world. We focus first on the *immutable* aspects of the wumpus world, leaving the mutable aspects for a later section. For now, we need the following symbols for each  $[x, y]$  location:

<sup>7</sup> Latin has a separate word, *aut*, for exclusive or.

 $P_{x,y}$  is true if there is a pit in [x, y].

 $W_{x,y}$  is true if there is a wumpus in  $[x, y]$ , dead or alive.

 $B_{x,y}$  is true if the agent perceives a breeze in [x, y].

 $S_{x,y}$  is true if the agent perceives a stench in  $[x, y]$ .

The sentences we write will suffice to derive  $\neg P_{1,2}$  (there is no pit in [1,2]), as was done informally in Section 7.3. We label each sentence  $R_i$  so that we can refer to them:

• There is no pit in  $[1,1]$ :

 $R_1: \neg P_{1,1}$ .

• A square is breezy if and only if there is a pit in a neighboring square. This has to be stated for each square; for now, we include just the relevant squares:

 $R_2: B_{1,1} \Leftrightarrow (P_{1,2} \vee P_{2,1}).$  $R_3: B_{2,1} \Leftrightarrow (P_{1,1} \vee P_{2,2} \vee P_{3,1}).$ 

• The preceding sentences are true in all wumpus worlds. Now we include the breeze percepts for the first two squares visited in the specific world the agent is in, leading up to the situation in Figure 7.3(b).

 $R_4: \neg B_{1,1}.$  $R_5: B_{2,1}.$ 

#### **7.4.4 A simple inference procedure**

Our goal now is to decide whether  $KB \models \alpha$  for some sentence  $\alpha$ . For example, is  $\neg P_{1,2}$ entailed by our KB? Our first algorithm for inference is a model-checking approach that is a direct implementation of the definition of entailment: enumerate the models, and check that  $\alpha$  is true in every model in which KB is true. Models are assignments of true or false to every proposition symbol. Returning to our wumpus-world example, the relevant proposition symbols are  $B_{1,1}$ ,  $B_{2,1}$ ,  $P_{1,1}$ ,  $P_{1,2}$ ,  $P_{2,1}$ ,  $P_{2,2}$ , and  $P_{3,1}$ . With seven symbols, there are  $2^7 = 128$  possible models; in three of these, KB is true (Figure 7.9). In those three models,  $\neg P_{1,2}$  is true, hence there is no pit in [1,2]. On the other hand,  $P_{2,2}$  is true in two of the three models and false in one, so we cannot yet tell whether there is a pit in [2,2].

Figure 7.9 reproduces in a more precise form the reasoning illustrated in Figure 7.5. A general algorithm for deciding entailment in propositional logic is shown in Figure 7.10. Like the BACKTRACKING-SEARCH algorithm on page 215, TT-ENTAILS? performs a recursive enumeration of a finite space of assignments to symbols. The algorithm is **sound** because it implements directly the definition of entailment, and **complete** because it works for any KB and  $\alpha$  and always terminates—there are only finitely many models to examine.

Of course, "finitely many" is not always the same as "few." If  $KB$  and  $\alpha$  contain n symbols in all, then there are  $2^n$  models. Thus, the time complexity of the algorithm is  $O(2^n)$ . (The space complexity is only  $O(n)$  because the enumeration is depth-first.) Later in this chapter we show algorithms that are much more efficient in many cases. Unfortunately, propositional entailment is co-NP-complete (i.e., probably no easier than NP-complete—see Appendix A), so *every known inference algorithm for propositional logic has a worst-case complexity that is exponential in the size of the input.*



**Figure 7.9** A truth table constructed for the knowledge base given in the text. *KB* is true if  $R_1$  through  $R_5$  are true, which occurs in just 3 of the 128 rows (the ones underlined in the right-hand column). In all 3 rows,  $P_{1,2}$  is false, so there is no pit in [1,2]. On the other hand, there might (or might not) be a pit in [2,2].

```
function TT-ENTAILS?(KB, \alpha) returns true or false
inputs: KB, the knowledge base, a sentence in propositional logic
        \alpha, the query, a sentence in propositional logic
 symbols \leftarrow a list of the proposition symbols in KB and \alphareturn TT-CHECK-ALL(KB, \alpha, symbols, \{\})function TT-CHECK-ALL(KB,α, symbols , model) returns true or false
if EMPTY?(symbols) then
    if PL-TRUE?(KB, model) then return PL-TRUE?(\alpha, model)
    else return true // when KB is false, always return true
else do
    P \leftarrow FIRST(symbols)
    rest \leftarrowREST(symbols)
    return (TT-CHECK-ALL(KB, \alpha, rest, model \cup {P = true})
            and
            TT-CHECK-ALL(KB, \alpha, rest, model \cup {P = false}))
```
**Figure 7.10** A truth-table enumeration algorithm for deciding propositional entailment. (TT stands for truth table.) PL-TRUE? returns *true* if a sentence holds within a model. The variable *model* represents a partial model—an assignment to some of the symbols. The keyword "**and**" is used here as a logical operation on its two arguments, returning true or false.

