#### **LEARNING II**

### Remember search

- We can describe a state space search model as:
  - a state space *S*;
  - an initial state  $s_0$ ;
  - a set of actions,  $A(s) \subseteq A$ , applicable in each state  $s \in S$ ;
  - transition function f(s, a) for  $s \in S$  and  $a \in A$ ;
  - action costs c(a, s) > 0; and
  - a set of goal states  $G \subseteq S$
- How do we choose which action to use?
- We have a heuristic h(n) and choose based on:

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

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#### Overview

- Another form of learning is *reinforcement learning*.
- Here we don't have examples, but do have feedback from the environment.
- The agent learns by doing, occasionally getting things right, and then getting a reward.
- Rather like animals learn.

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#### Now learn heuristics

- We will start by assuming that the agent knows the results and costs of each action.
- We will also assume that it can build the whole search tree.
- This is what we do when we study search
- We then set h(n) = 0 for all n and pick the best node.
- When the agent has expanded node  $n_i$  to give a set of children  $\delta(n_i)$ , it updates its  $h(n_i)$  to be:

$$h(n_i) := \min_{nj \in \delta(n_i)} \left[ h(n_j) + c(n_i, n_j) 
ight]$$

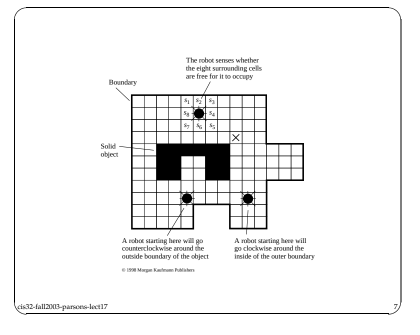
where  $c(n_i, n_j)$  is the cost of moving from  $n_i$  to  $n_j$ .

ullet We further assume that the agent can recognise the goal state and knows that h(goal) is 0.

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- This won't do much for the agent the first time—it is just uniform cost search.
- However, subsequent searches will "zoom in" on the right solution faster and faster.
- This happens as the  $h_T(n)$  values propagate back from the goal.
- (There are few enough values that these can be stored in a table.)
- Each run propagates the true cost of getting to the goal further back through the search.
- Eventually, the minimal cost path can just be read off the tree.

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### Learning without a model of action

- As described this kind of search is a "thought experiment" that an agent carries out.
- In the case of the navigating robot, it is planning its route across the grid.
- Alternatively it would be possible for the agent to actually carry out the operations to see what happens.
- In the case of the robot it could move through the room randomly at first, working out over a number of runs what the outcomes of actions were, and which were most useful at which point.
- (To do this, the agent will have to build a model of the state space in its "head").

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- What we assume is that:
  - The agent can distinguish the states it visits (and name them).
  - The agent knows how much actions cost once it has taken them.
- The process starts at the start state  $s_0$ .
- The agent then takes an action (maybe at random), and moves to another state. And repeats.
- As it visits each state, it names it and updates the heuristic value of this state as:

$$h(n_i) := [h(n_i) + c(n_i, n_i)]$$

where  $n_i$  is the node in which an action is taken,  $n_j$  is the node the action takes the agent to, and  $c(n_i, n_j)$  is the cost of the action.

•  $h(n_i)$  is zero if the node hasn't been reached before.

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• Whenever the agent has to choose an action *a*, it chooses it by:

$$a = \operatorname{argmin}_a \left[ h(\sigma(n_i, a)) + c(n_i, \sigma(n_i, a)) \right]$$

where  $\sigma(n_i, a)$  is the state reached from  $n_i$  after carrying out a.

- As before, the estimated minimum cost path to the goal is built up over repeated runs.
- However, allowing some randomness in the choice of actions increases the chance that the "estimated minimum cost path" really is the best path.

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- Potentially you could consider all the things it is possible to measure.
- Then:

$$h(n) = w_1 W(n) + w_2 P(n) + \dots$$

- We then learn which weights are best.
- One way to do this is as follows:
- After expanding  $n_i$  to  $\delta(n_i)$  we adjust the weights so that:

$$h(n_i) := h(n_i) + \beta \left( \min_{n_i \in \delta(n_i)} \left[ h(n_j) + c(n_i, n_j) \right] - h(n_i) \right)$$

• We modify  $h(n_i)$  by adding some proportion of (controlled by  $\beta$ ) of the difference between what we thought  $h(n_i)$  was before the expansion, and what we think it is after.

