

Reinforcement learning

Master CPS, Year 2 Semester 1

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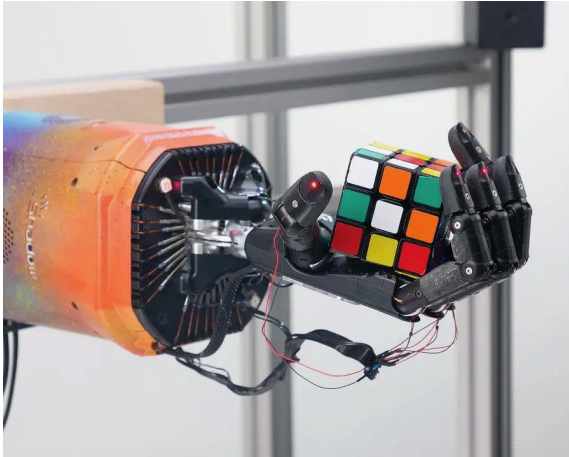


Part IV

Approximate dynamic programming and offline approximate reinforcement learning

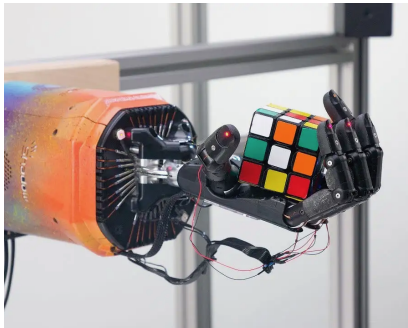


RL for manipulation of a Rubik's Cube (OpenAI)



The need for approximation

- Classical RL – representation in tabular form, e.g., $Q(x, u)$ separately for all values of x and u
- In real applications, x , u often **continuous** (or discrete with very many values)!



- **Tabular representation is impossible**

The need for approximation (continued)

In real applications, the functions of interest must often be
approximated



Part IV in plan

- Reinforcement learning problem
- Optimal solution
- Exact dynamic programming
- Exact reinforcement learning
- **Approximation techniques**
- **Approximate dynamic programming**
- **Offline approximate reinforcement learning**
- Online approximate reinforcement learning



Contents of part IV

- 1 Approximation techniques
- 2 Approximation in DP&RL
- 3 Q-iteration with interpolation
- 4 Fitted Q-iteration



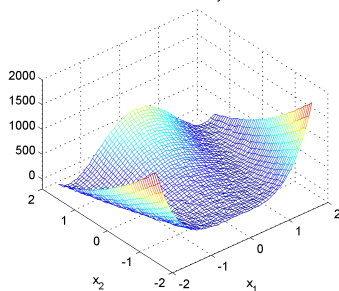
Approximation

Approximation:

a function with (uncountably) infinitely many values must be represented using a small number of values

 $f(x)$

True values y

 $\hat{f}(x)$

?

Parametric approximation

Parametric approximation: $\hat{f}$ has a fixed form,
its output is determined by a vector of **parameters** θ :

$$\hat{f}(x; \theta)$$

- 1 **Linear approximation** – weighted combination of **features** (basis functions) ϕ_i :

$$\begin{aligned}\hat{f}(x; \theta) &= \phi_1(x)\theta_1 + \phi_2(x)\theta_2 + \dots \phi_n(x)\theta_n \\ &= \sum_{i=1}^n \phi_i(x)\theta_i = \phi^\top(x)\theta\end{aligned}$$

Note: linear in parameters, can be nonlinear in x

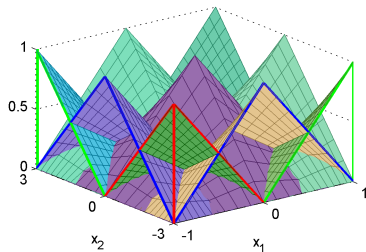
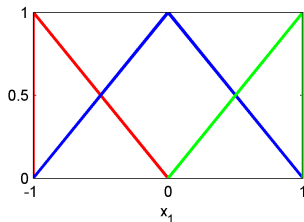
- 2 **Nonlinear approximation:** stays in the general form