 $(\alpha \wedge \beta) \equiv (\beta \wedge \alpha)$  commutativity of  $\wedge$  $(\alpha \vee \beta) \equiv (\beta \vee \alpha)$  commutativity of ∨  $((\alpha \wedge \beta) \wedge \gamma) \equiv (\alpha \wedge (\beta \wedge \gamma))$  associativity of  $\wedge$  $((\alpha \vee \beta) \vee \gamma) \equiv (\alpha \vee (\beta \vee \gamma))$  associativity of  $\vee$  $\neg(\neg \alpha) \equiv \alpha$  double-negation elimination  $(\alpha \Rightarrow \beta) \equiv (\neg \beta \Rightarrow \neg \alpha)$  contraposition  $(\alpha \Rightarrow \beta) \equiv (\neg \alpha \vee \beta)$  implication elimination  $(\alpha \Leftrightarrow \beta) \equiv ((\alpha \Rightarrow \beta) \wedge (\beta \Rightarrow \alpha))$  biconditional elimination  $\neg(\alpha \land \beta) \equiv (\neg \alpha \lor \neg \beta)$  De Morgan  $\neg(\alpha \lor \beta) \equiv (\neg \alpha \land \neg \beta)$  De Morgan  $(\alpha \wedge (\beta \vee \gamma)) \equiv ((\alpha \wedge \beta) \vee (\alpha \wedge \gamma))$  distributivity of  $\wedge$  over  $\vee$  $(\alpha \vee (\beta \wedge \gamma)) \equiv ((\alpha \vee \beta) \wedge (\alpha \vee \gamma))$  distributivity of  $\vee$  over  $\wedge$ 

**Figure 7.11** Standard logical equivalences. The symbols  $\alpha$ ,  $\beta$ , and  $\gamma$  stand for arbitrary sentences of propositional logic.

# 7.5 PROPOSITIONAL THEOREM PROVING

So far, we have shown how to determine entailment by *model checking*: enumerating models and showing that the sentence must hold in all models. In this section, we show how entail-THEOREM PROVING ment can be done by **theorem proving**—applying rules of inference directly to the sentences in our knowledge base to construct a proof of the desired sentence without consulting models. If the number of models is large but the length of the proof is short, then theorem proving can be more efficient than model checking.

**EQUIVALENCE** 



Before we plunge into the details of theorem-proving algorithms, we will need some additional concepts related to entailment. The first concept is **logical equivalence**: two sen- LOGICAL tences  $\alpha$  and  $\beta$  are logically equivalent if they are true in the same set of models. We write this as  $\alpha \equiv \beta$ . For example, we can easily show (using truth tables) that  $P \wedge Q$  and  $Q \wedge P$ are logically equivalent; other equivalences are shown in Figure 7.11. These equivalences play much the same role in logic as arithmetic identities do in ordinary mathematics. An alternative definition of equivalence is as follows: any two sentences  $\alpha$  and  $\beta$  are equivalent only if each of them entails the other:

 $\alpha \equiv \beta$  if and only if  $\alpha \models \beta$  and  $\beta \models \alpha$ .

VALIDITY The second concept we will need is **validity**. A sentence is valid if it is true in *all* models. For TAUTOLOGY example, the sentence  $P \vee \neg P$  is valid. Valid sentences are also known as **tautologies**—they are *necessarily* true. Because the sentence True is true in all models, every valid sentence is logically equivalent to True. What good are valid sentences? From our definition of entailment, we can derive the **deduction theorem**, which was known to the ancient Greeks:

*For any sentences*  $\alpha$  *and*  $\beta$ ,  $\alpha \models \beta$  *if and only if the sentence*  $(\alpha \Rightarrow \beta)$  *is valid.* 

(Exercise 7.5 asks for a proof.) Hence, we can decide if  $\alpha \models \beta$  by checking that  $(\alpha \Rightarrow \beta)$  is true in every model—which is essentially what the inference algorithm in Figure 7.10 doesor by proving that  $(\alpha \Rightarrow \beta)$  is equivalent to *True.* Conversely, the deduction theorem states that every valid implication sentence describes a legitimate inference.

SATISFIABILITY The final concept we will need is **satisfiability**. A sentence is satisfiable if it is true in, or satisfied by, *some* model. For example, the knowledge base given earlier,  $(R_1 \wedge R_2 \wedge R_3)$  $R_3 \wedge R_4 \wedge R_5$ , is satisfiable because there are three models in which it is true, as shown in Figure 7.9. Satisfiability can be checked by enumerating the possible models until one is found that satisfies the sentence. The problem of determining the satisfiability of sentences SAT in propositional logic—the **SAT** problem—was the first problem proved to be NP-complete. Many problems in computer science are really satisfiability problems. For example, all the constraint satisfaction problems in Chapter 6 ask whether the constraints are satisfiable by some assignment.

