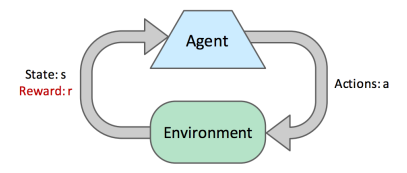
. These lecture notes are heavily based on notes originally written by Nikhil Sharma

Reinforcement Learning

In the previous note, we discussed Markov decision processes, which we solved using techniques such as value iteration and policy iteration to compute the optimal values of states and extract optimal policies. Solving Markov decision processes is an example of **offline planning**, where agents have full knowledge of both the **transition** function and the **reward** function, all the information they need to precompute optimal actions in the world encoded by the MDP without ever actually taking any actions. In this note, we'll discuss **online planning**, during which an agent has no prior knowledge of rewards or transitions in the world (still represented as a MDP). In online planning, an agent must try **exploration**, during which it performs actions and receives **feedback** in the form of the successor states it arrives in and the corresponding rewards it reaps. The agent uses this feedback to estimate an optimal policy through a process known as **reinforcement learning** before using this estimated policy for **exploitation** or reward maximization.

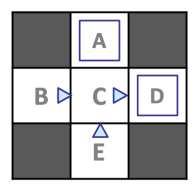


Let's start with some basic terminology. At each time step during online planning, an agent starts in a state s, then takes an action a and ends up in a successor state s', attaining some reward r. Each (s,a,s',r) tuple is known as a **sample**. Often, an agent continues to take actions and collect samples in succession until arriving at a terminal state. Such a collection of samples is known as an **episode**. Agents typically go through many episodes during exploration in order to collect sufficient data needed for learning.

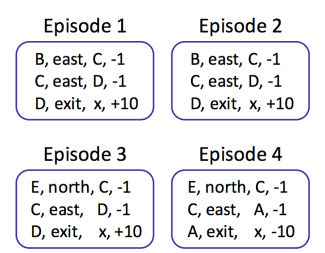
There are two types of reinforcement learning, **model-based learning** and **model-free learning**. Model-based learning attempts to estimate the transition and reward functions with the samples attained during exploration before using these estimates to solve the MDP normally with value or policy iteration. Model-free learning, on the other hand, attempts to estimate the values or Q-values of states directly, without ever using any memory to construct a model of the rewards and transitions in the MDP.

Model-Based Learning

In model-based learning an agent generates an approximation of the transition function, $\hat{T}(s,a,s')$, by keeping counts of the number of times it arrives in each state s' after entering each Q-state (s,a). The agent can then generate the the approximate transition function \hat{T} upon request by **normalizing** the counts it has collected - dividing the count for each observed tuple (s,a,s') by the sum over the counts for all instances where the agent was in Q-state (s,a). Normalization of counts scales them such that they sum to one, allowing them to be interpreted as probabilities. Consider the following example MDP with states $S = \{A,B,C,D,E,x\}$, with x representing the terminal state, and discount factor y = 1:



Assume we allow our agent to explore the MDP for four episodes under the policy $\pi_{explore}$ delineated above (a directional triangle indicates motion in the direction the triangle points, and a blue squares represents taking exit as the action of choice), and yield the following results:



We now have a collective 12 samples, 3 from each episode with counts as follows:

S	a	\mathbf{s}'	count	
A	exit	x	1	
В	east	C	2	
C	east	A	1	
C	east	D	3	
D	exit	х	3	
E	north	C	2	

Recalling that T(s,a,s') = P(s'|a,s), we can estimate the transition function with these counts by dividing the counts for each tuple (s,a,s') by the total number of times we were in Q-state (s,a) and the reward function directly from the rewards we reaped during exploration:

• Transition Function:
$$\hat{T}(s,a,s')$$

• Reward Function: $\hat{R}(s,a,s')$
• $\hat{T}(A,exit,x) = \frac{\#(A,exit,x)}{\#(A,exit)} = \frac{1}{1} = 1$
• $\hat{T}(B,east,C) = \frac{\#(B,east,C)}{\#(B,east)} = \frac{2}{2} = 1$
• $\hat{T}(C,east,A) = \frac{\#(C,east,A)}{\#(C,east)} = \frac{1}{4} = 0.25$
• $\hat{R}(A,exit,x) = -10$
• $\hat{R}(B,east,C) = -1$
• $\hat{R}(B,east,C) = -1$
• $\hat{R}(C,east,A) = -1$
• $\hat{R$