Linear parametric approximation: Interpolation

Interpolation:

- D-dimensional grid of points
- Multilinear interpolation between points
- Equivalent to **pyramidal** features



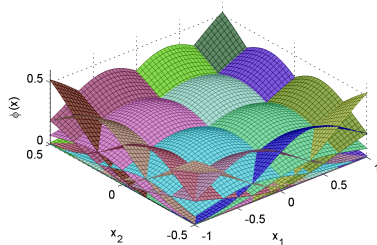
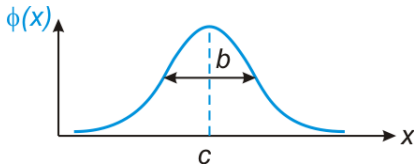
Linear parametric approximation: RBF

Radial basis function (Gaussian):

$$\phi(x) = \exp \left[-\frac{(x - c)^2}{b^2} \right] \quad (1\text{-dim});$$

$$= \exp \left[-\sum_{d=1}^D \frac{(x_d - c_d)^2}{b_d^2} \right] \quad (D\text{-dim})$$

Optionally, normalization: $\tilde{\phi}_i(x) = \frac{\phi_i(x)}{\sum_{i' \neq i} \phi_{i'}(x)}$



Training linear approximators: least squares

- n_s points $(x_j, f(x_j))$, objective described by system of equations:

$$\hat{f}(x_1; \theta) = \phi_1(x_1)\theta_1 + \phi_2(x_1)\theta_2 + \dots \phi_n(x_1)\theta_n = f(x_1)$$

...

$$\hat{f}(x_{n_s}; \theta) = \phi_1(x_{n_s})\theta_1 + \phi_2(x_{n_s})\theta_2 + \dots \phi_n(x_{n_s})\theta_n = f(x_{n_s})$$

- Matrix form:

$$\begin{bmatrix} \phi_1(x_1) & \phi_2(x_1) & \dots & \phi_n(x_1) \\ \dots & \dots & \dots & \dots \\ \phi_1(x_{n_s}) & \phi_2(x_{n_s}) & \dots & \phi_n(x_{n_s}) \end{bmatrix} \cdot \theta = \begin{bmatrix} f(x_1) \\ \dots \\ f(x_{n_s}) \end{bmatrix} \quad A\theta = b$$

- Linear regression



Least squares (continued)

- Overdetermined system ($n_s > n$), equations cannot all be satisfied with equality.
⇒ Solve in the least-squares sense:

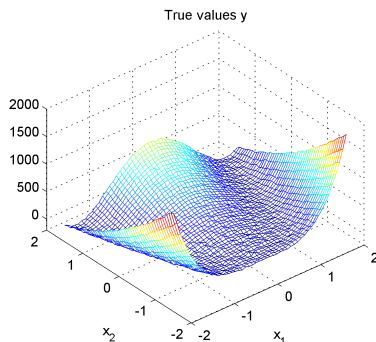
$$\min_{\theta} \sum_{j=1}^{n_s} \left[f(x_j) - \hat{f}(x_j; \theta) \right]^2$$

...linear algebra and analysis...

- $\theta = (A^{\top} A)^{-1} A^{\top} b \quad (\Leftarrow (A^{\top} A)\theta = A^{\top} b)$



Example: Rosenbrock's “banana” function

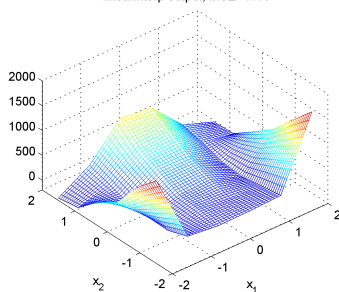


- $f(x) = (1 - x_1)^2 + 100[(x_2 + 1.5) - x_1^2]^2$, $x = [x_1, x_2]^T$
- **Training:** 200 points, uniformly randomly distributed
- **Validation:** 31×31 point grid