> Validity and satisfiability are of course connected:  $\alpha$  is valid iff  $\neg \alpha$  is unsatisfiable; contrapositively,  $\alpha$  is satisfiable iff  $\neg \alpha$  is not valid. We also have the following useful result:

 $\alpha \models \beta$  *if and only if the sentence*  $(\alpha \land \neg \beta)$  *is unsatisfiable.* 

Proving  $\beta$  from  $\alpha$  by checking the unsatisfiability of  $(\alpha \land \neg \beta)$  corresponds exactly to the **REDUCTION** standard mathematical proof technique of *reductio ad absurdum* (literally, "reduction to an REDUCTION REFUTATION absurd thing"). It is also called proof by **refutation** or proof by **contradiction**. One assumes a CONTRADICTION sentence  $\beta$  to be false and shows that this leads to a contradiction with known axioms  $\alpha$ . This contradiction is exactly what is meant by saying that the sentence  $(\alpha \land \neg \beta)$  is unsatisfiable.

#### **7.5.1 Inference and proofs**

INFERENCE RULES This section covers **inference rules** that can be applied to derive a **proof**—a chain of conclu-PROOF sions that leads to the desired goal. The best-known rule is called **Modus Ponens** (Latin for MODUS PONENS *mode that affirms*) and is written

$$
\frac{\alpha \Rightarrow \beta, \quad \alpha}{\beta}
$$

.

The notation means that, whenever any sentences of the form  $\alpha \Rightarrow \beta$  and  $\alpha$  are given, then the sentence  $\beta$  can be inferred. For example, if (WumpusAhead ∧WumpusAlive)  $\Rightarrow$  Shoot and (WumpusAhead  $\land$  WumpusAlive) are given, then Shoot can be inferred.

AND-ELIMINATION Another useful inference rule is **And-Elimination**, which says that, from a conjunction, any of the conjuncts can be inferred:

> $\alpha \wedge \beta$  $\frac{\wedge \nu}{\alpha}$  .

For example, from (WumpusAhead  $\land$  WumpusAlive), WumpusAlive can be inferred.

By considering the possible truth values of  $\alpha$  and  $\beta$ , one can show easily that Modus Ponens and And-Elimination are sound once and for all. These rules can then be used in any particular instances where they apply, generating sound inferences without the need for enumerating models.

All of the logical equivalences in Figure 7.11 can be used as inference rules. For example, the equivalence for biconditional elimination yields the two inference rules

 $\alpha \Leftrightarrow \beta$  $(\alpha \Rightarrow \beta) \wedge (\beta \Rightarrow \alpha)$ and  $\frac{(\alpha \Rightarrow \beta) \land (\beta \Rightarrow \alpha)}{\alpha \Leftrightarrow \beta}$ .



ABSURDUM

Not all inference rules work in both directions like this. For example, we cannot run Modus Ponens in the opposite direction to obtain  $\alpha \Rightarrow \beta$  and  $\alpha$  from  $\beta$ .

Let us see how these inference rules and equivalences can be used in the wumpus world. We start with the knowledge base containing  $R_1$  through  $R_5$  and show how to prove  $\neg P_{1,2}$ , that is, there is no pit in [1,2]. First, we apply biconditional elimination to  $R_2$  to obtain

 $R_6: (B_{1,1} \Rightarrow (P_{1,2} \vee P_{2,1})) \wedge ((P_{1,2} \vee P_{2,1}) \Rightarrow B_{1,1}).$ 

Then we apply And-Elimination to  $R_6$  to obtain

 $R_7: ((P_{1,2} \vee P_{2,1}) \Rightarrow B_{1,1})$ .

Logical equivalence for contrapositives gives

 $R_8: (\neg B_{1,1} \Rightarrow \neg (P_{1,2} \vee P_{2,1}))$ .

Now we can apply Modus Ponens with  $R_8$  and the percept  $R_4$  (i.e.,  $\neg B_{1,1}$ ), to obtain

 $R_9: \neg (P_{1,2} \vee P_{2,1})$ .

Finally, we apply De Morgan's rule, giving the conclusion

 $R_{10} : \neg P_{1,2} \wedge \neg P_{2,1}$ .

That is, neither [1,2] nor [2,1] contains a pit.

We found this proof by hand, but we can apply any of the search algorithms in Chapter 3 to find a sequence of steps that constitutes a proof. We just need to define a proof problem as follows:

- INITIAL STATE: the initial knowledge base.
- ACTIONS: the set of actions consists of all the inference rules applied to all the sentences that match the top half of the inference rule.
- RESULT: the result of an action is to add the sentence in the bottom half of the inference rule.
- GOAL: the goal is a state that contains the sentence we are trying to prove.