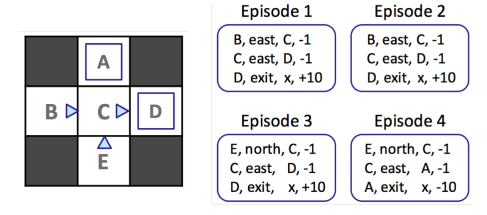
By the **law of large numbers**, as we collect more and more samples by having our agent experience more episodes, our models of \hat{T} and \hat{R} will improve, with \hat{T} converging towards T and \hat{R} acquiring knowledge of previously undiscovered rewards as we discover new (s,a,s') tuples. Whenever we see fit, we can end our agent's training to generate a policy $\pi_{exploit}$ by running value or policy iteration with our current models for \hat{T} and \hat{R} and use $\pi_{exploit}$ for exploitation, having our agent traverse the MDP taking actions seeking reward maximization rather than seeking learning. We'll soon discuss methods for how to allocate time between exploration and exploitation effectively. Model-based learning is very simple and intuitive yet remarkably effective, generating \hat{T} and \hat{R} with nothing more than counting and normalization. However, it can be expensive to maintain counts for every (s,a,s') tuple seen, and so in the next section on model-free learning we'll develop methods to bypass maintaining counts altogether and avoid the memory overhead required by model-based learning.

Model-Free Learning

Onward to model-free learning! There are several model-free learning algorithms, and we'll cover three of them: direct evaluation, temporal difference learning, and Q-learning. Direct evaluation and temporal difference learning fall under a class of algorithms known as **passive reinforcement learning**. In passive reinforcement learning, an agent is given a policy to follow and learns the value of states under that policy as it experiences episodes, which is exactly what is done by policy evaluation for MDPs when *T* and *R* are known. Q-learning falls under a second class of model-free learning algorithms known as **active reinforcement learning**, during which the learning agent can use the feedback it receives to iteratively update its policy while learning until eventually determining the optimal policy after sufficient exploration.

Direct Evaluation

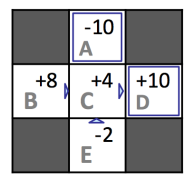
The first passive reinforcement learning technique we'll cover is known as **direct evaluation**, a method that's as boring and simple as the name makes it sound. All direct evaluation does is fix some policy π and have the agent experience several episodes while following π . As the agent collects samples through these episodes it maintains counts of the total utility obtained from each state and the number of times it visited each state. At any point, we can compute the estimated value of any state s by dividing the total utility obtained from s by the number of times s was visited. Let's run direct evaluation on our example from earlier, recalling that $\gamma = 1$.



Walking through the first episode, we can see that from state D to termination we acquired a total reward of 10, from state C we acquired a total reward of (-1) + 10 = 9, and from state B we acquired a total reward of (-1) + (-1) + 10 = 8. Completing this process yields the total reward across episodes for each state and the resulting estimated values as follows:

S	Total Reward	Times Visited	$V^{\pi}(s)$
A	-10	1	-10
В	16	2	8
C	16	4	4
D	30	3	10
E	-4	2	-2

Though direct evaluation eventually learns state values for each state, it's often unnecessarily slow to converge because it wastes information about transitions between states.



In our example, we computed $V^{\pi}(E) = -2$ and $V^{\pi}(B) = 8$, though based on the feedback we received both states only have C as a successor state and incur the same reward of -1 when transitioning to C. According to the Bellman equation, this means that both B and E should have the same value under π . However, of the 4 times our agent was in state C, it transitioned to D and reaped a reward of 10 three times and transitioned to D and reaped a reward of 10 three times and transitioned to D and reaped a reward of D are the interest of D and reaped a reward of D are the interest of D and D are the single time it received the D reward it started in state D and D will converge to their true values, but cases like this cause the process to take longer than we'd like. This issue can be mitigated by choosing to use our second passive reinforcement learning algorithm, temporal difference learning.

