# Batch RL Via Least Squares Policy Iteration

Alan Fern

<sup>\*</sup> Based in part on slides by Ronald Parr

#### **Overview**

Motivation

- LSPI
  - Derivation from LSTD
  - Experimental results

#### Online versus Batch RL

- Online RL: integrates data collection and optimization
  - Select actions in environment and at the same time update parameters based on each observed (s,a,s',r)
- Batch RL: decouples data collection and optimization
  - First generate/collect experience in the environment giving a data set of state-action-reward-state pairs {(s<sub>i</sub>,a<sub>i</sub>,r<sub>i</sub>,s<sub>i</sub>')}
  - Use the fixed set of experience to optimize/learn a policy
- Online vs. Batch:
  - Batch algorithms are often more "data efficient" and stable
  - Batch algorithms ignore the exploration-exploitation problem, and do their best with the data they have

#### **Batch RL Motivation**

 There are many applications that naturally fit the batch RL model

#### • Medical Treatment Optimization:

- ▲ <u>Input:</u> collection of treatment episodes for an ailment giving sequence of observations and actions including outcomes
- Ouput: a treatment policy, ideally better than current practice

#### • Emergency Response Optimization:

- Input: collection of emergency response episodes giving movement of emergency resources before, during, and after 911 calls
- Output: emergency response policy

#### **LSPI**

- LSPI is a model-free batch RL algorithm
  - Learns a linear approximation of Q-function
  - stable and efficient
  - Never diverges or gives meaningless answers
- LSPI can be applied to a dataset regardless of how it was collected
  - But garbage in, garbage out.

<u>Least-Squares Policy Iteration</u>, Michail Lagoudakis and Ronald Parr, *Journal of Machine Learning Research (JMLR)*, Vol. 4, 2003, pp. 1107-1149.

#### **Notation**

- S: state space, s: individual states
- R(s,a): reward for taking action a in state s
- $\gamma$ : discount factor
- V(s): state value function
- P(s' | s,a) = T(s,a,s'): transition function
- Q(s,a): state-action value function
- Policy:  $\pi(s) \rightarrow a$

Objective: Maximize expected, total, discounted reward

$$\mathbf{E} \left[ \sum_{t=0}^{\infty} \gamma^t r_t \right]$$

## **Projection Approach to Approximation**

• Recall the standard Bellman equation:

$$V^{*}(s) = \max_{a} R(s, a) + \gamma \sum_{s'} P(s'|s, a) V^{*}(s')$$

or equivalently  $V^* = T[V^*]$  where T[.] is the Bellman operator

 Recall from value iteration, the sub-optimality of a value function can be bounded in terms of the Bellman error:

 $||V-T[V]||_{\infty}$ 

 This motivates trying to find an approximate value function with small Bellman error

## **Projection Approach to Approximation**

- Suppose that we have a space of representable value functions
  - ♠ E.g. the space of linear functions over given features
- Let  $\Pi$  be a *projection* operator for that space
  - Projects any value function (in or outside of the space) to "closest" value function in the space
- "Fixed Point" Bellman Equation with approximation

$$\hat{V}^* = \prod \left( T[\hat{V}^*] \right)$$

- Depending on space this will have a small Bellman error
- LSPI will attempt to arrive at such a value function
  - Assumes linear approximation and least-squares projection

## **Projected Value Iteration**

- Naïve Idea: try computing projected fixed point using VI
- Exact VI: (iterate Bellman backups)

$$V^{i+1} = T[V^i]$$

Projected VI: (iterated projected Bellman backups):

$$\hat{V}^{i+1} = \prod \left( T[\hat{V}^i] \right)$$

Projects exact Bellman backup to closest function in our restricted function space

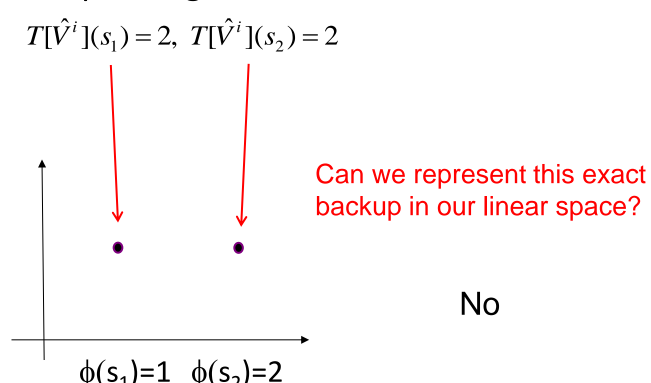
exact Bellman backup (produced value function)