Thus, searching for proofs is an alternative to enumerating models. In many practical cases *finding a proof can be more efficient because the proof can ignore irrelevant propositions, no matter how many of them there are.* For example, the proof given earlier leading to  $\neg P_{1,2} \land$  $\neg P_{2,1}$  does not mention the propositions  $B_{2,1}$ ,  $P_{1,1}$ ,  $P_{2,2}$ , or  $P_{3,1}$ . They can be ignored because the goal proposition,  $P_{1,2}$ , appears only in sentence  $R_2$ ; the other propositions in  $R_2$ appear only in  $R_4$  and  $R_2$ ; so  $R_1$ ,  $R_3$ , and  $R_5$  have no bearing on the proof. The same would hold even if we added a million more sentences to the knowledge base; the simple truth-table algorithm, on the other hand, would be overwhelmed by the exponential explosion of models.

MONOTONICITY One final property of logical systems is **monotonicity**, which says that the set of entailed sentences can only *increase* as information is added to the knowledge base.<sup>8</sup> For any sentences  $\alpha$  and  $\beta$ ,

if  $KB \models \alpha$  then  $KB \land \beta \models \alpha$ .

<sup>&</sup>lt;sup>8</sup> **Nonmonotonic** logics, which violate the monotonicity property, capture a common property of human reasoning: changing one's mind. They are discussed in Section 12.6.

For example, suppose the knowledge base contains the additional assertion  $\beta$  stating that there are exactly eight pits in the world. This knowledge might help the agent draw *additional* conclusions, but it cannot invalidate any conclusion  $\alpha$  already inferred—such as the conclusion that there is no pit in [1,2]. Monotonicity means that inference rules can be applied whenever suitable premises are found in the knowledge base—the conclusion of the rule must follow *regardless of what else is in the knowledge base*.

# **7.5.2 Proof by resolution**

We have argued that the inference rules covered so far are *sound*, but we have not discussed the question of *completeness* for the inference algorithms that use them. Search algorithms such as iterative deepening search (page 89) are complete in the sense that they will find any reachable goal, but if the available inference rules are inadequate, then the goal is not reachable—no proof exists that uses only those inference rules. For example, if we removed the biconditional elimination rule, the proof in the preceding section would not go through. The current section introduces a single inference rule, **resolution**, that yields a complete inference algorithm when coupled with any complete search algorithm.

We begin by using a simple version of the resolution rule in the wumpus world. Let us consider the steps leading up to Figure 7.4(a): the agent returns from [2,1] to [1,1] and then goes to [1,2], where it perceives a stench, but no breeze. We add the following facts to the knowledge base:

$$
R_{11} : \neg B_{1,2} .
$$
  
\n
$$
R_{12} : B_{1,2} \Leftrightarrow (P_{1,1} \vee P_{2,2} \vee P_{1,3}).
$$

By the same process that led to  $R_{10}$  earlier, we can now derive the absence of pits in [2,2] and  $[1,3]$  (remember that  $[1,1]$  is already known to be pitless):

$$
R_{13} : \neg P_{2,2} .
$$
  

$$
R_{14} : \neg P_{1,3} .
$$

We can also apply biconditional elimination to  $R_3$ , followed by Modus Ponens with  $R_5$ , to obtain the fact that there is a pit in  $[1,1]$ ,  $[2,2]$ , or  $[3,1]$ :

$$
R_{15}:\quad P_{1,1}\vee P_{2,2}\vee P_{3,1} .
$$

Now comes the first application of the resolution rule: the literal  $\neg P_{2,2}$  in  $R_{13}$  *resolves with* RESOLVENT the literal  $P_{2,2}$  in  $R_{15}$  to give the **resolvent** 

$$
R_{16}:\quad P_{1,1}\vee P_{3,1} .
$$

In English; if there's a pit in one of  $[1,1]$ ,  $[2,2]$ , and  $[3,1]$  and it's not in  $[2,2]$ , then it's in  $[1,1]$ or [3,1]. Similarly, the literal  $\neg P_{1,1}$  in  $R_1$  resolves with the literal  $P_{1,1}$  in  $R_{16}$  to give

$$
R_{17}:\quad P_{3,1} \ .
$$

In English: if there's a pit in  $[1,1]$  or  $[3,1]$  and it's not in  $[1,1]$ , then it's in  $[3,1]$ . These last UNIT RESOLUTION two inference steps are examples of the **unit resolution** inference rule,

$$
\frac{\ell_1 \vee \dots \vee \ell_k, \quad m}{\ell_1 \vee \dots \vee \ell_{i-1} \vee \ell_{i+1} \vee \dots \vee \ell_k}
$$

LITERALS

,

CLAUSE of the other). Thus, the unit resolution rule takes a **clause**—a disjunction of literals—and a literal and produces a new clause. Note that a single literal can be viewed as a disjunction of UNIT CLAUSE one literal, also known as a **unit clause**.