Temporal Difference Learning

Temporal difference learning (TD learning) uses the idea of *learning from every experience*, rather than simply keeping track of total rewards and number of times states are visited and learning at the end as direct evaluation does. In policy evaluation, we used the system of equations generated by our fixed policy and the Bellman equation to determine the values of states under that policy (or used iterative updates like with value iteration).

$$V^{\pi}(s) = \sum_{s'} T(s, \pi(s), s') [R(s, \pi(s), s') + \gamma V^{\pi}(s')]$$

Each of these equations equates the value of one state to the weighted average over the discounted values of that state's successors plus the rewards reaped in transitioning to them. TD learning tries to answer the question of how to compute this weighted average without the weights, cleverly doing so with an **exponential moving average**. We begin by initializing $\forall s, V^{\pi}(s) = 0$. At each time step, an agent takes an action $\pi(s)$ from a state s, transitions to a state s', and receives a reward $R(s, \pi(s), s')$. We can obtain a **sample value** by summing the received reward with the discounted current value of s' under π :

sample =
$$R(s, \pi(s), s') + \gamma V^{\pi}(s')$$

This sample is a new estimate for $V^{\pi}(s)$. The next step is to incorporate this sampled estimate into our existing model for $V^{\pi}(s)$ with the exponential moving average, which adheres to the following update rule:

$$V^{\pi}(s) \leftarrow (1-\alpha)V^{\pi}(s) + \alpha \cdot \text{sample}$$

Above, α is a parameter constrained by $0 \le \alpha \le 1$ known as the **learning rate** that specifies the weight we want to assign our existing model for $V^{\pi}(s)$, $1-\alpha$, and the weight we want to assign our new sampled estimate, α . It's typical to start out with learning rate of $\alpha = 1$, accordingly assigning $V^{\pi}(s)$ to whatever the first sample happens to be, and slowly shrinking it towards 0, at which point all subsequent samples will be zeroed out and stop affecting our model of $V^{\pi}(s)$.

Let's stop and analyze the update rule for a minute. Annotating the state of our model at different points in time by defining $V_k^{\pi}(s)$ and sample_k as the estimated value of state s after the k^{th} update and the k^{th} sample respectively, we can reexpress our update rule:

$$V_k^{\pi}(s) \leftarrow (1-\alpha)V_{k-1}^{\pi}(s) + \alpha \cdot \text{sample}_k$$

This recursive definition for $V_k^{\pi}(s)$ happens to be very interesting to expand:

$$\begin{array}{lll} V_k^\pi(s) & \leftarrow & (1-\alpha)V_{k-1}^\pi(s) + \alpha \cdot \mathrm{sample}_k \\ V_k^\pi(s) & \leftarrow & (1-\alpha)[(1-\alpha)V_{k-2}^\pi(s) + \alpha \cdot \mathrm{sample}_{k-1}] + \alpha \cdot \mathrm{sample}_k \\ V_k^\pi(s) & \leftarrow & (1-\alpha)^2V_{k-2}^\pi(s) + (1-\alpha) \cdot \alpha \cdot \mathrm{sample}_{k-1} + \alpha \cdot \mathrm{sample}_k \\ & \vdots \\ V_k^\pi(s) & \leftarrow & (1-\alpha)^kV_0^\pi(s) + \alpha \cdot [(1-\alpha)^{k-1} \cdot \mathrm{sample}_1 + \ldots + (1-\alpha) \cdot \mathrm{sample}_{k-1} + \mathrm{sample}_k] \\ V_k^\pi(s) & \leftarrow & \alpha \cdot [(1-\alpha)^{k-1} \cdot \mathrm{sample}_1 + \ldots + (1-\alpha) \cdot \mathrm{sample}_{k-1} + \mathrm{sample}_k] \end{array}$$

Because $0 \le (1 - \alpha) \le 1$, as we raise the quantity $(1 - \alpha)$ to increasingly larger powers, it grows closer and closer to 0. By the update rule expansion we derived, this means that older samples are given exponentially less weight, exactly what we want since these older samples are computed using older (and hence worse) versions of our model for $V^{\pi}(s)$! This is the beauty of temporal difference learning - with a single straightforward update rule, we are able to:

- learn at every time step, hence using information about state transitions as we get them since we're using iteratively updated versions of $V^{\pi}(s')$ in our samples rather than waiting until the end to perform any computation.
- give exponentially less weight to older, potentially less accurate samples.
- converge to learning true state values much faster with fewer episodes than direct evaluation.