Rosenbrock function: Linear approximator results

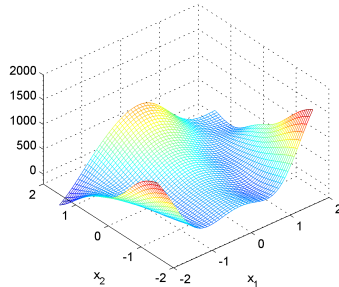
Interpolation on a 6x6 grid:

linearinterp output; MSE=4175



6x6 RBFs:

RBFs output; MSE=5399

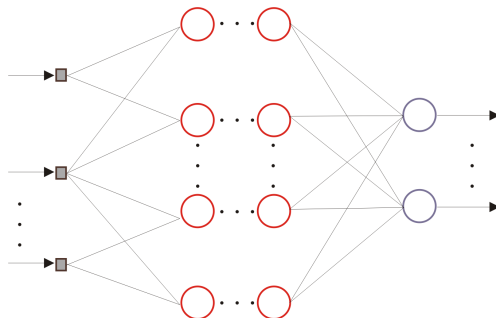


- Interpolation = collection of multilinear surfaces
- RBF approximation is smoother (wide RBFs)

Nonlinear parametric approximation: neural network

Neural network:

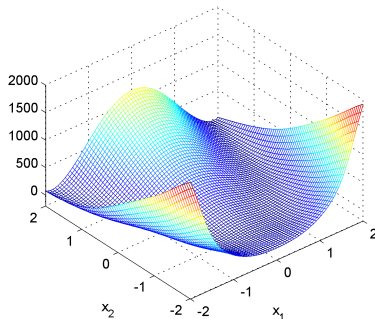
- **Neurons** with (non)linear activation functions
- **Interconnected** in multiple layers, through weighted connections + biases



Rosenbrock function: Neural network result

One hidden layer with 10 neurons and tangent-sigmoid activation functions + linear output layer. 500 training epochs.

NN output, MSE=300.83



Thanks to the greater flexibility of the neural network, the results are better than those with linear approximators.

Neural network to find features

Usually, the last layer of a neural network is linear, leading again to the feature-based approximator:

$$\hat{f}(x; \theta) = \phi^\top(x; \theta_{\text{feat}}) \theta_{\text{lin}}$$

where $\theta = [\theta_{\text{feat}}^\top, \theta_{\text{lin}}^\top]^\top$.

Key difference from linear approximation: features ϕ are now themselves parametrized – by θ_{feat} – and **automatically found by the neural network**. In linear approximation, the features are manually designed.

Note that overall the approximator is nonlinear!



Neural network features

Example: Convolutional neural net features:



Details to follow later (with Florin)

Comparison between approximators

- Linear approximators are easier to handle theoretically than nonlinear ones
- Nonlinear approximators are more flexible than linear ones



- 1 Approximation techniques
- 2 Approximation in DP&RL**
- 3 Q-iteration with interpolation
- 4 Fitted Q-iteration



Approximation in RL

Problems to be solved:

- 1 **Representation:** $Q(x, u)$, $V(x)$, $h(x)$
Using the approximation techniques discussed earlier
- 2 **Maximization:** e.g., $\max_u Q(x, u)$



Option 1: h implicit

- The policy is implicitly represented...
- ...by computing greedy actions on demand from $\hat{Q}$:

$$h(x) = \arg \max_u \hat{Q}(x, u)$$

- ⇒ Main problem: **approximating the Q-function**
- Approximator must ensure **efficient solution for** $\arg \max$
 - Will be the focus in this lecture



Option 2: h explicit

- The policy is explicitly approximated: $\hat{h}(x)$

Advantages:

- **Continuous actions** are easier to handle
- The representation can more easily incorporate **prior knowledge**



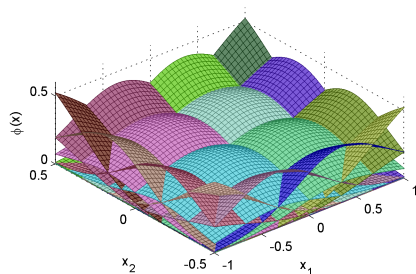
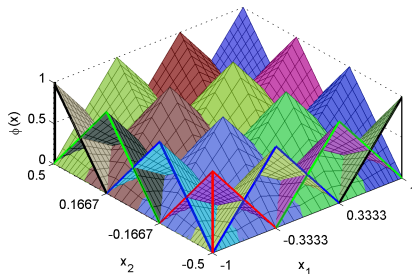
Action discretization