RESOLUTION The unit resolution rule can be generalized to the full **resolution** rule,

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

where  $\ell_i$  and  $m_j$  are complementary literals. This says that resolution takes two clauses and produces a new clause containing all the literals of the two original clauses *except* the two complementary literals. For example, we have

$$
\frac{P_{1,1} \vee P_{3,1}, \quad \neg P_{1,1} \vee \neg P_{2,2}}{P_{3,1} \vee \neg P_{2,2}}.
$$

There is one more technical aspect of the resolution rule: the resulting clause should contain FACTORING only one copy of each literal.<sup>9</sup> The removal of multiple copies of literals is called **factoring**. For example, if we resolve  $(A \vee B)$  with  $(A \vee \neg B)$ , we obtain  $(A \vee A)$ , which is reduced to just A.

> The *soundness* of the resolution rule can be seen easily by considering the literal  $l_i$  that is complementary to literal  $m_j$  in the other clause. If  $\ell_i$  is true, then  $m_j$  is false, and hence  $m_1 \vee \cdots \vee m_{j-1} \vee m_{j+1} \vee \cdots \vee m_n$  must be true, because  $m_1 \vee \cdots \vee m_n$  is given. If  $\ell_i$  is false, then  $\ell_1 \vee \cdots \vee \ell_{i-1} \vee \ell_{i+1} \vee \cdots \vee \ell_k$  must be true because  $\ell_1 \vee \cdots \vee \ell_k$  is given. Now  $\ell_i$  is either true or false, so one or other of these conclusions holds—exactly as the resolution rule states.

> What is more surprising about the resolution rule is that it forms the basis for a family of *complete* inference procedures. *A resolution-based theorem prover can, for any sentences*  $\alpha$  *and*  $\beta$  *in propositional logic, decide whether*  $\alpha \models \beta$ . The next two subsections explain how resolution accomplishes this.

#### **Conjunctive normal form**



The resolution rule applies only to clauses (that is, disjunctions of literals), so it would seem to be relevant only to knowledge bases and queries consisting of clauses. How, then, can it lead to a complete inference procedure for all of propositional logic? The answer is that *every sentence of propositional logic is logically equivalent to a conjunction of clauses.* A sentence expressed as a conjunction of clauses is said to be in **conjunctive normal form** or **CNF** (see Figure 7.14). We now describe a procedure for converting to CNF. We illustrate the procedure by converting the sentence  $B_{1,1} \Leftrightarrow (P_{1,2} \vee P_{2,1})$  into CNF. The steps are as follows:

1. Eliminate  $\Leftrightarrow$ , replacing  $\alpha \Leftrightarrow \beta$  with  $(\alpha \Rightarrow \beta) \wedge (\beta \Rightarrow \alpha)$ .

$$
(B_{1,1} \Rightarrow (P_{1,2} \vee P_{2,1})) \wedge ((P_{1,2} \vee P_{2,1}) \Rightarrow B_{1,1}).
$$

2. Eliminate  $\Rightarrow$ , replacing  $\alpha \Rightarrow \beta$  with  $\neg \alpha \lor \beta$ :

$$
(\neg B_{1,1} \vee P_{1,2} \vee P_{2,1}) \wedge (\neg (P_{1,2} \vee P_{2,1}) \vee B_{1,1}) \ .
$$

,

<sup>&</sup>lt;sup>9</sup> If a clause is viewed as a *set* of literals, then this restriction is automatically respected. Using set notation for clauses makes the resolution rule much cleaner, at the cost of introducing additional notation.

3. CNF requires  $\neg$  to appear only in literals, so we "move  $\neg$  inwards" by repeated application of the following equivalences from Figure 7.11:

$$
\neg(\neg \alpha) \equiv \alpha \text{ (double-negation elimination)}
$$
  

$$
\neg(\alpha \land \beta) \equiv (\neg \alpha \lor \neg \beta) \text{ (De Morgan)}
$$
  

$$
\neg(\alpha \lor \beta) \equiv (\neg \alpha \land \neg \beta) \text{ (De Morgan)}
$$

In the example, we require just one application of the last rule:

 $(\neg B_{1,1} \lor P_{1,2} \lor P_{2,1}) \land ((\neg P_{1,2} \land \neg P_{2,1}) \lor B_{1,1}).$ 

4. Now we have a sentence containing nested  $\wedge$  and  $\vee$  operators applied to literals. We apply the distributivity law from Figure 7.11, distributing  $\vee$  over  $\wedge$  wherever possible.

 $(\neg B_{1,1} \lor P_{1,2} \lor P_{2,1}) \land (\neg P_{1,2} \lor B_{1,1}) \land (\neg P_{2,1} \lor B_{1,1})$ .

