

Deep Reinforcement Learning

Function approximation

Julien Vitay

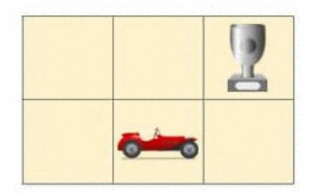
Professur für Künstliche Intelligenz - Fakultät für Informatik

1 - Limits of tabular RL

Tabular reinforcement learning

- All the methods seen so far belong to **tabular RL**.
- Q-learning necessitates to store in a **Q-table** one Q-value per state-action pair (s,a).

Game Board:



Current state (s): 0 0 0

Q Table:

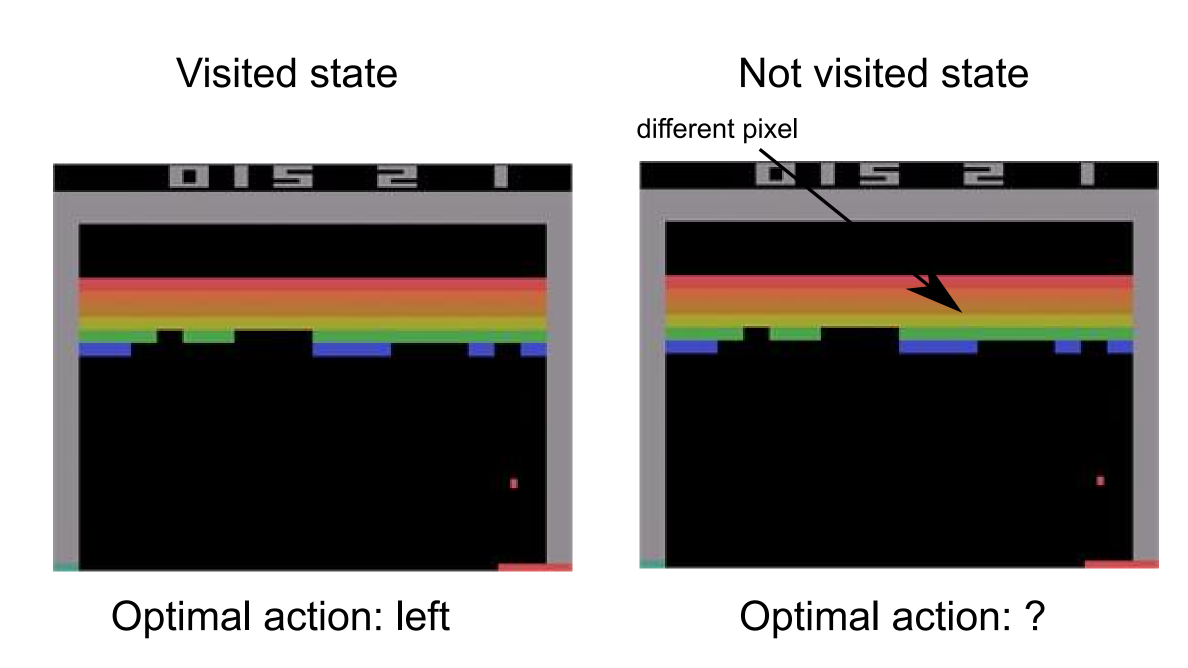
	000	0 0 0 0 1 0	0 0 0 0 0 1	100	0 1 0 0 0 0	0 0 1 0 0 0
Î	0.2	0.3	1.0	-0.22	-0.3	0.0
Ţ	-0.5	-0.4	-0.2	-0.04	-0.02	0.0
\Rightarrow	0.21	0.4	-0.3	0.5	1.0	0.0
	-0.6	-0.1	-0.1	-0.31	-0.01	0.0

y = 0.95

Source: https://towardsdatascience.com/qrash-course-deep-q-networks-from-the-ground-up-1bbda41d3677