### Learning without a search graph

- For many interesting problems, it is not possible to store all the states/nodes and build the entire search graph.
- Provided we have a model of the effects of actions, we can still search with an evaluation function.
- We start by assembling a heuristic as a linear combination of some set of plausible functions.
- For the 8-puzzle these might be:
  - -W(n): number of tiles out of place.
  - -P(n): sum of distance each tile is from home.
- Plus any additional functions you can think of.

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• We can rewrite this as:

$$h(n_i) := (1 - \beta)h(n_i) + \beta \min_{n_i \in \delta(n_i)} [h(n_j) + c(n_i, n_j)]$$

- ullet eta controls how fast the agent learns—how much weight we give to the new estimate of the heuristic.
- If  $\beta = 0$  there is no adjustment.
- If  $\beta = 1$ ,  $h(n_i)$  is thrown away immediately.
- $\bullet$  Low values of  $\beta$  lead to slow learning, and high values mean that performance is erratic.
- Note that this *temporal difference approach* can also work without a model of the effects of actions (with suitable modification).

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## Rewards not goals

- For many tasks agents don't have short term goals, but instead accrue *rewards* over a period of time.
- ullet Instead of a plan, we want a *policy*  $\pi$  which says how the agent should act over time.
- Typically this is expressed as what action should be carried out in a given state.
- We express the reward an agent gets as  $r(n_i, a)$ , and if doing a in  $n_i$  takes the agent to  $n_j$ , then:

$$r(n_i, a) = -c(n_i, n_j) + \rho(n_j)$$

where  $\rho(n_j)$  is a reward for being in state  $n_j$ .

• We want an optimal policy  $\pi^*$  which maximises the (discounted) reward at every node.

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- Given a policy  $\pi$ , we can compute the value of each node—the reward the agent will get if it starts at that node and follows the policy.
- If the agent is at  $n_i$  and follows  $\pi$  to  $n_j$  then the agent will get reward:

$$V^{\pi}(n_i) = r(n_i, \pi(n_i)) + \gamma V^{\pi}(n_j)$$

where  $\gamma$  is the discount factor (think of it as the opposite of bank interest).

• The optimum policy then gives us the action that maximises this reward:

$$V^{\pi^*}(n_i) = \max_{a} \left[ r(n_i, a) + \gamma V^{\pi^*}(n_j) \right]$$

- One way to find the optimum policy is by searching through all possible policies.
- Start with a random policy and calculate its reward.
- Then guess another policy and see if it has a better reward (kind of slow).
- Better would be to tweak the policy by swapping some a in  $n_i$  for an a' with a higher  $r(n_i, a')$ .
- Again there is no guarantee of success.
- But there are better approaches.

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• If we knew what the values of the nodes were under  $\pi^*$ , then we could easily compute the optimal policy:

$$\pi^*(n_i) = \operatorname{argmax}_a \left[ r(n_i, a) + \gamma V^{\pi^*}(n_j) \right]$$

- The problem is that we don't know these values.
- But we can find them out using value iteration.
- ullet We start by guessing (randomly is fine) an estimated value V(n) for each node.

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• Then when we are at  $n_i$  we pick the action to maximise:

$$\operatorname{argmax}_{a} [r(n_i, a) + \gamma V(n_i)]$$

that is the best thing given what we currently know.

• We then update  $V(n_i)$  by:

$$V(n_i) := (1 - \beta)V(n_i) + \beta \left[ r(n_i, a), \gamma V(n_i) \right]$$

- ullet Progressive iterations of this calculation make V(n) a closer and closer approximation to  $V^{\pi^*}(n_i)$ .
- Intuitively this is because we replace the estimate with the actual reward we get for the next state (and the next state and the next state).

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# Summary

- This lecture has looked at a number of approaches to learning heuristic functions.
- We started assuming that the agent knew everything but the heuristic, and progressively relaxed assumptions.
- This created a battery of reinforcement learning methods that can be applied in a wide variety of situations.
- These models are not the best that we can do. Next week we will look at models for nondeterministic environments.

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