## **Example: Projected Bellman Backup**

Restrict space to linear functions over a single feature  $\phi$ :

$$\hat{V}(s) = w \cdot \phi(s)$$

Suppose just two states  $s_1$  and  $s_2$  with:  $\phi(s_1)=1$ ,  $\phi(s_2)=2$  Suppose exact backup of  $V^i$  gives:



## **Example: Projected Bellman Backup**

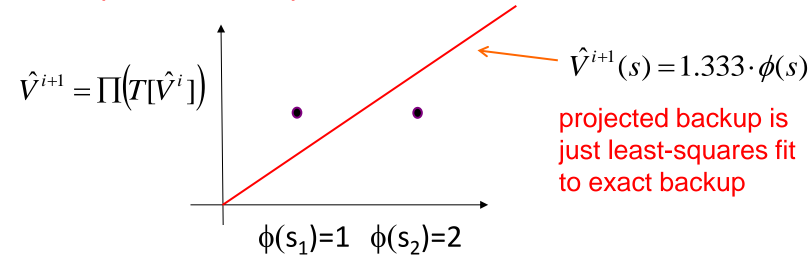
Restrict space to linear functions over a single feature  $\phi$ :

$$\hat{V}(s) = w \cdot \phi(s)$$

Suppose just two states  $s_1$  and  $s_2$  with:  $\phi(s_1)=1$ ,  $\phi(s_2)=2$  Suppose exact backup of  $V^i$  gives:

$$T[\hat{V}^i](s_1) = 2$$
,  $T[\hat{V}^i](s_2) = 2$ 

The backup can't be represented via our linear function:



## **Problem: Stability**

 Exact value iteration stability ensured by contraction property of Bellman backups:

$$V^{i+1} = T[V^i]$$

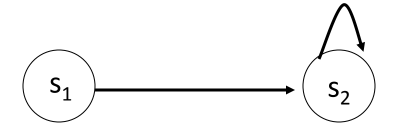
Is the "projected" Bellman backup a contraction:

$$\hat{V}^{i+1} = \prod \left( T[\hat{V}^i] \right)$$



#### **Example: Stability Problem** [Bertsekas & Tsitsiklis 1996]

<u>Problem</u>: Most projections lead to backups that are not contractions and unstable

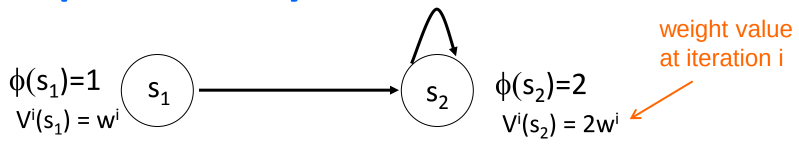


Rewards all zero, single action,  $\gamma = 0.9$ :  $V^* = 0$ 

Consider linear approx. w/ single feature  $\phi$  with weight w.