The original sentence is now in CNF, as a conjunction of three clauses. It is much harder to read, but it can be used as input to a resolution procedure.

#### **A resolution algorithm**

Inference procedures based on resolution work by using the principle of proof by contradiction introduced on page 250. That is, to show that  $KB \models \alpha$ , we show that  $(KB \land \neg \alpha)$  is unsatisfiable. We do this by proving a contradiction.

A resolution algorithm is shown in Figure 7.12. First,  $(KB \wedge \neg \alpha)$  is converted into CNF. Then, the resolution rule is applied to the resulting clauses. Each pair that contains complementary literals is resolved to produce a new clause, which is added to the set if it is not already present. The process continues until one of two things happens:

- there are no new clauses that can be added, in which case KB does not entail  $\alpha$ ; or,
- two clauses resolve to yield the *empty* clause, in which case  $KB$  entails  $\alpha$ .

The empty clause—a disjunction of no disjuncts—is equivalent to False because a disjunction is true only if at least one of its disjuncts is true. Another way to see that an empty clause represents a contradiction is to observe that it arises only from resolving two complementary unit clauses such as P and  $\neg P$ .

We can apply the resolution procedure to a very simple inference in the wumpus world. When the agent is in [1,1], there is no breeze, so there can be no pits in neighboring squares. The relevant knowledge base is

$$
KB = R_2 \land R_4 = (B_{1,1} \Leftrightarrow (P_{1,2} \lor P_{2,1})) \land \neg B_{1,1}
$$

and we wish to prove  $\alpha$  which is, say,  $\neg P_{1,2}$ . When we convert  $(KB \wedge \neg \alpha)$  into CNF, we obtain the clauses shown at the top of Figure 7.13. The second row of the figure shows clauses obtained by resolving pairs in the first row. Then, when  $P_{1,2}$  is resolved with  $\neg P_{1,2}$ , we obtain the empty clause, shown as a small square. Inspection of Figure 7.13 reveals that many resolution steps are pointless. For example, the clause  $B_{1,1} \vee \neg B_{1,1} \vee P_{1,2}$  is equivalent to  $True \vee P_{1,2}$  which is equivalent to  $True$ . Deducing that  $True$  is true is not very helpful. Therefore, any clause in which two complementary literals appear can be discarded.

**function** PL-RESOLUTION(KB,α) **returns** true or false **inputs**: KB, the knowledge base, a sentence in propositional logic  $\alpha$ , the query, a sentence in propositional logic clauses  $\leftarrow$  the set of clauses in the CNF representation of  $KB \wedge \neg \alpha$  $new \leftarrow \{\}$ **loop do for each** pair of clauses C<sup>i</sup> , C<sup>j</sup> **in** clauses **do**  $resolvents \leftarrow \text{PL-RESOLVE}(C_i, C_j)$ **if** resolvents contains the empty clause **then return** true  $new \leftarrow new \cup \ resolvents$ **if** new ⊆ clauses **then return** false  $clauses \leftarrow clauses \cup new$ 

**Figure 7.12** A simple resolution algorithm for propositional logic. The function PL-RESOLVE returns the set of all possible clauses obtained by resolving its two inputs.



world.  $\neg P_{1,2}$  is shown to follow from the first four clauses in the top row.

#### **Completeness of resolution**

CLOSURE

To conclude our discussion of resolution, we now show why PL-RESOLUTION is complete. RESOLUTION To do this, we introduce the **resolution closure**  $RC(S)$  of a set of clauses S, which is the set of all clauses derivable by repeated application of the resolution rule to clauses in  $S$  or their derivatives. The resolution closure is what PL-RESOLUTION computes as the final value of the variable *clauses*. It is easy to see that  $RC(S)$  must be finite, because there are only finitely many distinct clauses that can be constructed out of the symbols  $P_1, \ldots, P_k$  that appear in S. (Notice that this would not be true without the factoring step that removes multiple copies of literals.) Hence, PL-RESOLUTION always terminates.

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

If a set of clauses is unsatisfiable, then the resolution closure of those clauses contains the empty clause.

This theorem is proved by demonstrating its contrapositive: if the closure RC(S) does *not*

GROUND **RESOLUTION** THEOREM

contain the empty clause, then  $S$  is satisfiable. In fact, we can construct a model for  $S$  with suitable truth values for  $P_1, \ldots, P_k$ . The construction procedure is as follows:

For  $i$  from 1 to  $k$ ,

- If a clause in  $RC(S)$  contains the literal  $\neg P_i$  and all its other literals are false under the assignment chosen for  $P_1, \ldots, P_{i-1}$ , then assign false to  $P_i$ .
- $-$  Otherwise, assign *true* to  $P_i$ .