Tabular reinforcement learning

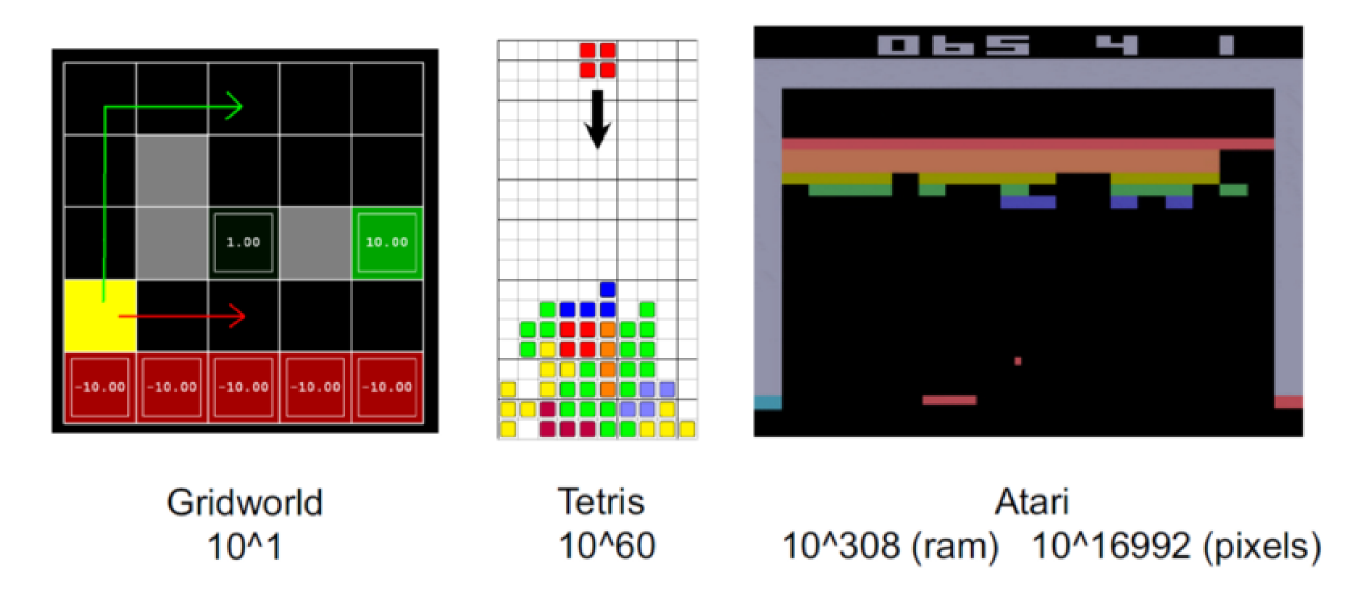
• If a state has never been visited during learning, the Q-values will still be at their initial value (0.0), no policy can be derived.



• Similar states likely have the same optimal action: we want to be able to **generalize** the policy between states.

Tabular reinforcement learning

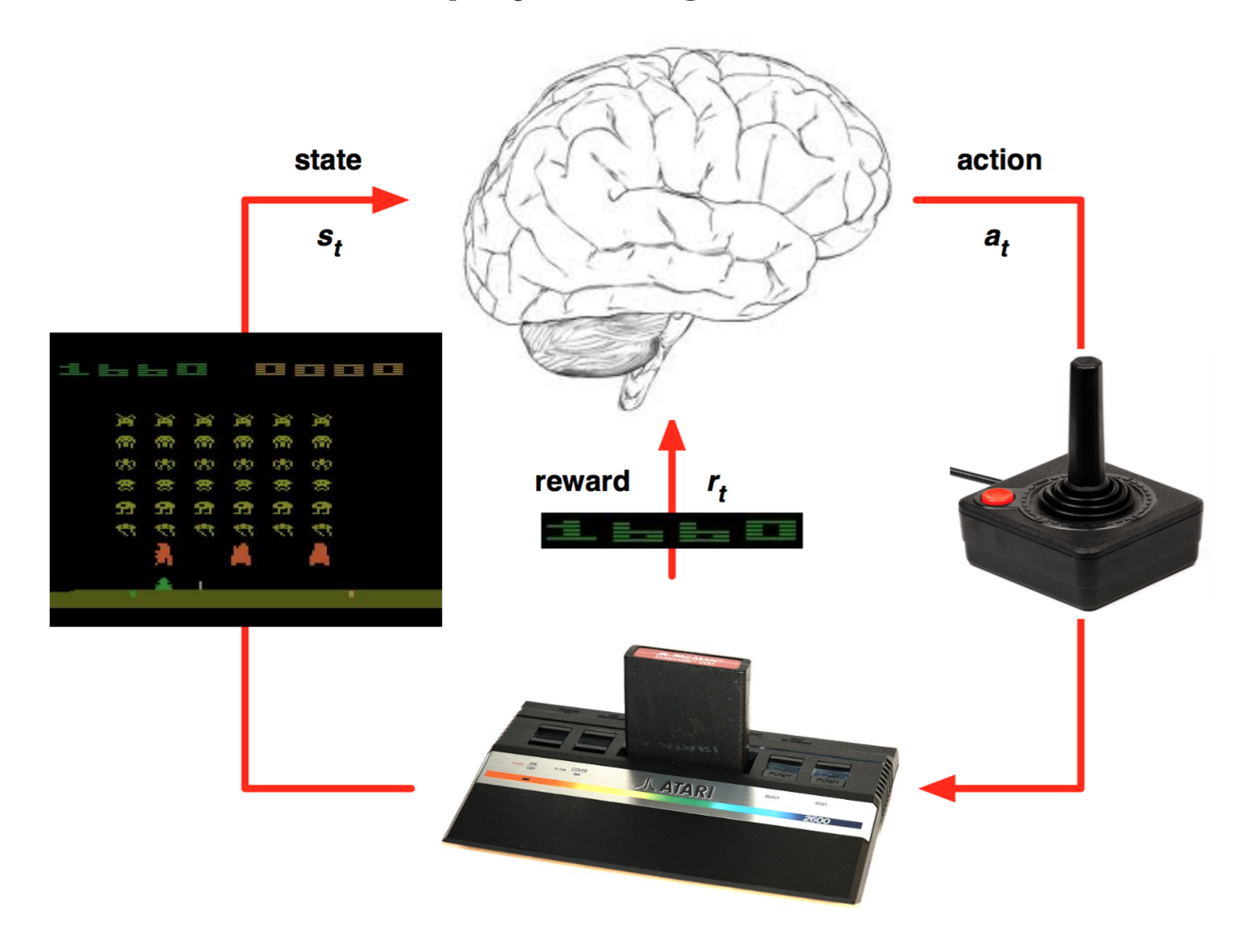
• For most realistic problems, the size of the Q-table becomes quickly untractable.