- Approximator must ensure efficient solution for $\arg \max$
- ⇒ Typically: **discretize actions**
- Select M discrete actions $u_1, \dots, u_j, \dots, u_M \in U$
Compute “arg max” using explicit enumeration
- Example: discretization on a grid



State-space approximation

- Often features $\phi_1, \dots, \phi_N : X \rightarrow [0, \infty)$
- Ex. **pyramidal**, **RBF**



or perhaps found by neural network

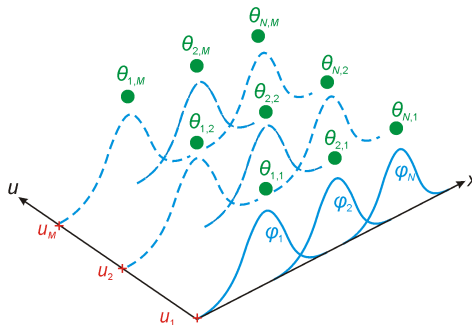
Approximating Q with discrete actions

Given:

- 1 N features $\phi_1, \dots, \phi_N$
- 2 M discrete actions $u_1, \dots, u_M$

Store:

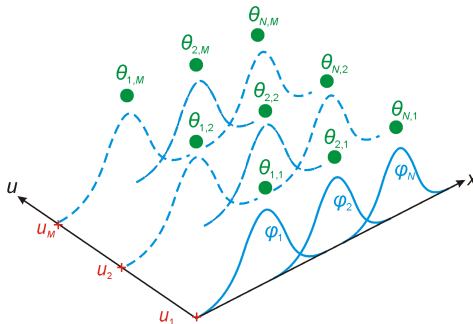
- 3 $N \cdot M$ parameters θ
(for each feature–discrete action pair)



Approximating Q with discrete actions (continued)

Approximate Q -function:

$$\hat{Q}(x, u_j; \theta) = \sum_{i=1}^N \phi_i(x) \theta_{i,j} = [\phi_1(x) \dots \phi_N(x)] \begin{bmatrix} \theta_{1,j} \\ \vdots \\ \theta_{N,j} \end{bmatrix}$$

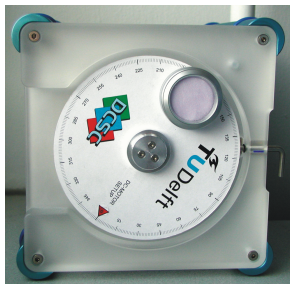


Benefit of approximation in RL

Approximation allows applying RL
to realistic problems



Simple control example: Inverted pendulum



- State $x = [\alpha, \dot{\alpha}]^\top$ with angle $\alpha \in [-\pi, \pi)$ rad, velocity $\dot{\alpha} \in [-15\pi, 15\pi]$ rad/s
- Action $u \in [-3, 3]$ V
- Dynamics:

$$\ddot{\alpha} = 1/J \cdot \left[mgl \sin(\alpha) - (b + \frac{K^2}{R})\dot{\alpha} + \frac{K}{R}u \right]$$

- **Objective:** stabilize pointing up, encoded by reward:

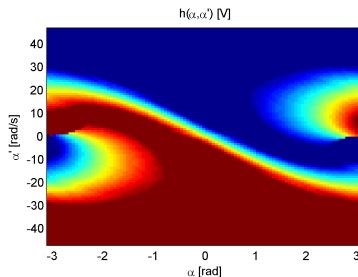
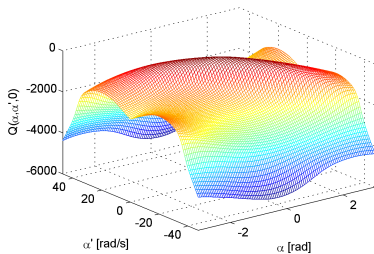
$$\rho(x, u) = -5\alpha^2 - 0.1\dot{\alpha}^2 - u^2$$

normalized to $[0, 1]$

- Discount factor $\gamma = 0.98$
- Insufficient power $\Rightarrow$ swing back & forth before stabilizing