This assignment to  $P_1, \ldots, P_k$  is a model of S. To see this, assume the opposite—that, at some stage i in the sequence, assigning symbol  $P_i$  causes some clause C to become false. For this to happen, it must be the case that all the *other* literals in C must already have been falsified by assignments to  $P_1, \ldots, P_{i-1}$ . Thus, C must now look like either (false  $\vee$  false  $\vee$  $\cdots false \vee P_i$ ) or like  $(false \vee false \vee \cdots false \vee \neg P_i)$ . If just one of these two is in  $RC(S)$ , then the algorithm will assign the appropriate truth value to  $P_i$  to make C true, so C can only be falsified if *both* of these clauses are in  $RC(S)$ . Now, since  $RC(S)$  is closed under resolution, it will contain the resolvent of these two clauses, and that resolvent will have all of its literals already falsified by the assignments to  $P_1, \ldots, P_{i-1}$ . This contradicts our assumption that the first falsified clause appears at stage  $i$ . Hence, we have proved that the construction never falsifies a clause in  $RC(S)$ ; that is, it produces a model of  $RC(S)$  and thus a model of S itself (since S is contained in  $RC(S)$ ).

#### **7.5.3 Horn clauses and definite clauses**

The completeness of resolution makes it a very important inference method. In many practical situations, however, the full power of resolution is not needed. Some real-world knowledge bases satisfy certain restrictions on the form of sentences they contain, which enables them to use a more restricted and efficient inference algorithm.

DEFINITE CLAUSE One such restricted form is the **definite clause**, which is a disjunction of literals of which *exactly one is positive*. For example, the clause  $(\neg L_{1,1} \lor \neg Breeze \lor B_{1,1})$  is a definite clause, whereas  $(\neg B_{1,1} \lor P_{1,2} \lor P_{2,1})$  is not.

HORN CLAUSE Slightly more general is the **Horn clause**, which is a disjunction of literals of which *at most one is positive*. So all definite clauses are Horn clauses, as are clauses with no positive GOAL CLAUSES literals; these are called **goal clauses**. Horn clauses are closed under resolution: if you resolve two Horn clauses, you get back a Horn clause.

Knowledge bases containing only definite clauses are interesting for three reasons:

1. Every definite clause can be written as an implication whose premise is a conjunction of positive literals and whose conclusion is a single positive literal. (See Exercise 7.13.) For example, the definite clause  $(\neg L_{1,1} \lor \neg Breeze \lor B_{1,1})$  can be written as the implication  $(L_{1,1} \wedge Breeze) \Rightarrow B_{1,1}$ . In the implication form, the sentence is easier to understand: it says that if the agent is in [1,1] and there is a breeze, then [1,1] is breezy. BODY In Horn form, the premise is called the **body** and the conclusion is called the **head**. A HEAD sentence consisting of a single positive literal, such as  $L_{1,1}$ , is called a **fact**. It too can FACT be written in implication form as  $True \Rightarrow L_{1,1}$ , but it is simpler to write just  $L_{1,1}$ .



**Figure 7.14** A grammar for conjunctive normal form, Horn clauses, and definite clauses. A clause such as  $A \wedge B \Rightarrow C$  is still a definite clause when it is written as  $\neg A \vee \neg B \vee C$ , but only the former is considered the canonical form for definite clauses. One more class is the *k*-CNF sentence, which is a CNF sentence where each clause has at most *k* literals.

BACKWARD-<br>CHAINING

- FORWARD-CHAINING 2. Inference with Horn clauses can be done through the **forward-chaining** and **backwardchaining** algorithms, which we explain next. Both of these algorithms are natural, in that the inference steps are obvious and easy for humans to follow. This type of inference is the basis for **logic programming**, which is discussed in Chapter 9.
	- 3. Deciding entailment with Horn clauses can be done in time that is *linear* in the size of the knowledge base—a pleasant surprise.

# **7.5.4 Forward and backward chaining**

The forward-chaining algorithm PL-FC-ENTAILS?  $(KB, q)$  determines if a single proposition symbol  $q$ —the query—is entailed by a knowledge base of definite clauses. It begins from known facts (positive literals) in the knowledge base. If all the premises of an implication are known, then its conclusion is added to the set of known facts. For example, if  $L_{1,1}$ and Breeze are known and  $(L_{1,1} \wedge Breeze) \Rightarrow B_{1,1}$  is in the knowledge base, then  $B_{1,1}$  can be added. This process continues until the query  $q$  is added or until no further inferences can be made. The detailed algorithm is shown in Figure 7.15; the main point to remember is that it runs in linear time.