$$\hat{V}(s) = w \cdot \phi(s)$$
 Optimal  $w = 0$  since  $V^* = 0$ 

### **Example: Stability Problem**



From V<sup>i</sup> perform projected backup for each state

$$T[\hat{V}^i](s_1) = \gamma \hat{V}^i(s_2) = 1.8w^i$$

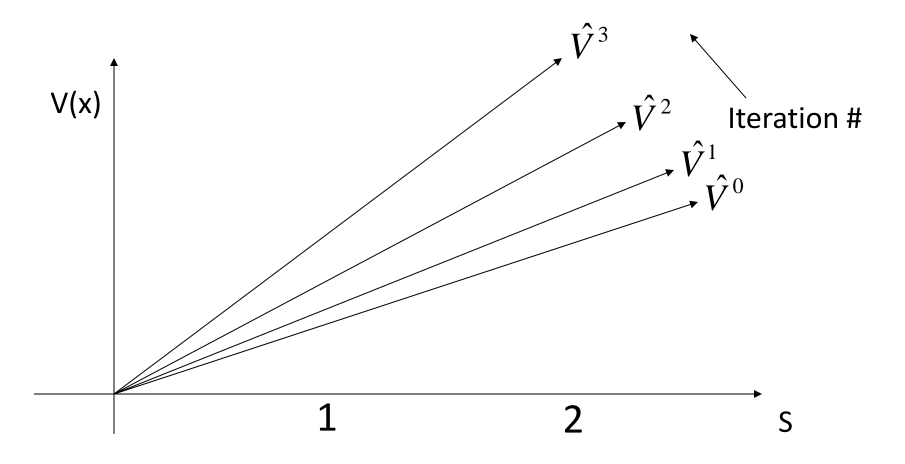
$$T[\hat{V}^i](s_2) = \gamma \hat{V}^i(s_2) = 1.8w^i$$

Can't be represented in our space so find wi+1 that gives least-squares approx. to exact backup

After some math we can get:  $w^{i+1} = 1.2 w^{i}$ 

What does this mean?

### **Example: Stability Problem**



Each iteration of Bellman backup makes approximation worse! Even for this trivial problem "projected" VI diverges.

### **Understanding the Problem**

- What went wrong?
  - Exact Bellman backups reduces error in max-norm
  - ▲ Least squares (= projection) non-expansive in L₂ norm
  - May increase max-norm distance

 Conclusion: Alternating value iteration and function approximation is risky business

#### **Overview**

Motivation

- LSPI
  - ◆ Derivation from Least-Squares Temporal Difference Learning
  - **▲** Experimental results

#### **How does LSPI fix these?**

- LSPI performs approximate policy iteration
  - PI involves policy evaluation and policy improvement
  - ◆ Uses a variant of least-squares temporal difference learning (LSTD) for approx. policy evaluation [Bratdke & Barto '96]

#### • Stability:

- ▲ LSTD directly solves for the fixed point of the approximate Bellman equation for policy values
- With singular-value decomposition (SVD), this is always well defined

#### Data efficiency

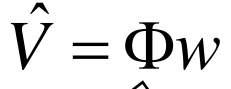
- ▲ LSTD finds best approximation for any finite data set
- Makes a single pass over the data for each policy
- Can be implemented incrementally

### OK, What's LSTD?

- Least Squares Temporal Difference Learning
- Assumes linear value function approximation of K features  $\hat{V}(s) = \sum_{k} w_{k} \phi_{k}(s)$
- The  $\phi_k$  are arbitrary feature functions of states
- Some vector notation

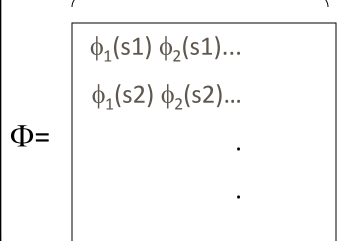
$$\hat{V} = \begin{bmatrix} \hat{V}(s_1) \\ \vdots \\ \hat{V}(s_n) \end{bmatrix} \quad w = \begin{bmatrix} w_1 \\ \vdots \\ w_k \end{bmatrix} \quad \phi_k = \begin{bmatrix} \phi_k(s_1) \\ \vdots \\ \phi_k(s_n) \end{bmatrix} \quad \Phi = \begin{bmatrix} \phi_1 & \cdots & \phi_K \end{bmatrix}$$

#### **Deriving LSTD**



 $V=\Phi w$  assigns a value to every state





# states

 $\hat{V}$  is a linear function in the column space of  $\phi_1...\phi_k$ , that is,

$$\hat{V} = w_1 \cdot \phi_1 + \dots + w_K \cdot \phi_K$$

## Suppose we know value of policy

• Want:  $\Phi_W pprox V^\pi$ 

Least squares weights minimizes squared error

$$w = (\Phi^T \Phi)^{-1} \Phi^T V^{\pi}$$

Sometimes called pseudoinverse

Least squares projection is then

$$\hat{V} = \Phi w = \Phi (\Phi^T \Phi)^{-1} \Phi^T V^{\pi}$$

Textbook least squares projection operator

#### But we don't know V...

Recall fixed-point equation for policies

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

• Will solve a projected fixed-point equation:

$$\hat{V}^{\pi} = \prod \left( R + \gamma P \hat{V}^{\pi} \right)$$

$$R = \begin{bmatrix} R(s_1, \pi(s_1)) \\ \vdots \\ R(s_n, \pi(s_n)) \end{bmatrix}, P = \begin{bmatrix} P(s_1 | s_1, \pi(s_1)) & \cdots & P(s_n | s_1, \pi(s_1)) \\ \vdots & \vdots & \vdots \\ P(s_1 | s_n, \pi(s_n)) & \cdots & P(s_1 | s_n, \pi(s_n)) \end{bmatrix}$$

Substituting least squares projection into this gives:

$$\Phi w = \Phi(\Phi^T \Phi)^{-1} \Phi^T (R + \gamma P \Phi w)$$

• Solving for w:  $w = (\Phi^T \Phi - \gamma \Phi^T P \Phi)^{-1} \Phi^T R$ 

#### Almost there...

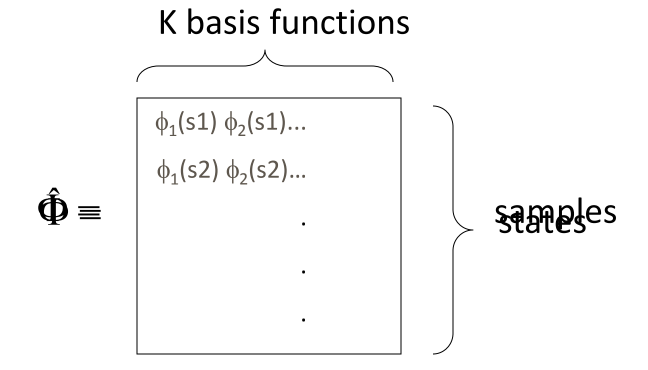
$$w = (\Phi^T \Phi - \gamma \Phi^T P \Phi)^{-1} \Phi^T R$$

- Matrix to invert is only K x K
- But...
  - ♠ Expensive to construct matrix (e.g. P is |S|x|S|)
  - We don't know P
  - We don't know R

## Using Samples for $\Phi$

Suppose we have state transition samples of the policy running in the MDP:  $\{(s_i,a_i,r_i,s_i')\}$ 

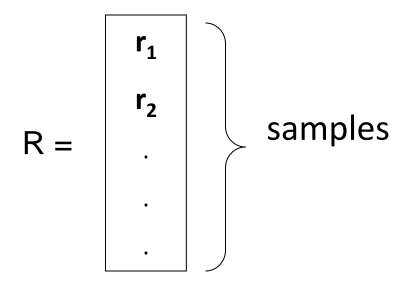
Idea: Replace enumeration of states with sampled states



## **Using Samples for R**

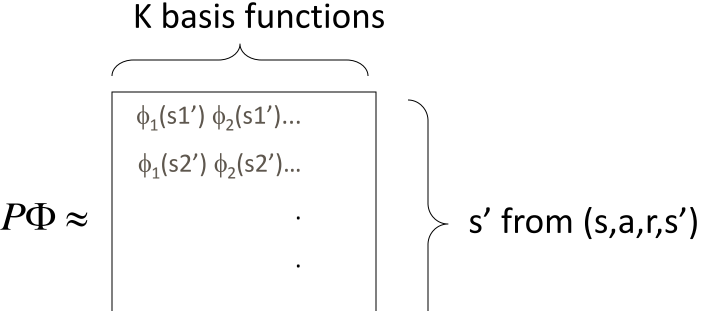
Suppose we have state transition samples of the policy running in the MDP:  $\{(s_i,a_i,r_i,s_i')\}$ 

Idea: Replace enumeration of reward with sampled rewards



## Using Samples for P $\Phi$

Idea: Replace expectation over next states with sampled next states.