Inverted pendulum: Optimal solution

Left: Q-function for $u = 0$ Right: policy



► Replay

New questions raised by approximation

- 1 **Convergence**: does the algorithm remain convergent?
- 2 **Near-optimality**: is the solution at a controlled distance from the optimum?
- 3 **Consistency**: for an ideal approximator with infinite precision, is the optimal solution recovered?



Algorithm landscape

By model usage:

- **Model-based**: f, ρ known a priori
- **Model-free**: f, ρ unknown (reinforcement learning)

By interaction level:

- **Offline**: algorithm runs in advance
- **Online**: algorithm runs with the system

Exact vs. approximate:

- **Exact**: x, u small number of discrete values
- **Approximate**: x, u continuous (or many discrete values)

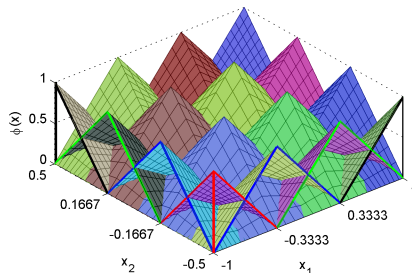


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- 4 Fitted Q-iteration



Q-function approximator

- Interpolation = pyramidal features



- Each feature i has center x_i
- $\theta_{i,j}$ can be viewed as $\hat{Q}(x_i, u_j)$, since: $\phi_i(x_i) = 1, \phi_{i'}(x_i) = 0$ for $i' \neq i$

Q-iteration with interpolation

Recall classical Q-iteration:

repeat at each iteration ℓ

for all x, u **do**

$$Q_{\ell+1}(x, u) \leftarrow \rho(x, u) + \gamma \max_{u'} Q_{\ell}(f(x, u), u')$$

end for

until convergence

Q-iteration with interpolation

repeat at each iteration ℓ

for all centers x_i , discrete actions u_j **do**

$$\theta_{\ell+1,i,j} \leftarrow \rho(x_i, u_j) + \gamma \max_{j'} \hat{Q}(f(x_i, u_j), u_{j'}; \theta_{\ell})$$

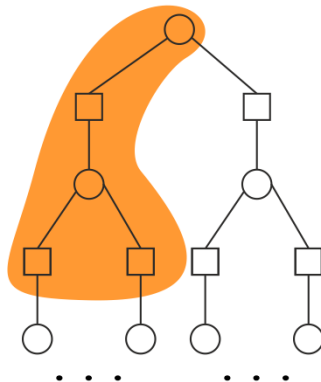
end for

until convergence

Stochastic version exists, but here we only consider the deterministic case



Illustration



Policy

- Recall the optimal policy:

$$h^*(x) = \underset{u}{\text{arg max}} \, Q^*(x, u)$$

- When Q is approximated using action discretization (e.g. the interpolating approximator above):

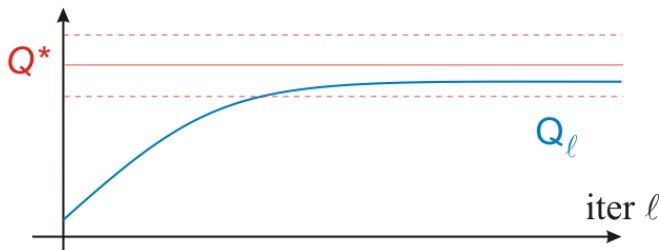
$$\hat{h}^*(x) = \underset{u_j, j=1, \dots, M}{\text{arg max}} \, \hat{Q}(x, u_j; \theta^*)$$