The best way to understand the algorithm is through an example and a picture. Figure 7.16(a) shows a simple knowledge base of Horn clauses with  $A$  and  $B$  as known facts. Figure 7.16(b) shows the same knowledge base drawn as an **AND–OR graph** (see Chapter 4). In AND–OR graphs, multiple links joined by an arc indicate a conjunction—every link must be proved—while multiple links without an arc indicate a disjunction—any link can be proved. It is easy to see how forward chaining works in the graph. The known leaves (here,  $A$  and  $B$ ) are set, and inference propagates up the graph as far as possible. Wherever a conjunction appears, the propagation waits until all the conjuncts are known before proceeding. The reader is encouraged to work through the example in detail.

```
function PL-FC-ENTAILS?(KB, q) returns true or false
inputs: KB, the knowledge base, a set of propositional definite clauses
         q, the query, a proposition symbol
count \leftarrow a table, where count[c] is the number of symbols in c's premise
inferred \leftarrow a table, where inferred[s] is initially false for all symbols
a\ddot{g} \leftarrow a queue of symbols, initially symbols known to be true in KB
while agenda is not empty do
    p \leftarrow POP(aqenda)if p = q then return true
    if inferred[p] = false then
        \text{inferced}[p] \leftarrow \text{true}for each clause c in KB where p is in c.PREMISE do
            decrement count[c]if count[c] = 0 then add c. CONCLUSION to agenda
return false
```
**Figure 7.15** The forward-chaining algorithm for propositional logic. The *agenda* keeps track of symbols known to be true but not yet "processed." The count table keeps track of how many premises of each implication are as yet unknown. Whenever a new symbol  $p$  from the agenda is processed, the count is reduced by one for each implication in whose premise p appears (easily identified in constant time with appropriate indexing.) If a count reaches zero, all the premises of the implication are known, so its conclusion can be added to the agenda. Finally, we need to keep track of which symbols have been processed; a symbol that is already in the set of inferred symbols need not be added to the agenda again. This avoids redundant work and prevents loops caused by implications such as  $P \Rightarrow Q$  and  $Q \Rightarrow P$ .

It is easy to see that forward chaining is **sound**: every inference is essentially an appli-



cation of Modus Ponens. Forward chaining is also **complete**: every entailed atomic sentence will be derived. The easiest way to see this is to consider the final state of the *inferred* table FIXED POINT (after the algorithm reaches a **fixed point** where no new inferences are possible). The table contains true for each symbol inferred during the process, and false for all other symbols. We can view the table as a logical model; moreover, *every definite clause in the original KB is true in this model.* To see this, assume the opposite, namely that some clause  $a_1 \wedge \ldots \wedge a_k \Rightarrow b$ is false in the model. Then  $a_1 \wedge \ldots \wedge a_k$  must be true in the model and b must be false in the model. But this contradicts our assumption that the algorithm has reached a fixed point! We can conclude, therefore, that the set of atomic sentences inferred at the fixed point defines a model of the original KB. Furthermore, any atomic sentence  $q$  that is entailed by the KB must be true in all its models and in this model in particular. Hence, every entailed atomic sentence  $q$  must be inferred by the algorithm.

DATA-DRIVEN Forward chaining is an example of the general concept of **data-driven** reasoning—that is, reasoning in which the focus of attention starts with the known data. It can be used within an agent to derive conclusions from incoming percepts, often without a specific query in mind. For example, the wumpus agent might TELL its percepts to the knowledge base using



an incremental forward-chaining algorithm in which new facts can be added to the agenda to initiate new inferences. In humans, a certain amount of data-driven reasoning occurs as new information arrives. For example, if I am indoors and hear rain starting to fall, it might occur to me that the picnic will be canceled. Yet it will probably not occur to me that the seventeenth petal on the largest rose in my neighbor's garden will get wet; humans keep forward chaining under careful control, lest they be swamped with irrelevant consequences.

The backward-chaining algorithm, as its name suggests, works backward from the query. If the query  $q$  is known to be true, then no work is needed. Otherwise, the algorithm finds those implications in the knowledge base whose conclusion is  $q$ . If all the premises of one of those implications can be proved true (by backward chaining), then  $q$  is true. When applied to the query  $Q$  in Figure 7.16, it works back down the graph until it reaches a set of known facts,  $A$  and  $B$ , that forms the basis for a proof. The algorithm is essentially identical to the AND-OR-GRAPH-SEARCH algorithm in Figure 4.11. As with forward chaining, an efficient implementation runs in linear time.

REASONING

Backward chaining is a form of **goal-directed reasoning**. It is useful for answering G<sub>RASONING</sub> specific questions such as "What shall I do now?" and "Where are my keys?" Often, the cost of backward chaining is *much less* than linear in the size of the knowledge base, because the process touches only relevant facts.

# 7.6 EFFECTIVE PROPOSITIONAL MODEL CHECKING

In this section, we describe two families of efficient algorithms for general propositional inference based on model checking: One approach based on backtracking search, and one on local hill-climbing search. These algorithms are part of the "technology" of propositional logic. This section can be skimmed on a first reading of the chapter.