# A Guided Tour of Chapter 9: Reinforcement Learning for Prediction

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#### RL does not have access to a probability model

- ullet DP/ADP assume access to probability model (knowledge of  $\mathcal{P}_R$ )
- Often in real-world, we do not have access to these probabilities
- Which means we'd need to interact with the actual environment
- Actual Environment serves up individual experiences, not probabilities
- Even if MDP model is available, model updates can be challenging
- Often real-world models end up being too large or too complex
- Sometimes estimating a sampling model is much more feasible
- So RL interacts with either actual or simulated environment
- Either way, we receive individual experiences of next state and reward
- RL learns Value Functions from a stream of individual experiences
- How does RL solve Prediction and Control with such limited access?

## The RL Approach

- Like humans/animals, RL doesn't aim to estimate probability model
- Rather, RL is a "trial-and-error" approach to linking actions to returns
- This is hard because actions have overlapping reward sequences
- Also, sometimes actions result in delayed rewards
- The key is incrementally updating Q-Value Function from experiences
- Appropriate Approximation of Q-Value Function is also key to success
- RL algorithms are founded on the Bellman Equations
- Moreover, RL Control is based on Generalized Policy Iteration
- This lecture/chapter focuses on RL for Prediction

#### **RL** for Prediction

- ullet Prediction: Problem of estimating MDP Value Function for a policy  $\pi$
- ullet Equivalently, problem of estimating  $\pi$ -implied MRP's Value Function
- Assume interface serves an atomic experience of (next state, reward)
- Interacting with this interface repeatedly provides a trace experience

$$S_0, R_1, S_1, R_2, S_2, \dots$$

• Value Function  $V: \mathcal{N} \to \mathbb{R}$  of an MRP is defined as:

$$V(s) = \mathbb{E}[G_t | S_t = s]$$
 for all  $s \in \mathcal{N}, \text{ for all } t = 0, 1, 2, \dots$ 

where the *Return G*<sup>t</sup> for each t = 0, 1, 2, ... is defined as:

$$G_{t} = \sum_{i=t+1}^{\infty} \gamma^{i-t-1} \cdot R_{i} = R_{t+1} + \gamma \cdot R_{t+2} + \gamma^{2} \cdot R_{t+3} + \dots = R_{t+1} + \gamma \cdot G_{t+1}$$

#### Code interface for RL Prediction

```
An atomic experience is represented as a TransitionStep[S]
@dataclass(frozen=True)
class TransitionStep(Generic[S]):
    state: S
    next_state: S
    reward: float
```

Input to RL prediction can be either of:

- Atomic Experiences as Iterable [TransitionStep [S]], or
- Trace Experiences as Iterable [Iterable [TransitionStep [S]]]

Note that Iterable can be either a Sequence or an Iterator (i.e., stream)

## Monte-Carlo (MC) Prediction

- Supervised learning with states and returns from trace experiences
- Incremental estimation with update method of FunctionApprox
- x-values are states  $S_t$ , y-values are returns  $G_t$
- Note that updates can be done only at the end of a trace experience
- Returns calculated with a backward walk:  $G_t = R_{t+1} + \gamma \cdot G_{t+1}$

$$\mathcal{L}_{(S_t,G_t)}(\boldsymbol{w}) = \frac{1}{2} \cdot (V(S_t; \boldsymbol{w}) - G_t)^2$$

$$\nabla_{\mathbf{w}} \mathcal{L}_{(S_t,G_t)}(\mathbf{w}) = (V(S_t; \mathbf{w}) - G_t) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

$$\Delta \mathbf{w} = \alpha \cdot (G_t - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

## Structure of the parameters update formula

$$\Delta \mathbf{w} = \alpha \cdot (G_t - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

The update  $\Delta w$  to parameters w should be seen as product of:

- Learning Rate  $\alpha$
- Return Residual of the observed return  $G_t$  relative to the estimated conditional expected return  $V(S_t; \mathbf{w})$
- Estimate Gradient of the conditional expected return  $V(S_t; \mathbf{w})$  with respect to the parameters  $\mathbf{w}$

This structure (as product of above 3 entities) will be a repeated pattern.