θ^* = parameters at convergence



Q-iteration w/ interpolation: Illustration of properties

Monotonic convergence to a **near-optimal solution**



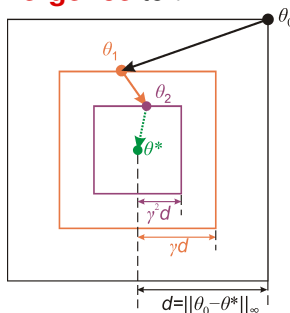
Convergence

Similar to classical Q-iteration:

- Each iteration is a contraction with factor γ :

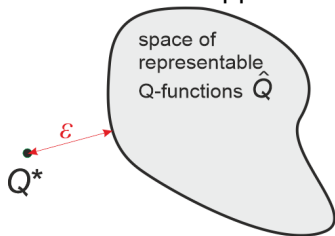
$$\|\theta_{\ell+1} - \theta^*\|_{\infty} \leq \gamma \|\theta_{\ell} - \theta^*\|_{\infty}$$

⇒ **Monotonic convergence** to θ^*



Near-optimality

Characterize approximator by the minimum distance to Q^* :



$$\epsilon = \min_{\theta} \left\| Q^*(x, u) - \hat{Q}(x, u; \theta) \right\|_{\infty}$$

Then:

- 1 Suboptimality of resulting $\hat{Q}(x, u; \theta^*)$ is bounded:

$$\left\| Q^*(x, u) - \hat{Q}(x, u; \theta^*) \right\|_{\infty} \leq \frac{2\epsilon}{1 - \gamma}$$

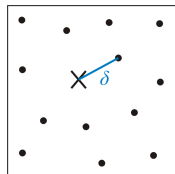
- 2 Suboptimality of policy $\hat{h}^*$ is bounded: by

$$\left\| Q^*(x, u) - Q^{\hat{h}^*}(x, u) \right\|_{\infty} \leq \frac{4\epsilon}{(1 - \gamma)^2}$$

Consistency

- Consistency: $\hat{Q}^{\theta^*} \rightarrow Q^*$ as resolution increases

- Resolution:
$$\begin{cases} \delta_x = \max_x \min_i \|x - x_i\|_2 \\ \delta_u = \max_u \min_j \|u - u_j\|_2 \end{cases}$$

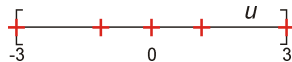
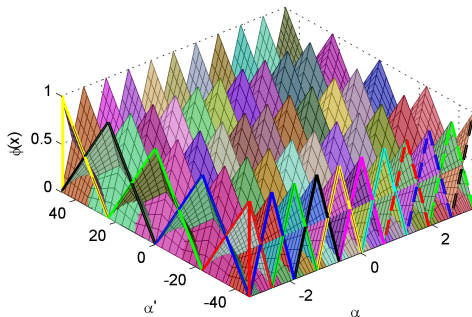


- Given certain technical conditions,
 $\Rightarrow \lim_{\delta_x \rightarrow 0, \delta_u \rightarrow 0} \hat{Q}^{\theta^*} = Q^*$ — consistency

Inverted pendulum: Q-iteration w/ interpolation, demo

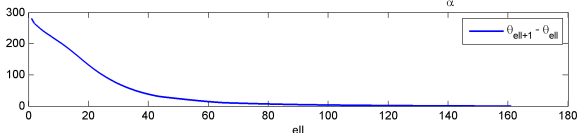
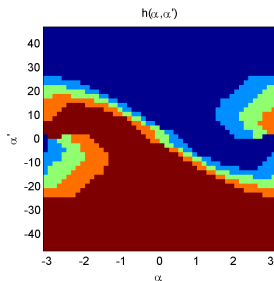
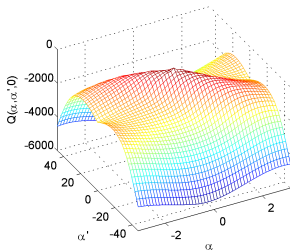
Features: equidistant grid 41×21

Discretization: 5 actions, distributed around 0



Inverted pendulum: Q-iteration w/ interpolation, demo

Fuzzy Q-iteration, ell=161



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Fitted Q-iteration

Extend Q-iteration with interpolation so that it works:

- with other approximators
- **model-free** – RL



Intermediate model-based algorithm

Recall Q-iteration with interpolation:

for all x_i, u_j $\theta_{\ell+1,i,j} \leftarrow \rho(x_i, u_j) + \gamma \max_{j'} \hat{Q}(f(x_i, u_j), u_{j'}; \theta_\ell)$ **end for**

- 1 Use **arbitrary** state-action samples
- 2 Extend to **generic approximator**
- 3 Find parameters using **least-squares**

get state-action samples (x_s, u_s) , $s = 1, \dots, n_s$

repeat at each iteration ℓ

for $s = 1, \dots, n_s$ **do**

 compute bootstrapped target for $\hat{Q}(x_s, u_s; \theta)$:

$\hat{R}_s \leftarrow \rho(x_s, u_s) + \gamma \max_{u'} \hat{Q}(f(x_s, u_s), u'; \theta_\ell)$

end for

$\theta_{\ell+1} \leftarrow \arg \min \sum_{s=1}^{n_s} \left[\hat{R}_s - \hat{Q}(x_s, u_s; \theta) \right]^2$

until termination

Q-iteration with interpolation is equivalent to this generalized algorithm if samples = all combinations x_i, u_j



Fitted Q-iteration: Algorithm

- ④ Use **transitions** instead of model

Fitted Q-iteration

get or collect dataset $\mathcal{D} = \{(x_s, u_s, r_s, x'_s), s = 1, \dots, n_s\}$

repeat at each iteration ℓ

for $s = 1, \dots, n_s$ **do**

 compute bootstrapped target for $\hat{Q}(x_s, u_s; \theta)$:

$$\hat{R}_s \leftarrow r_s + \gamma \max_{u'} \hat{Q}(x'_s, u'; \theta_\ell)$$

end for

$$\theta_{\ell+1} \leftarrow \arg \min \sum_{s=1}^{n_s} [\hat{R}_s - \hat{Q}(x_s, u_s; \theta)]^2 =: \mathcal{L}(\theta)$$

until termination

$\mathcal{L}$ is called the **loss function**



Deterministic versus stochastic

- In the deterministic case, $x'_s = f(x_s, u_s)$, $r_s = \rho(x_s, u_s)$
– substitutions are exact
- In the stochastic case, $x'_s \sim \tilde{f}(x_s, u_s, \cdot)$, $r_s = \tilde{\rho}(x_s, u_s, x'_s)$

⇒ Algorithm **remains valid**; intuition:

- Ideally, $Q(x, u) \leftarrow E_{x'} \left\{ r + \gamma \max_{u'} \hat{Q}(x', u'; \theta_\ell) \right\}$
- Assuming for the moment all samples are at $(x_s, u_s) = (x, u)$,

$$\min_{\theta} \sum_{s=1}^{n_s} \left| r_s + \gamma \max_{u'} \hat{Q}(x'_s, u'; \theta_\ell) - \hat{Q}(x, u; \theta) \right|^2$$

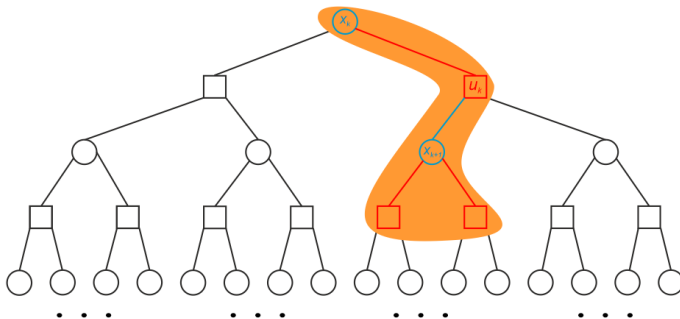
leads to $\hat{Q}(x, u; \theta) \approx E \{ \dots \}$ (like in Monte-Carlo)

- Even if (x_s, u_s) do not repeat, least-squares still approximate the expected value