Source: https://medium.com/@twt446/a-summary-of-deep-reinforcement-learning-rl-bootcamp-lecture-2-c3a15db5934e

- ullet If you use black-and-white 256x256 images as inputs, you have $2^{256*256}=10^{19728}$ possible states!
- Tabular RL is limited to toy problems.

Tabular RL cannot learn to play video games



Source: David Silver. http://www0.cs.ucl.ac.uk/staff/d.silver/web/Teaching.html

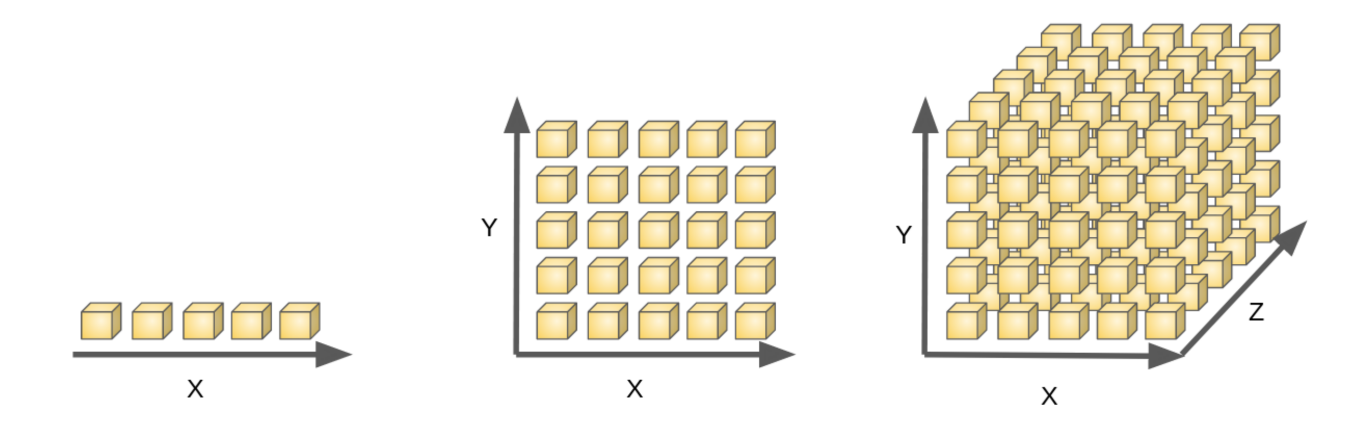
Continuous action spaces

- Tabular RL only works for small discrete action spaces.
- Robots have continuous action spaces, where the actions are changes in joint angles or torques.
- A joint angle could take any value in $[0, \pi]$.



Continuous action spaces

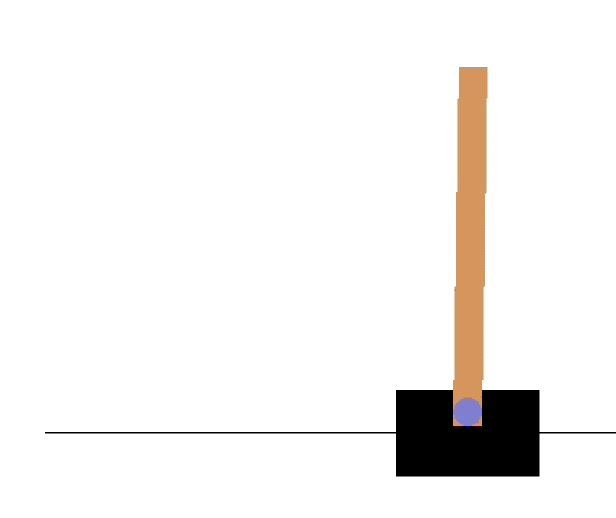
 A solution would be to discretize the action space (one action per degree), but we would fall into the curse of dimensionality.



- The more degrees of freedom, the more discrete actions, the more entries in the Q-table...
- Tabular RL cannot deal with continuous action spaces, unless we approximate the policy with an **actor-critic** architecture.

2 - Function approximation

Feature vectors



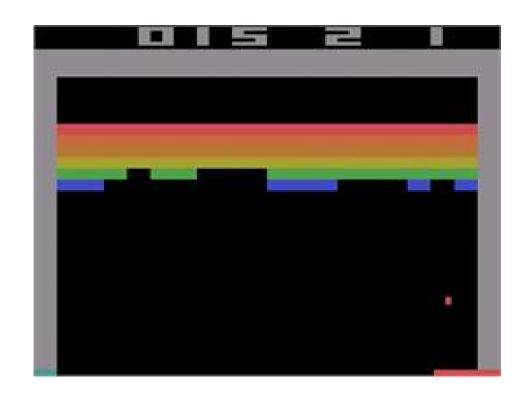
- Let's represent a state s by a vector of d features $\phi(s) = [\phi_1(s), \phi_2(s), \dots, \phi_d(s)]^T.$
- For the cartpole, the feature vector would be:

$$\phi(s) = egin{bmatrix} x \ \dot{x} \ heta \ \dot{ heta} \end{bmatrix}$$

- x is the position, θ the angle, \dot{x} and $\dot{\theta}$ their derivatives.
- We are able to represent **any state** *s* using these four variables.