#### Tabular MC Prediction

- ullet Finite state space, let's say non-terminal states  $\mathcal{N} = \{s_1, s_2, \dots, s_m\}$
- Denote  $V_n(s_i)$  as estimate of VF after the *n*-th occurrence of  $s_i$
- Denote  $Y_i^{(1)}, Y_i^{(2)}, \dots, Y_i^{(n)}$  as returns for first n occurrences of  $s_i$
- ullet Denote count\_to\_weight\_func attribute of Tabular as  $f(\cdot)$
- Then the Tabular update at the n-th occurrence of  $s_i$  is:

$$V_n(s_i) = (1 - f(n)) \cdot V_{n-1}(s_i) + f(n) \cdot Y_i^{(n)}$$
  
=  $V_{n-1}(s_i) + f(n) \cdot (Y_i^{(n)} - V_{n-1}(s_i))$ 

- So update to VF for s<sub>i</sub> is Latest Weight times Return Residual
- For default setting of count\_to\_weight\_func as  $f(n) = \frac{1}{n}$ :

$$V_n(s_i) = \frac{n-1}{n} \cdot V_{n-1}(s_i) + \frac{1}{n} \cdot Y_i^{(n)} = V_{n-1}(s_i) + \frac{1}{n} \cdot (Y_i^{(n)} - V_{n-1}(s_i))$$

#### Tabular MC Prediction

Expanding the incremental updates across values of n, we get:

$$V_n(s_i) = f(n) \cdot Y_i^{(n)} + (1 - f(n)) \cdot f(n-1) \cdot Y_i^{(n-1)} + \dots \dots + (1 - f(n)) \cdot (1 - f(n-1)) \cdot \dots \cdot (1 - f(2)) \cdot f(1) \cdot Y_i^{(1)}$$

• For default setting of count\_to\_weight\_func as  $f(n) = \frac{1}{n}$ :

$$V_{n}(s_{i}) = \frac{1}{n} \cdot Y_{i}^{(n)} + \frac{n-1}{n} \cdot \frac{1}{n-1} \cdot Y_{i}^{(n-1)} + \dots$$
$$\dots + \frac{n-1}{n} \cdot \frac{n-2}{n-1} \cdot \dots \cdot \frac{1}{2} \cdot \frac{1}{1} \cdot Y_{i}^{(1)} = \frac{\sum_{k=1}^{n} Y_{i}^{(k)}}{n}$$

- Tabular MC is simply incremental calculation of averages of returns
- Exactly the calculation in the update method of Tabular class
- View Tabular MC as an application of Law of Large Numbers

## Tabular MC as a special case of Linear Func Approximation

- Features functions are indicator functions for states
- Linear-approx parameters are Value Function estimates for states
- count\_to\_weight\_func plays the role of learning rate
- So tabular Value Function update can be written as:

$$w_i^{(n)} = w_i^{(n-1)} + \alpha_n \cdot (Y_i^{(n)} - w_i^{(n-1)})$$

- $Y_i^{(n)} w_i^{(n-1)}$  represents the gradient of the loss function
- For non-stationary problems, algorithm needs to "forget" distant past
- With constant learning rate  $\alpha$ , time-decaying weights:

$$V_n(s_i) = \alpha \cdot Y_i^{(n)} + (1 - \alpha) \cdot \alpha \cdot Y_i^{(n-1)} + \ldots + (1 - \alpha)^{n-1} \cdot \alpha \cdot Y_i^{(1)}$$
$$= \sum_{j=1}^n \alpha \cdot (1 - \alpha)^{n-j} \cdot Y_i^{(j)}$$

• Weights sum to 1 asymptotically:  $\lim_{n\to\infty}\sum_{j=1}^n \alpha\cdot(1-\alpha)^{n-j}=1$ 

#### Each-Visit MC and First-Visit MC

- The MC algorithm we covered is known as Each-Visit Monte-Carlo
- Because we include each occurrence of a state in a trace experience
- Alternatively, we can do First-Visit Monte-Carlo
- Only the first occurrence of a state in a trace experience is considered
- Keep track of whether a state has been visited in a trace experience
- MC Prediction algorithms are easy to understand and implement
- MC produces unbiased estimates but can be slow to converge
- Key disadvantage: MC requires complete trace experiences