## **Putting it Together**

LSTD needs to compute:

$$w = (\Phi^T \Phi - \gamma \Phi^T P \Phi)^{-1} \Phi^T R = B^{-1} b$$

$$B = \Phi^T \Phi - \gamma \Phi^T (P \Phi)$$

$$b = \Phi^T R$$
from previous slide

- The hard part of which is B the kxk matrix:
- Both B and b can be computed incrementally for each (s,a,r,s') sample: (initialize to zero)

$$B_{ij} \leftarrow B_{ij} + \phi_i(s)\phi_j(s) - \gamma\phi_i(s)\phi_j(s')$$
$$b_i \leftarrow b_i + r \cdot \phi_i(s)$$

# **LSTD Algorithm**

- Collect data by executing trajectories of current policy
- For each (s,a,r,s') sample:

$$B_{ij} \leftarrow B_{ij} + \phi_i(s)\phi_j(s) - \gamma\phi_i(s)\phi_j(s')$$
$$b_i \leftarrow b_i + r \cdot \phi_i(s, a)$$

$$w \leftarrow B^{-1}b$$

## **LSTD Summary**

- Does O(k²) work per datum
  - Linear in amount of data.
- Approaches model-based answer in limit
- Finding fixed point requires inverting matrix

- Fixed point almost always exists
- Stable; efficient

## **Approximate Policy Iteration with LSTD**

**Policy Iteration:** iterates between policy improvement and policy evaluation

Idea: use LSTD for approximate policy evaluation in PI

Start with random weights w (i.e. value function)

Repeat Until Convergence

$$\pi(s) = \operatorname{greedy}(\hat{V}(s, \mathbf{w}))$$
 // policy improvement

Evaluate  $\pi$  using LSTD

- lacktriangle Generate sample trajectories of  $\pi$
- Use LSTD to produce new weights **w** (**w** gives an approx. value function of  $\pi$ )

#### What Breaks?

- No way to execute greedy policy without a model
- Approximation is biased by current policy
  - We only approximate values of states we see when executing the current policy
  - ▲ LSTD is a weighted approximation toward those states
- Can result in Learn-forget cycle of policy iteration
  - Drive off the road; learn that it's bad
  - New policy never does this; forgets that it's bad
- Not truly a batch method
  - Data must be collected from current policy for LSTD

#### **LSPI**

- LSPI is similar to previous loop but replaces LSTD with a new algorithm LSTDQ
- LSTD: produces a value function
  - Requires sample from policy under consideration
- LSTDQ: produces a Q-function for current policy
  - ◆ Can learn Q-function for policy from any (reasonable) set of samples---sometimes called an off-policy method
  - No need to collect samples from current policy
- Disconnects policy evaluation from data collection
  - Permits reuse of data across iterations!
  - ▲ Truly a batch method.

# Implementing LSTDQ

- Both LSTD and LSTDQ compute:  $B = \Phi^T \Phi \lambda \Phi^T (P\Phi)$
- But LSTDQ basis functions are indexed by actions

$$\hat{Q}_w(s,a) = \sum_k w_k \cdot \phi_k(s,a)$$
 defines greedy policy:  $\pi_w(s) = \arg\max_a \hat{Q}_w(s,a)$ 

• For each (s,a,r,s') sample:

$$B_{ij} \leftarrow B_{ij} + \phi_i(s, a)\phi_j(s, a) - \lambda\phi_i(s, a)\phi_j(s', \pi_w(s'))$$