Feature vectors

• For more complex problems, the feature vector should include all the necessary information (Markov property).



$$x$$
 position of the paddle x position of the ball y position of the ball x speed of the ball y speed of the position presence of brick 1 presence of brick 2 x

• In deep RL, we will **learn** these feature vectors, but let's suppose for now that we have them.

Feature vectors

Note that we can always fall back to the tabular case using one-hot encoding of the states:

$$\phi(s_1) = egin{bmatrix} 1 \ 0 \ 0 \ \cdots \ 0 \end{bmatrix} \qquad \phi(s_2) = egin{bmatrix} 0 \ 1 \ 0 \ \cdots \ 0 \end{bmatrix} \qquad \phi(s_3) = egin{bmatrix} 0 \ 0 \ 1 \ \cdots \ 0 \end{bmatrix} \qquad \ldots$$

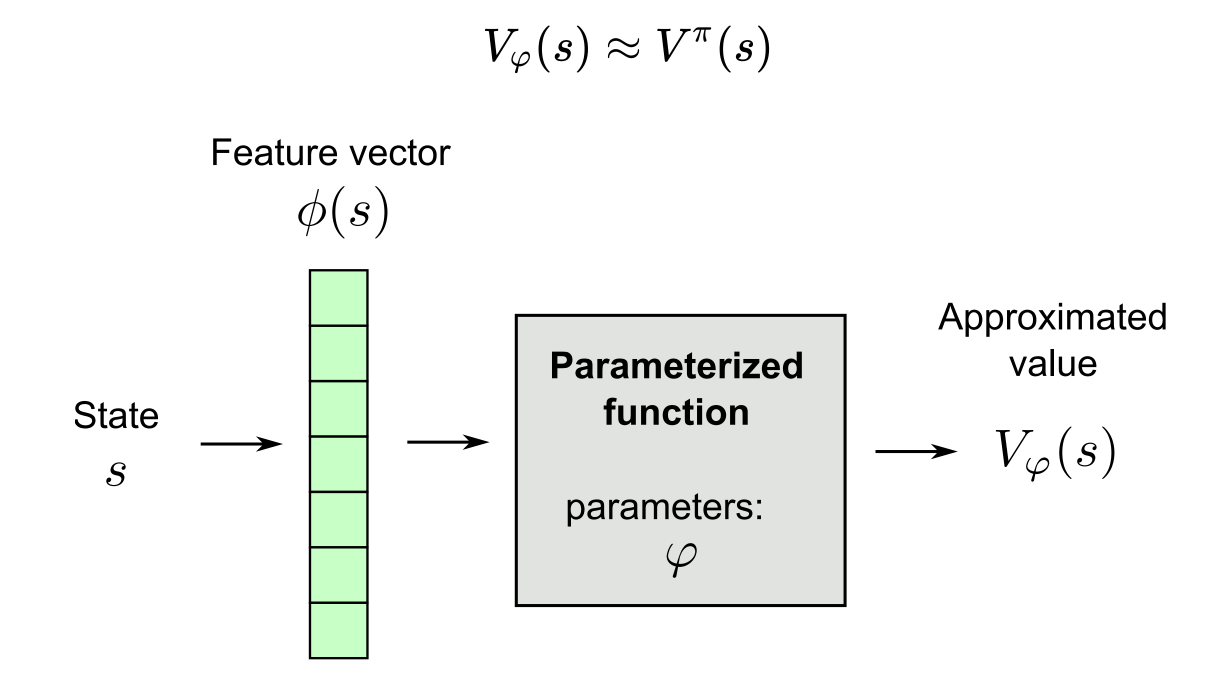
• But the idea is that we can represent states with much less values than the number of states:

$$d \ll |\mathcal{S}|$$

• We can also represent continuous state spaces with feature vectors.

State value approximation

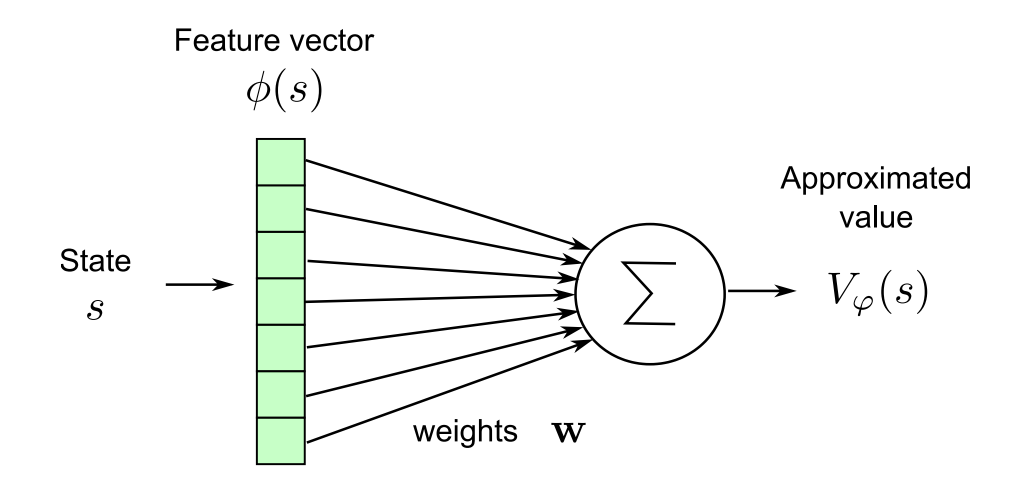
• In state value approximation, we want to approximate the state value function $V^\pi(s)$ with a parameterized function $V_\varphi(s)$:



• The parameterized function can have any form. Its has a set of parameters φ used to transform the feature vector $\phi(s)$ into an approximated value $V_{\varphi}(s)$.