## Temporal-Difference (TD) Prediction

- To understand TD, we start with Tabular TD Prediction
- Key: Exploit recursive structure of VF in MRP Bellman Equation
- Replace  $G_t$  with  $R_{t+1} + \gamma \cdot V(S_{t+1})$  using atomic experience data
- So we are bootstrapping the VF ("estimate from estimate")
- ullet The tabular MC Prediction update (for constant lpha) is modified from:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot (G_t - V(S_t))$$

to:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - V(S_t))$$

- $R_{t+1} + \gamma \cdot V(S_{t+1})$  known as TD target
- $\delta_t = R_{t+1} + \gamma \cdot V(S_{t+1}) V(S_t)$  known as *TD Error*
- TD Error is the crucial quantity it represents "sample Bellman Error"
- VF is adjusted so as to bridge TD error (on an expected basis)

#### TD updates after each atomic experience

- Unlike MC, we can use TD when we have incomplete traces
- Often in real-world situations, experiments gets curtailed/disrupted
- Also, we can use TD in non-episodic (known as continuing) traces
- TD updates VF after each atomic experience ("continuous learning")
- So TD can be run on any stream of atomic experiences
- This means we can chop up the input stream and serve in any order

## TD Prediction with Function Approximation

- Each atomic experience leads to a parameters update
- To understand how parameters update work, consider:

$$\mathcal{L}_{(S_t, S_{t+1}, R_{t+1})}(\mathbf{w}) = \frac{1}{2} \cdot (V(S_t; \mathbf{w}) - (R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w})))^2$$

- Above formula replaces  $G_t$  (of MC) with  $R_{t+1} + \gamma \cdot V(S_{t+1}, \mathbf{w})$
- Unlike MC, in TD, we don't take the gradient of this loss function
- ullet "Cheat" in gradient calc by ignoring dependency of  $V(S_{t+1}; oldsymbol{w})$  on  $oldsymbol{w}$
- This "gradient with cheating" calculation is known as semi-gradient
- So we pretend the only dependency on  ${\it w}$  is through  $V(S_t; {\it w})$

$$\Delta \mathbf{w} = \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w}) - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

#### Structure of the parameters update formula

$$\Delta \mathbf{w} = \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w}) - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

The update  $\Delta w$  to parameters w should be seen as product of:

- Learning Rate  $\alpha$
- TD Error  $\delta_t = R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w}) V(S_t; \mathbf{w})$
- Estimate Gradient of the conditional expected return  $V(S_t; \mathbf{w})$  with respect to the parameters  $\mathbf{w}$

So parameters update formula has same product-structure as MC

## TD's many benefits

- "TD is the most significant and innovative idea in RL" Rich Sutton
- Blends the advantages of DP and MC
- Like DP, TD learns by bootstrapping (drawing from Bellman Eqn)
- Like MC, TD learns from experiences without access to probabilities
- So TD overcomes curse of dimensionality and curse of modeling
- TD also has the advantage of not requiring entire trace experiences
- Most significantly, TD is akin to human (continuous) learning

# Bias, Variance and Convergence of TD versus MC

- ullet MC uses  $G_t$  is an unbiased estimate of the Value Function
- This helps MC with convergence even with function approximation
- TD uses  $R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w})$  as a biased estimate of the VF
- ullet Tabular TD prediction converges to true VF in the mean for const lpha
- And converges to true VF under Robbins-Monro learning rate schedule

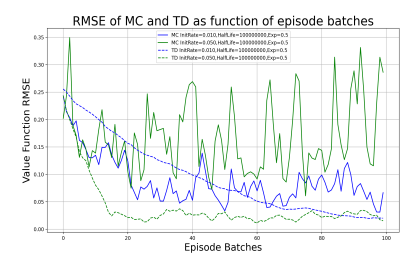
$$\sum_{n=1}^{\infty} \alpha_n = \infty \text{ and } \sum_{n=1}^{\infty} \alpha_n^2 < \infty$$

- However, Robbins-Monro schedule is not so useful in practice
- TD Prediction with func-approx does not always converge to true VF
- Most convergence proofs are for Tabular, some for linear func-approx
- TD Target  $R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w})$  has much lower variance that  $G_t$
- ullet G<sub>t</sub> depends on many random rewards whose variances accumulate
- TD Target depends on only the next reward, so lower variance

# Speed of Convergence of TD versus MC

- We typically compare algorithms based on:
  - Speed of convergence
  - Efficiency in use of limited set of experiences data
- There are no formal proofs for MC v/s TD on above criterion
- MC and TD have significant differences in their:
  - Usage of data
  - Nature of updates
  - Frequency of updates
- So unclear exactly how to compare them apples to apples
- ullet Typically, MC and TD are compared with constant lpha
- ullet Practically/empirically, TD does better than MC with constant lpha
- Also, MC is not very sensitive to initial Value Function, but TD is