Illustration

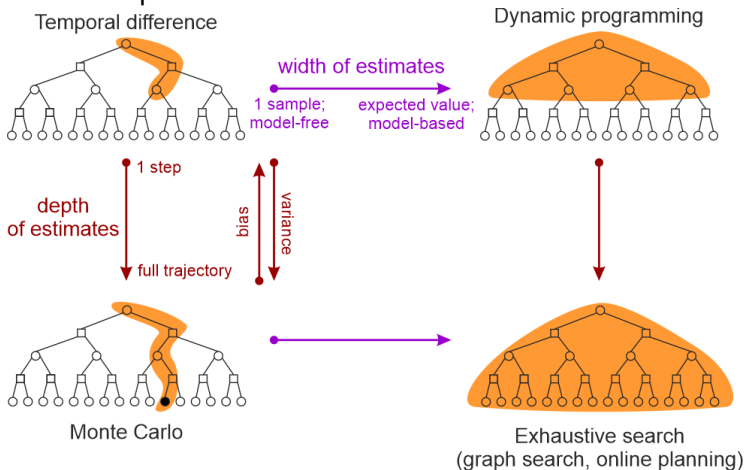
Targets are bootstrapping estimates for Q^* :



and therefore related to temporal differences.

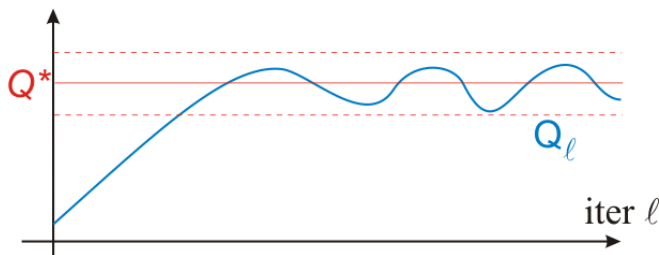
Fitted Q-iteration in the unified perspective

Belongs in the TD corner, but similar to DP in that it updates synchronously the complete Q-function across the entire state-action space.



Fitted Q-iteration: Illustration of properties

Convergence to a **sequence** of solutions,
each of them **near-optimal**

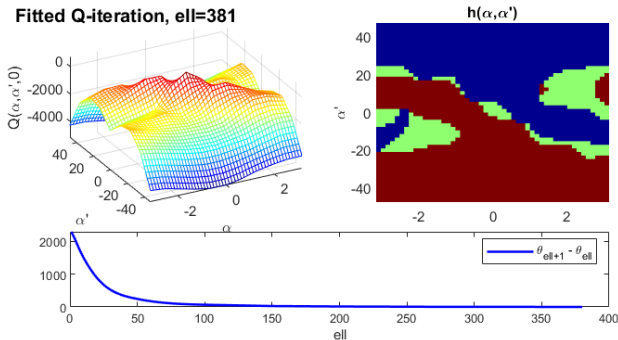


Inv. pend.: Fitted Q-iteration w/ interpolation, demo

Features: equidistant grid 11×9

Action discretization: 3 values, +/- max voltage and 0

Dataset: 10000 samples, uniformly distributed in the continuous x - discretized u space



Key terms in this part

- function approximation
- features / basis functions and parameters
- linear and nonlinear approximation
- interpolation, radial basis functions, neural network
- action discretization
- approximate dynamic programming
- fitted Q-iteration
- convergence, near-optimality, consistency



Exercises

- 1 Prove that the updates of Q-iteration with interpolation:

$$\theta_{\ell+1,i,j} \leftarrow \rho(x_i, u_j) + \gamma \max_{j'} \hat{Q}(f(x_i, u_j), u_{j'}; \theta_\ell)$$

are contractive with factor γ . Hint: it is essential that the features sum up to 1, and therefore the approximate Q-value is an average of the parameters!

- 2 What is the limit of the distance ε between Q^* and the space of representable Q-functions when $\delta_x \rightarrow 0, \delta_u \rightarrow 0$? Explain why.
- 3 Write an approximate V-iteration method with interpolation and prove that its updates are contractive.
- 4 Write a fitted V-iteration method. Is this method model-free?