Linear approximation of state value functions

• The simplest function approximator (FA) is the linear approximator.

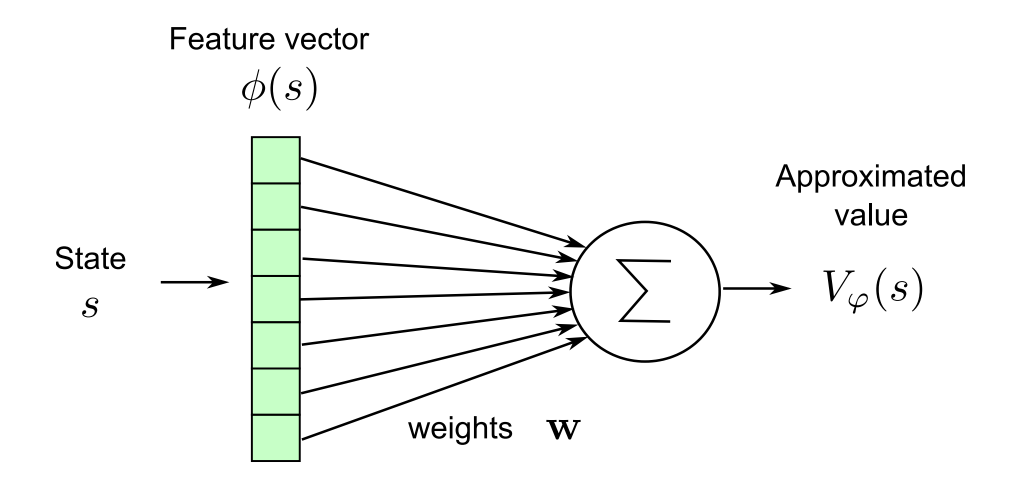


The approximated value is a linear combination of the features:

$$V_{arphi}(s) = \sum_{i=1}^d w_i \, \phi_i(s) = \mathbf{w}^T imes \phi(s)$$

- ullet The **weight vector** $\mathbf{w} = [w_1, w_2, \dots, w_d]^T$ is the set of parameters arphi of the function.
- A linear approximator is a single artificial neuron (linear regression) without a bias.

- Regardless the form of the function approximator, we want to find the parameters φ making the approximated values $V_{\varphi}(s)$ as close as possible from the true values $V^{\pi}(s)$ for all states s.
 - This is a regression problem.

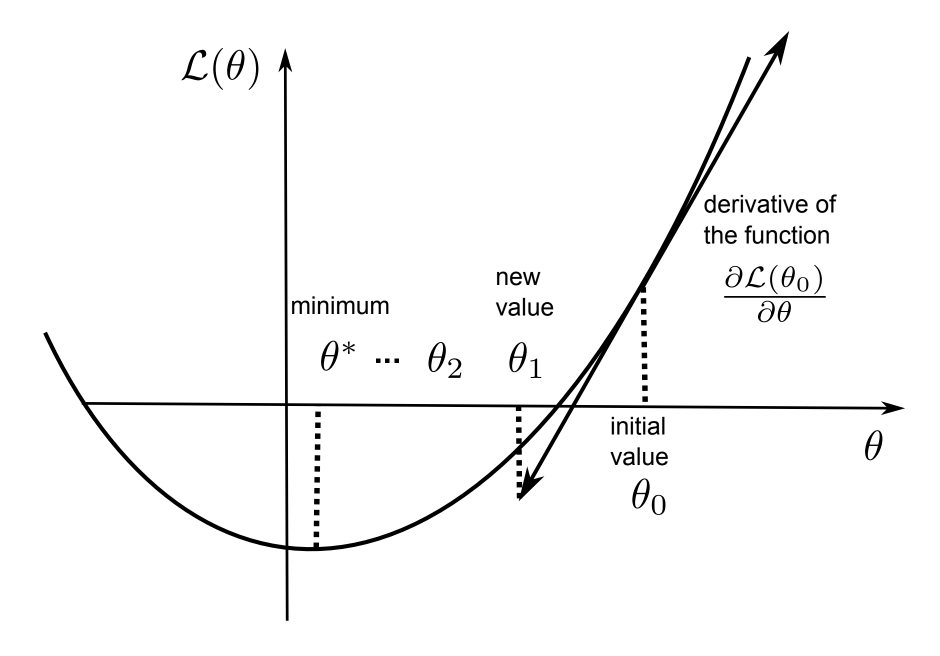


We want to minimize the mean square error between the two quantities:

$$\min_{arphi} \mathcal{L}(arphi) = \mathbb{E}_{s \in \mathcal{S}}[(V^{\pi}(s) - V_{arphi}(s))^2]$$

• The **loss function** $\mathcal{L}(\varphi)$ is minimal when the predicted values are close to the true ones on average for all states.