#### Convergence of MC versus TD with constant $\alpha$



#### RMSE of MC versus TD as function of episodes

- Symmetric random walk with barrier B = 10 and no discounting
- Graph depicts RMSE after every 7th episode (700 episodes in all)
- Blue curves for constant  $\alpha = 0.01$ , green for constant  $\alpha = 0.05$
- Notice how MC has significantly more variance
- RMSE progression is quite slow on blue curves (small learning rate)
- MC progresses quite fast initially but then barely progresses
- TD gets to fairly small RMSE quicker than corresponding MC
- $\bullet$  This performance of TD versus MC is typical for constant  $\alpha$

## Fixed-Data Experience Replay on TD versus MC

- So far, we've understood how TD learns versus how MC learns
- Now we want to understand what TD learns versus what MC learns
- To illustrate, we consider a finite set of trace experiences
- The agent can tap into this finite set of traces experiences endlessly
- But everything is ultimately sourced from this finite data set
- So we'd end up tapping into these experiences repeatedly
- We call this technique Experience Replay

#### MC and TD learn different Value Functions

- It is quite obvious what MC Prediction algorithm would learn
- MC Prediction is simply supervised learning with (state, return) pairs
- But here those pairs ultimately come from the given finite pairs
- So, MC estimates Value Function as average returns in the finite data
- Running MC Prediction algo matches explicit average returns calc
- But running TD Prediction algo gives significantly different answer
- So what is TD Prediction algorithm learning?
- TD drives towards VF of MRP implied by the finite experiences
- ullet Specifically, learns MLE for  $\mathcal{P}_R$  from the given finite data

$$\mathcal{P}_{R}(s, r, s') = \frac{\sum_{i=1}^{N} \mathbb{I}_{S_{i} = s, R_{i+1} = r, S_{i+1} = s'}}{\sum_{i=1}^{N} \mathbb{I}_{S_{i} = s}}$$

• TD is advantageous in Markov environments, MC in non-Markov

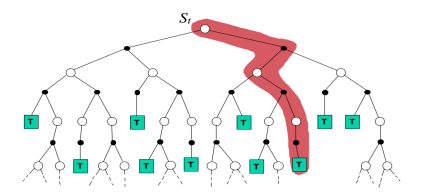
## Bootstrapping and Experiencing

- We summarize MC, TD and DP in terms of whether they:
  - Bootstrap: Update to VF utilizes a current or prior estimate of the VF
  - Experience: Interaction with actual or simulated environment
- TD and DP do bootstrap (updates use current/prior estimate of VF)
- MC does not bootstrap (updates use trace experience returns)
- MC and TD do experience (actual/simulated environment interaction)
- DP does not experience (updates use transition probabilities)
- Bootstrapping means backups are shallow (MC backups are deep)
- Experiencing means backups are narrow (DP backups are wide)

# MC backup Diagram

#### **Monte Carlo (Supervised Learning) (MC)**

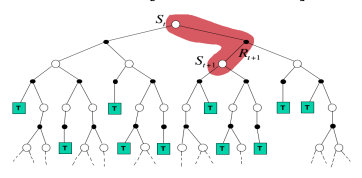
$$V(S_t) \leftarrow V(S_t) + \alpha \left[ G_t - V(S_t) \right]$$



## TD backup Diagram

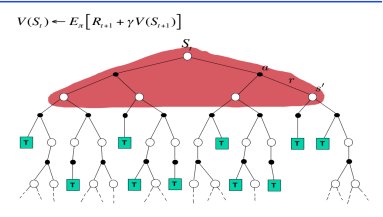
#### **Simplest TD Method**

$$V(S_t) \leftarrow V(S_t) + \alpha \left[ R_{t+1} + \gamma V(S_{t+1}) - V(S_t) \right]$$



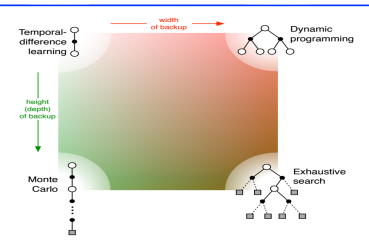
## DP backup Diagram

#### cf. Dynamic Programming



#### Unified View of RL

#### **Unified View**



#### Tabular *n*-step Bootstrapping

Tabular TD Prediction bootstraps the Value Function with update:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - V(S_t))$$