Q-Learning

Both direct evaluation and TD learning will eventually learn the true value of all states under the policy they follow. However, they both have a major inherent issue - we want to find an optimal *policy* for our agent, which requires knowledge of the Q-values of states. To compute Q-values from the values we have, we require a transition function and reward function as dictated by the Bellman equation.

$$Q^*(s,a) = \sum_{s'} T(s,a,s') [R(s,a,s') + \gamma V^*(s')]$$

As a result, TD learning or direct evaluation are typically used in tandem with some model-based learning to acquire estimates of *T* and *R* in order to effectively update the policy followed by the learning agent. This became avoidable by a revolutionary new idea known as **Q-learning**, which proposed learning the Q-values of states directly, bypassing the need to ever know any values, transition functions, or reward functions. As a result, Q-learning is entirely model-free. Q-learning uses the following update rule to perform what's known as **Q-value iteration**:

$$Q_{k+1}(s,a) \leftarrow \sum_{s'} T(s,a,s') [R(s,a,s') + \gamma \max_{a'} Q_k(s',a')]$$

Note that this update is only a slight modification over the update rule for value iteration. Indeed, the only real difference is that the position of the max operator over actions has been changed since we select an action before transitioning when we're in a state, but we transition before selecting a new action when we're in a Q-state.

With this new update rule under our belt, Q-learning is derived essentially the same way as TD learning, by acquiring **Q-value samples**:

sample =
$$R(s, a, s') + \gamma \max_{a'} Q(s', a')$$

and incorporating them into an exponential moving average.

$$Q(s,a) \leftarrow (1-\alpha)Q(s,a) + \alpha \cdot \text{sample}$$

As long as we spend enough time in exploration and decrease the learning rate α at an appropriate pace, Q-learning learns the optimal Q-values for every Q-state. This is what makes Q-learning so revolutionary - while TD learning and direct evaluation learn the values of states under a policy by following the policy before determining policy optimality via other techniques, Q-learning can learn the optimal policy directly even by taking suboptimal or random actions. This is called **off-policy learning** (contrary to direct evaluation and TD learning, which are examples of **on-policy learning**).

Approximate Q-Learning

Q-learning is an incredible learning technique that continues to sit at the center of developments in the field of reinforcement learning. Yet, it still has some room for improvement. As it stands, Q-learning just stores

all Q-values for states in tabular form, which is not particularly efficient given that most applications of reinforcement learning have several thousands or even millions of states. This means we can't visit all states during training and can't store all Q-values even if we could for lack of memory.







Figure 1

Figure 2

Figure 3

Above, if Pacman learned that Figure 1 is unfavorable after running vanilla Q-learning, it would still have no idea that Figure 2 or even Figure 3 are unfavorable as well. **Approximate Q-learning** tries to account for this by learning about a few general situations and extrapolating to many similar situations. The key to generalizing learning experiences is the **feature-based representation** of states, which represents each state as a vector known as a **feature vector**. For example, a feature vector for Pacman may encode

- the distance to the closest ghost.
- the distance to the closest food pellet.
- the number of ghosts.
- is Pacman trapped? 0 or 1

With feature vectors, we can treat values of states and Q-states as linear value functions:

$$V(s) = w_1 \cdot f_1(s) + w_2 \cdot f_2(s) + \dots + w_n \cdot f_n(s) = \vec{w} \cdot \vec{f}(s)$$

$$Q(s,a) = w_1 \cdot f_1(s,a) + w_2 \cdot f_2(s,a) + \dots + w_n \cdot f_n(s,a) = \vec{w} \cdot \vec{f}(s,a)$$

where $\vec{f}(s) = \begin{bmatrix} f_1(s) & f_2(s) & \dots & f_n(s) \end{bmatrix}^T$ and $\vec{f}(s,a) = \begin{bmatrix} f_1(s,a) & f_2(s,a) & \dots & f_n(s,a) \end{bmatrix}^T$ represent the feature vectors for state s and Q-state (s,a) respectively and $\vec{w} = \begin{bmatrix} w_1 & w_2 & \dots & w_n \end{bmatrix}$ represents a weight vector. Defining difference as

difference =
$$[R(s, a, s') + \gamma \max_{a'} Q(s', a')] - Q(s, a)$$

approximate Q-learning works almost identically to Q-learning, using the following update rule:

$$w_i \leftarrow w_i + \alpha \cdot \text{difference} \cdot f_i(s, a)$$

Rather than storing Q-values for each and every state, with approximate Q-learning we only need to store a single weight vector and can compute Q-values on-demand as needed. As a result, this gives us not only a more generalized version of Q-learning, but a significantly more memory-efficient one as well.

As a final note on Q-learning, we can reexpress the update rule for exact Q-learning using difference as follows:

$$Q(s,a) \leftarrow Q(s,a) + \alpha \cdot \text{difference}$$

This second notation gives us a slightly different but equally valuable interpretation of the update: it's computing the difference between the sampled estimated and the current model of Q(s,a), and shifting the model in the direction of the estimate with the magnitude of the shift being proportional to the magnitude of the difference.