• Let's suppose that we know the true state values $V^{\pi}(s)$ for all states and that the parameterized function is **differentiable**.



 We can find the minimum of the loss function by applying gradient descent (GD) iteratively:

$$\Delta arphi = -\eta \,
abla_arphi \mathcal{L}(arphi)$$

• $\nabla_{\varphi} \mathcal{L}(\varphi)$ is the gradient of the loss function w.r.t to the parameters φ .

$$abla_{arphi}\mathcal{L}(arphi) = egin{bmatrix} rac{\partial \mathcal{L}(arphi)}{\partial arphi_1} \ rac{\partial \mathcal{L}(arphi)}{\partial arphi_2} \ rac{\partial \mathcal{L}(arphi)}{\partial arphi_K} \end{bmatrix}$$

• When applied repeatedly, GD converges to a local minimum of the loss function.

To minimize the mean square error,

$$\min_{arphi} \mathcal{L}(arphi) = \mathbb{E}_{s \in \mathcal{S}}[(V^{\pi}(s) - V_{arphi}(s))^2]$$

we will iteratively modify the parameters φ according to:

$$egin{aligned} \Delta arphi &= arphi_{k+1} - arphi_n = -\eta \,
abla_arphi \mathcal{L}(arphi) = -\eta \,
abla_arphi \mathbb{E}_{s \in \mathcal{S}}[(V^\pi(s) - V_arphi(s))^2] \ &= \mathbb{E}_{s \in \mathcal{S}}[-\eta \,
abla_arphi(V^\pi(s) - V_arphi(s))^2] \ &= \mathbb{E}_{s \in \mathcal{S}}[\eta \, (V^\pi(s) - V_arphi(s)) \,
abla_arphi V_arphi(s)] \end{aligned}$$

 As it would be too slow to compute the expectation on the whole state space (batch algorithm), we will sample the quantity:

$$\delta_arphi = \eta \left(V^\pi(s) - V_arphi(s)
ight)
abla_arphi V_arphi(s)$$

and update the parameters with stochastic gradient descent (SGD).

Gradient of the mse:

$$\Delta arphi = \mathbb{E}_{s \in \mathcal{S}}[\eta\left(V^{\pi}(s) - V_{arphi}(s)
ight)
abla_{arphi}V_{arphi}(s)]$$

• If we sample K states s_i from the state space:

$$\Delta arphi = \eta \, rac{1}{K} \sum_{k=1}^K (V^\pi(s_k) - V_arphi(s_k)) \,
abla_arphi V_arphi(s_k)$$

• We can also sample a single state s (online algorithm):

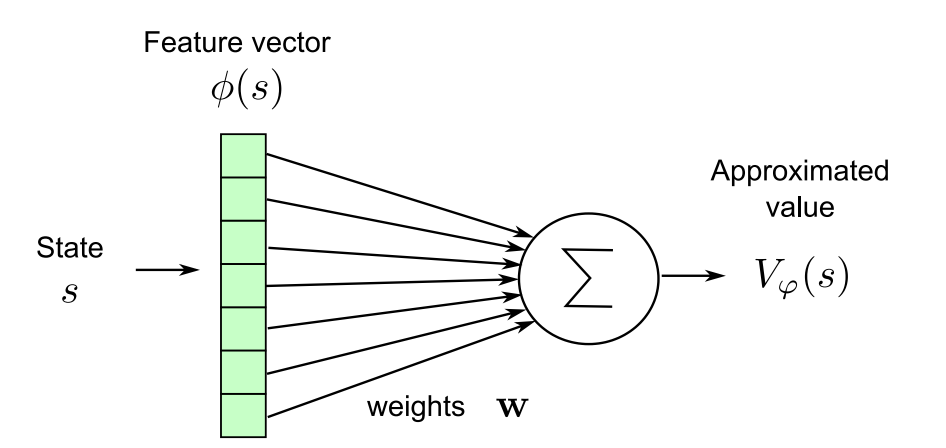
$$\Delta arphi = \eta \left(V^{\pi}(s) - V_{arphi}(s)
ight)
abla_{arphi} V_{arphi}(s)$$

• Unless stated otherwise, we will sample single states in this section, but the parameter updates will be noisy (high variance).

Linear approximation

 The approximated value is a linear combination of the features:

$$V_{arphi}(s) = \sum_{i=1}^d w_i \, \phi_i(s) = \mathbf{w}^T imes \phi(s)$$



The weights are updated using stochastic gradient descent:

$$\Delta \mathbf{w} = \eta \left(V^{\pi}(s) - V_{arphi}(s)
ight) \phi(s)$$

• This is the **delta learning rule** of linear regression and classification, with $\phi(s)$ being the input vector and $V^{\pi}(s) - V_{\varphi}(s)$ the prediction error.