• So it's natural to extend this to bootstrapping with 2 steps ahead:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot (R_{t+1} + \gamma \cdot R_{t+2} + \gamma^2 \cdot V(S_{t+2}) - V(S_t))$$

• Generalize to bootstrapping with  $n \ge 1$  time steps ahead:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot (G_{t,n} - V(S_t))$$

•  $G_{t,n}$  (known as *n*-step bootstrapped return) is defined as:

$$G_{t,n} = \sum_{i=t+1}^{t+n} \gamma^{i-t-1} \cdot R_i + \gamma^n \cdot V(S_{t+n})$$
  
=  $R_{t+1} + \gamma \cdot R_{t+2} + \gamma^2 \cdot R_{t+3} + \ldots + \gamma^{n-1} \cdot R_{t+n} + \gamma^n \cdot V(S_{t+n})$ 

#### n-step Bootstrapping with Function Approximation

Generalizing this to the case of Function Approximation, we get:

$$\Delta \mathbf{w} = \alpha \cdot (G_{t,n} - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

- This looks similar to formula for parameters update for MC and TD
- In terms of conceptualizing the change in parameters as product of:
  - Learning Rate  $\alpha$
  - *n-step Bootstrapped Error*  $G_{t,n} V(S_t; \mathbf{w})$
  - Estimate Gradient of the conditional expected return  $V(S_t; \boldsymbol{w})$  with respect to the parameters  $\boldsymbol{w}$
- n serves as a parameter taking us across the spectrum from TD to MC
- n = 1 is the case of TD while sufficiently large n is the case of MC

#### $\lambda$ -Return Prediction Algorithm

• Instead of  $G_{t,n}$ , a valid target is a weighted-average target:

$$\sum_{n=1}^{N} u_n \cdot G_{t,n} + u \cdot G_t \text{ where } u + \sum_{n=1}^{N} u_n = 1$$

- ullet Any of the  $u_n$  or u can be 0, as long as they all sum up to 1
- The  $\lambda$ -Return target is a special case of weights  $u_n$  and u

$$u_n = (1 - \lambda) \cdot \lambda^{n-1}$$
 for all  $n = 1, \dots, T - t - 1$   $u_n = 0$  for all  $n \ge T - t$  and  $u = \lambda^{T - t - 1}$ 

• We denote the  $\lambda$ -Return target as  $G_t^{(\lambda)}$ , defined as:

$$G_t^{(\lambda)} = (1 - \lambda) \cdot \sum_{n=1}^{T-t-1} \lambda^{n-1} \cdot G_{t,n} + \lambda^{T-t-1} \cdot G_t$$
$$\Delta \mathbf{w} = \alpha \cdot (G_t^{(\lambda)} - V(S_t; \mathbf{w})) \cdot \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

#### Online versus Offline

- Note that for  $\lambda=0$ , the  $\lambda$ -Return target reduces to the TD target
- Note that for  $\lambda=1$ , the  $\lambda$ -Return target reduces to the MC target  $\mathit{G}_{t}$
- ullet  $\lambda$  parameter enables us to finely tune from TD ( $\lambda=0$ ) to MC ( $\lambda=1$ )
- ullet Note that for  $\lambda>0$ , updates are made only at the end of an episode
- Algorithms updating at end of episodes known as Offline Algorithms
- Online algorithms (updates after each time step) are appealing:
  - Updated VF can be utilized immediately for next time step's update
  - This facilitates continuous/fast learning
- Can we have a similar  $\lambda$ -tunable online algorithm for Prediction?
- Yes this is known as the  $TD(\lambda)$  Prediction algorithm

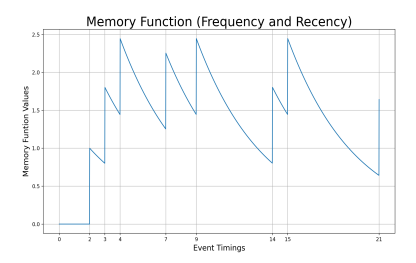
# Memory Function

- ullet TD( $\lambda$ ) algorithm is based on the concept of *Eligibility Traces*
- ullet We introduce the concept by defining a Memory Function M(t)
- ullet Assume an event occurs at times  $t_1 < t_2 < \ldots < t_n \in \mathbb{R}_{\geq 0}$
- We want M(t) to remember the # of times the event has occurred
- But we also want it to have an element of "forgetfulness"
- Recent event-occurrences remembered better than older occurrences
- We want  $M(\cdot)$  to give us a time-decayed count of event-occurrences

$$M(t) = \begin{cases} \mathbb{I}_{t=t_1} & \text{if } t \leq t_1, \\ M(t_i) \cdot \theta^{t-t_i} + \mathbb{I}_{t=t_{i+1}} & \text{if } t_i < t \leq t_{i+1} \text{ for any } 1 \leq i < n, \\ M(t_n) \cdot \theta^{t-t_n} & \text{otherwise (i.e., } t > t_n) \end{cases}$$