$$b_i \leftarrow b_i + r \cdot \phi_i(s, a)$$

$$w \leftarrow B^{-1}b$$

$$arg \max_a \hat{Q}_w(s', a)$$

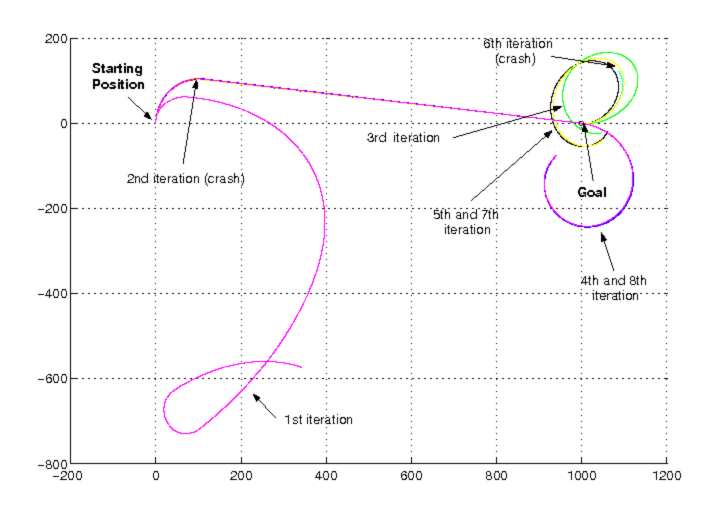
### **Running LSPI**

- There is a Matlab implementation available!
- Collect a database of (s,a,r,s') experiences (this is the magic step)
- Start with random weights (= random policy)
- 3. Repeat
  - Evaluate current policy against database
    - Run LSTDQ to generate new set of weights
    - New weights imply new Q-function and hence new policy
  - Replace current weights with new weights
- Until convergence

## **Results: Bicycle Riding**

- Watch random controller operate bike
- Collect ~40,000 (s,a,r,s') samples
- Pick 20 simple feature functions (×5 actions)
- Make 5-10 passes over data (PI steps)
- Reward was based on distance to goal + goal achievement
- Result: Controller that balances and rides to goal

# **Bicycle Trajectories**

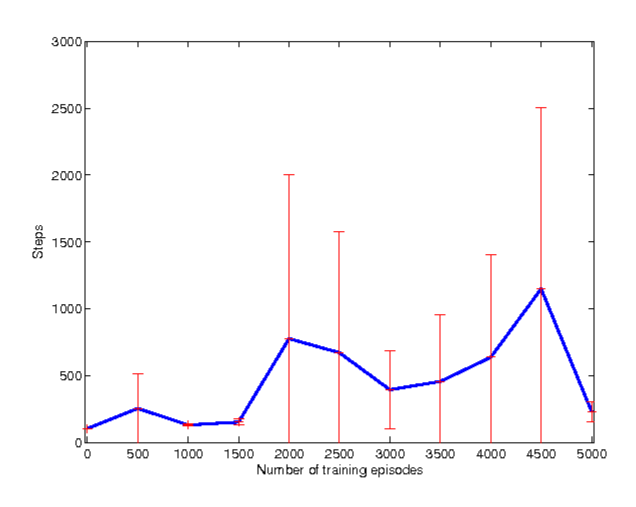


# What about Q-learning?

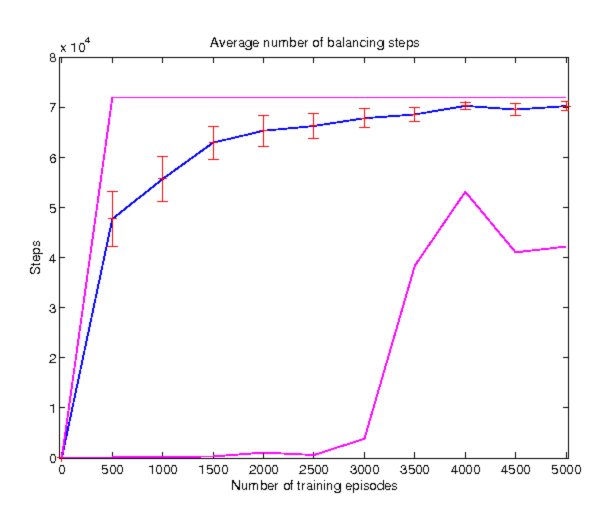
Ran Q-learning with same features

Used experience replay for data efficiency

# **Q-learning Results**



#### **LSPI Robustness**



## Some key points

- LSPI is a batch RL algorithm
  - Can generate trajectory data anyway you want
  - Induces a policy based on global optimization over full dataset
- Very stable with no parameters that need tweaking

#### So, what's the bad news?

- LSPI does not address the exploration problem
  - ▲ It decouples data collection from policy optimization
  - ↑ This is often not a major issue, but can be in some cases
- k<sup>2</sup> can sometimes be big
  - Lots of storage
  - Matrix inversion can be expensive
- Bicycle needed "shaping" rewards
- Still haven't solved
  - Feature selection (issue for all machine learning, but RL seems even more sensitive)