Function approximation with sampling

• The rule can be used with any function approximator, we only need to be able to differentiate it:

$$\Delta arphi = \eta \left(V^{\pi}(s) - V_{arphi}(s)
ight)
abla_{arphi} V_{arphi}(s)$$

- ullet The problem is that we do not know $V^\pi(s)$, as it is what we are trying to estimate.
- ullet We can replace $V^\pi(s)$ by a sampled estimate using Monte Carlo or TD:
 - Monte Carlo function approximation:

$$\Delta arphi = \eta \left(R_t - V_arphi(s)
ight)
abla_arphi V_arphi(s)$$

■ **Temporal Difference** function approximation:

$$\Delta arphi = \eta \left(r_{t+1} + \gamma \, V_{arphi}(s') - V_{arphi}(s)
ight)
abla_{arphi} V_{arphi}(s)$$

 Note that for Temporal Difference, we actually want to minimize the TD reward-prediction error for all states, i.e. the surprise:

$$\mathcal{L}(arphi) = \mathbb{E}_{s \in \mathcal{S}}[(r_{t+1} + \gamma \, V_{arphi}(s') - V_{arphi}(s))^2] = \mathbb{E}_{s \in \mathcal{S}}[\delta_t^2]$$

Gradient Monte Carlo Algorithm for value estimation

- Algorithm:
 - Initialize the parameter φ to 0 or randomly.
 - while not converged:
 - 1. Generate an episode according to the current policy π until a terminal state s_T is reached.

$$au=(s_o,a_o,r_1,s_1,a_1,\ldots,s_T)$$

- 2. For all encountered states $s_0, s_1, \ldots, s_{T-1}$:
 - 1. Compute the return $R_t = \sum_k \gamma^k r_{t+k+1}$.
 - 2. Update the parameters using function approximation:

$$\Delta arphi = \eta \left(R_t - V_arphi(s_t)
ight)
abla_arphi V_arphi(s_t)$$

• Gradient Monte Carlo has no bias (real returns) but a high variance.

Semi-gradient Temporal Difference Algorithm for value estimation

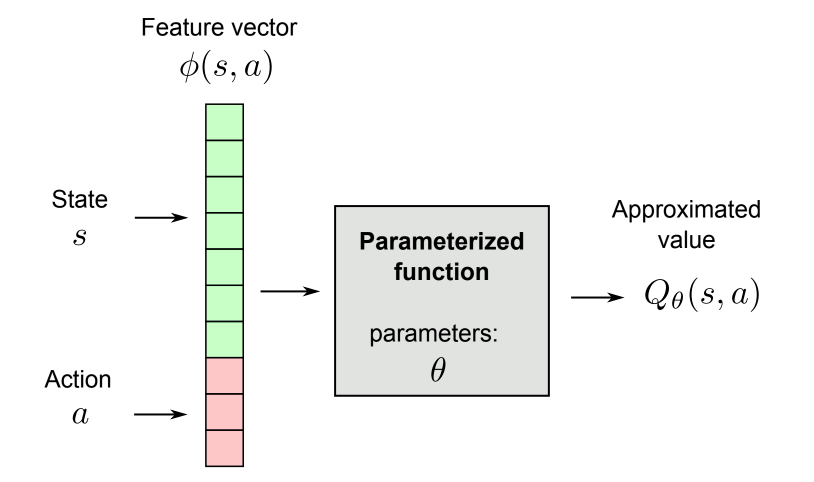
- Algorithm:
 - Initialize the parameter φ to 0 or randomly.
 - while not converged:
 - \circ Start from an initial state s_0 .
 - \circ **foreach** step t of the episode:
 - Select a_t using the current policy π in state s_t .
 - \circ Observe r_{t+1} and s_{t+1} .
 - Update the parameters using function approximation:

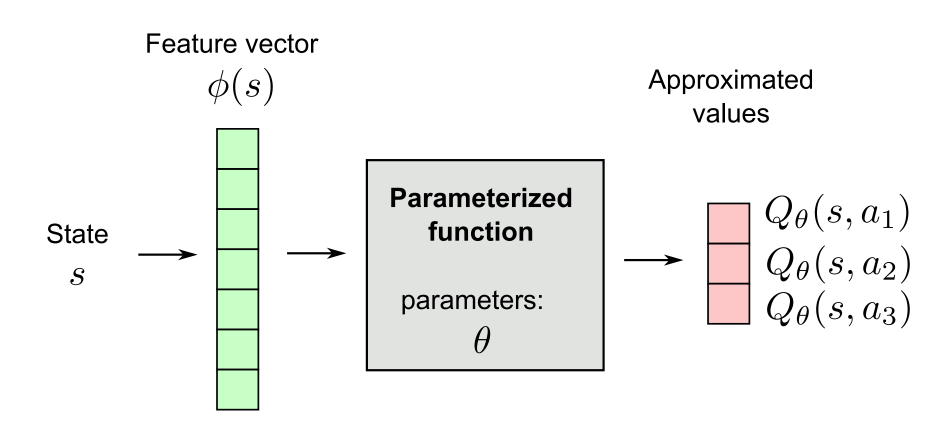
$$\Delta arphi = \eta \left(r_{t+1} + \gamma \, V_{arphi}(s_{t+1}) - V_{arphi}(s_t)
ight)
abla_{arphi} V_{arphi}(s_t)$$

- \circ if s_{t+1} is terminal: break
- Semi-gradient TD has less variance, but a significant bias as $V_{\varphi}(s_{t+1})$ is initially wrong. You can never trust these estimates completely.