- ullet There's an uptick of 1 each time the event occurs, but it decays by a factor of  $heta^{\Delta t}$  over any interval  $\Delta t$  where the event doesn't occur
- ullet Thus,  $M(\cdot)$  captures the notion of frequency as well as recency

## Memory Function with $\theta = 0.8$



# Eligibility Traces and Tabular $TD(\lambda)$ Prediction

- ullet Assume a finite state space with non-terminals  $\mathcal{N} = \{s_1, s_2, \dots, s_m\}$
- Eligibility Trace for each state  $s \in S$  is defined as the Memory function  $M(\cdot)$  with  $\theta = \gamma \cdot \lambda$ , and the event timings are the time steps at which the state s occurs in a trace experience
- Eligibility trace for a given trace experience at time t is a function

$$E_t: \mathcal{N} \to \mathbb{R}_{\geq 0}$$

$$E_0(s) = 0$$
, for all  $s \in \mathcal{N}$ 

$$E_t(s) = \gamma \cdot \lambda \cdot E_{t-1}(s) + \mathbb{I}_{S_t=s}$$
, for all  $s \in \mathcal{N}$ , for all  $t = 1, 2, ...$ 

• Tabular  $TD(\lambda)$  Prediction algorithm performs following update at each time step t in each trace experience:

$$V(s) \leftarrow V(s) + \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - V(S_t)) \cdot E_t(s), \text{ for all } s \in \mathcal{N}$$

# "Equivalence" of $TD(\lambda)$ and $\lambda$ -Return

- TD( $\lambda$ ) is an online algorithm, similar to TD
- But unlike TD, we update the VF for all states at each time step
- VF update for each state is proportional to TD-Error  $\delta_t$  (like TD)
- But here,  $\delta_t$  is scaled by  $E_t(s)$  for each state s at each t

$$V(s) \leftarrow V(s) + \alpha \cdot \delta_t \cdot E_t(s)$$
, for all  $s \in \mathcal{N}$ 

- But how is  $TD(\lambda)$  Prediction linked to the  $\lambda$ -Return Prediction?
- It turns out that if we made all updates in an offline manner, then sum of updates for a fixed state  $s \in \mathcal{N}$  over entire trace experience equals (offline) update for s in the  $\lambda$ -Return prediction algorithm

#### Theorem

$$\sum_{t=0}^{T-1} \alpha \cdot \delta_t \cdot E_t(s) = \sum_{t=0}^{T-1} \alpha \cdot (G_t^{(\lambda)} - V(S_t)) \cdot \mathbb{I}_{S_t = s}, \text{ for all } s \in \mathcal{N}$$

$$G_{t}^{(\lambda)} = (1 - \lambda) \cdot \lambda^{0} \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}))$$

$$+ (1 - \lambda) \cdot \lambda^{1} \cdot (R_{t+1} + \gamma \cdot R_{t+2} + \gamma^{2} \cdot V(S_{t+2}))$$

$$+ (1 - \lambda) \cdot \lambda^{2} \cdot (R_{t+1} + \gamma \cdot R_{t+2} + \gamma^{2} \cdot R_{t+3} + \gamma^{3} \cdot V(S_{t+2}))$$

$$+ \dots$$

$$= (\gamma \lambda)^{0} \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - \gamma \lambda \cdot V(S_{t+1}))$$

$$+ (\gamma \lambda)^{1} \cdot (R_{t+2} + \gamma \cdot V(S_{t+2}) - \gamma \lambda \cdot V(S_{t+2}))$$

$$+ (\gamma \lambda)^{2} \cdot (R_{t+3} + \gamma \cdot V(S_{t+3}) - \gamma \lambda \cdot V(S_{t+3}))$$

$$+ \dots$$

$$G_t^{(\lambda)} = (\gamma \lambda)^0 \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - \gamma \lambda \cdot V(S_{t+1}))$$

$$+ (\gamma \lambda)^1 \cdot (R_{t+2} + \gamma \cdot V(S_{t+2}) - \gamma \lambda \cdot V(S_{t+2}))$$

$$+ (\gamma \lambda)^2 \cdot (R_{t+3} + \gamma \cdot V(S_{t+3}) - \gamma \lambda \cdot V(S_{t+3}))$$

$$+ \dots$$

$$G_{t}^{(\lambda)} - V(S_{t}) = (\gamma \lambda)^{0} \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}) - V(S_{t}))$$

$$+ (\gamma \lambda)^{1} \cdot (R_{t+2} + \gamma \cdot V(S_{t+2}) - V(S_{t+1}))$$

$$+ (\gamma \lambda)^{2} \cdot (R_{t+3} + \gamma \cdot V(S_{t+3}) - V(S_{t+2}))$$

$$+ \dots$$

$$= \delta_{t} + \gamma \lambda \cdot \delta_{t+1} + (\gamma \lambda)^{2} \cdot \delta_{t+2} + \dots$$

#### Proof

Now assume that a specific non-terminal state s appears at time steps  $t_1, t_2, \ldots, t_n$ . Then,

$$\sum_{t=0}^{T-1} \alpha \cdot (G_t^{(\lambda)} - V(S_t)) \cdot \mathbb{I}_{S_t = s} = \sum_{i=1}^{n} \alpha \cdot (G_{t_i}^{(\lambda)} - V(S_{t_i}))$$

$$= \sum_{i=1}^{n} \alpha \cdot (\delta_{t_i} + \gamma \lambda \cdot \delta_{t_i + 1} + (\gamma \lambda)^2 \cdot \delta_{t_i + 2} + \dots)$$

$$= \sum_{t=0}^{T-1} \alpha \cdot \delta_t \cdot E_t(s)$$

# TD(0) and TD(1) with Offline Updates

- To be clear,  $TD(\lambda)$  Prediction is an online algorithm
- So not the same as offline  $\lambda$ -Return Prediction
- If we modified  $TD(\lambda)$  to be offline, they'd be equivalent
- Offline version of  $TD(\lambda)$  would not update VF at each step
- Accumulate changes in buffer, update VF offline with buffer contents
- If we set  $\lambda = 0$ ,  $E_t(s) = \mathbb{I}_{S_t = s}$  and so, the update reduces to:

$$V(S_t) \leftarrow V(S_t) + \alpha \cdot \delta_t$$

- ullet This is exactly the TD update. So, TD is often refered to as TD(0)
- ullet If we set  $\lambda=1$  with episodic traces, sum of all VF updates for a state over a trace experience is equal to it's VF update in Every-Visit MC
- Hence, Offline TD(1) is equivalent to Every-Visit MC

# $\mathsf{TD}(\lambda)$ Prediction with Function Approximation

- Generalize  $TD(\lambda)$  to the case of function approximation
- Data-Type of eligibility traces same as func-approx parameters w
- ullet So here we denote eligibility traces at time t as simply  $oldsymbol{\mathcal{E}}_t$
- Initialize  $E_0$  to 0 for each component in it's data type
- For each time step t > 0,  $E_t$  is calculated recursively:

$$\mathbf{E}_t = \gamma \lambda \cdot \mathbf{E}_{t-1} + \nabla_{\mathbf{w}} V(S_t; \mathbf{w})$$

VF approximation update at each time step t is as follows:

$$\Delta \mathbf{w} = \alpha \cdot (R_{t+1} + \gamma \cdot V(S_{t+1}; \mathbf{w}) - V(S_t; \mathbf{w})) \cdot \mathbf{E}_t$$

ullet Expressed more succinctly in terms of function-approx TD-Error  $\delta_t$ :

$$\Delta \mathbf{w} = \alpha \cdot \delta_t \cdot \mathbf{E}_t$$

## Key Takeaways from this Chapter

- Bias-Variance tradeoff of TD versus MC
- MC learns the mean of the observed returns while TD learns something "deeper" - it implicitly estimates an MRP from given data and produces the Value Function of the implicitly-estimated MRP
- Understanding TD versus MC versus DP from the perspectives of:
  - "Bootstrapping"
  - "Experiencing"
- "Equivalence" of  $\lambda$ -Return Prediction and TD( $\lambda$ ) Prediction
- TD is equivalent to TD(0) and MC is "equivalent" to TD(1)