Function approximation for Q-values

- ullet Q-values can be approximated by a parameterized function $Q_{ heta}(s,a)$ in the same manner.
- There are basically two options for the structure of the function approximator:
- The FA takes a feature vector for both the state s and the action a (which can be continuous) as inputs, and outputs a single Q-value $Q_{\theta}(s,a)$.
- The FA takes a feature vector for the state s as input, and outputs one Q-value $Q_{\theta}(s,a)$ per possible action (the action space must be discrete).





• In both cases, we minimize the mse between the true value $Q^{\pi}(s,a)$ and the approximated value $Q_{\theta}(s,a)$.

Q-learning with function approximation

- Initialize the parameters θ .
- while True:
 - Start from an initial state s_0 .
 - foreach step t of the episode:
 - \circ Select a_t using the behavior policy b (e.g. derived from π).
 - \circ Take a_t , observe r_{t+1} and s_{t+1} .
 - \circ Update the parameters θ :

$$\Delta heta = \eta \left(r_{t+1} + \gamma \, \max_{a} Q_{ heta}(s_{t+1}, a) - Q_{ heta}(s_{t}, a_{t})
ight)
abla_{ heta} Q_{ heta}(s_{t}, a_{t})$$

Improve greedily the learned policy:

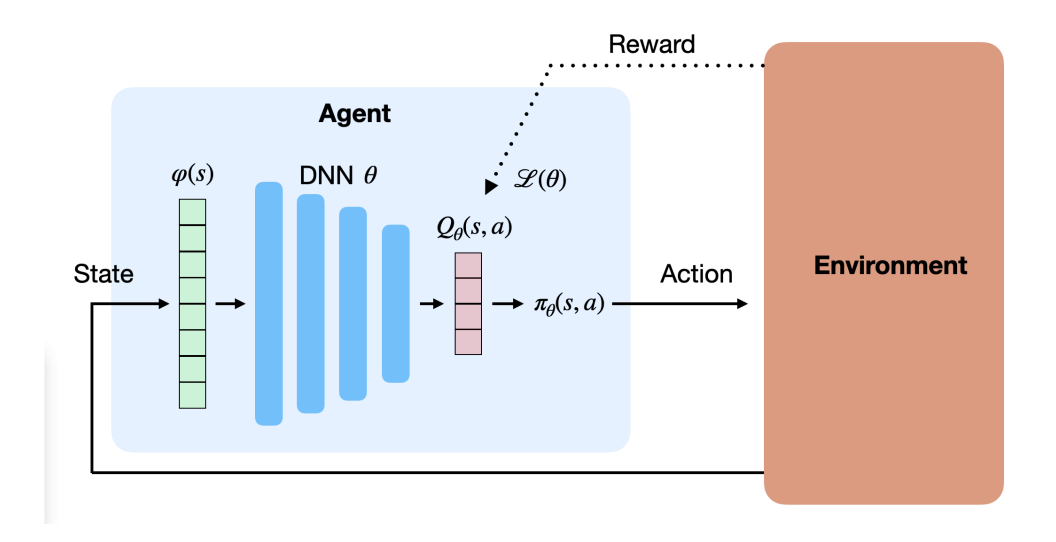
$$\pi(s_t,a) = \operatorname{Greedy}(Q_{ heta}(s_t,a))$$

 \circ if s_{t+1} is terminal: break

3 - Feature construction

Feature construction

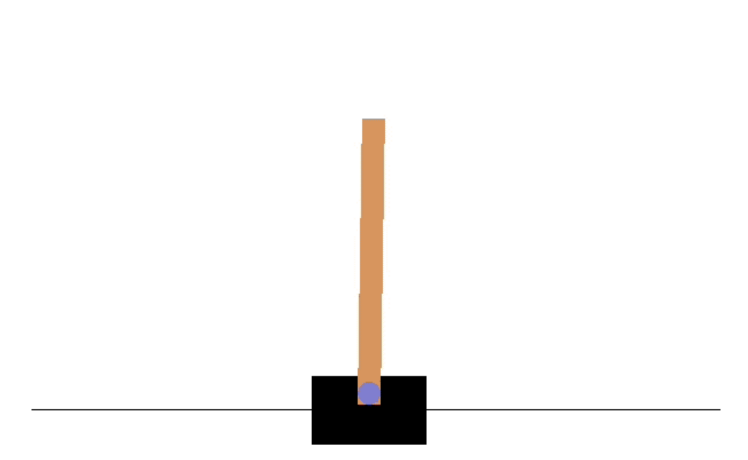
• Before we dive into deep RL (i.e. RL with non-linear FA), let's see how we can design good **feature vectors** for linear function approximation.



- The problem with deep NN is that they need a lot of samples to converge, what worsens the fundamental problem of RL: sample efficiency.
- By engineering the right features, we could use linear approximators, which converge much faster.
- The convergence of linear FA is **guaranteed**, not (always) non-linear ones.

Why do we need to choose features?





$$\phi(s) = egin{bmatrix} x \ \dot{x} \ heta \ \dot{ heta} \end{bmatrix}$$

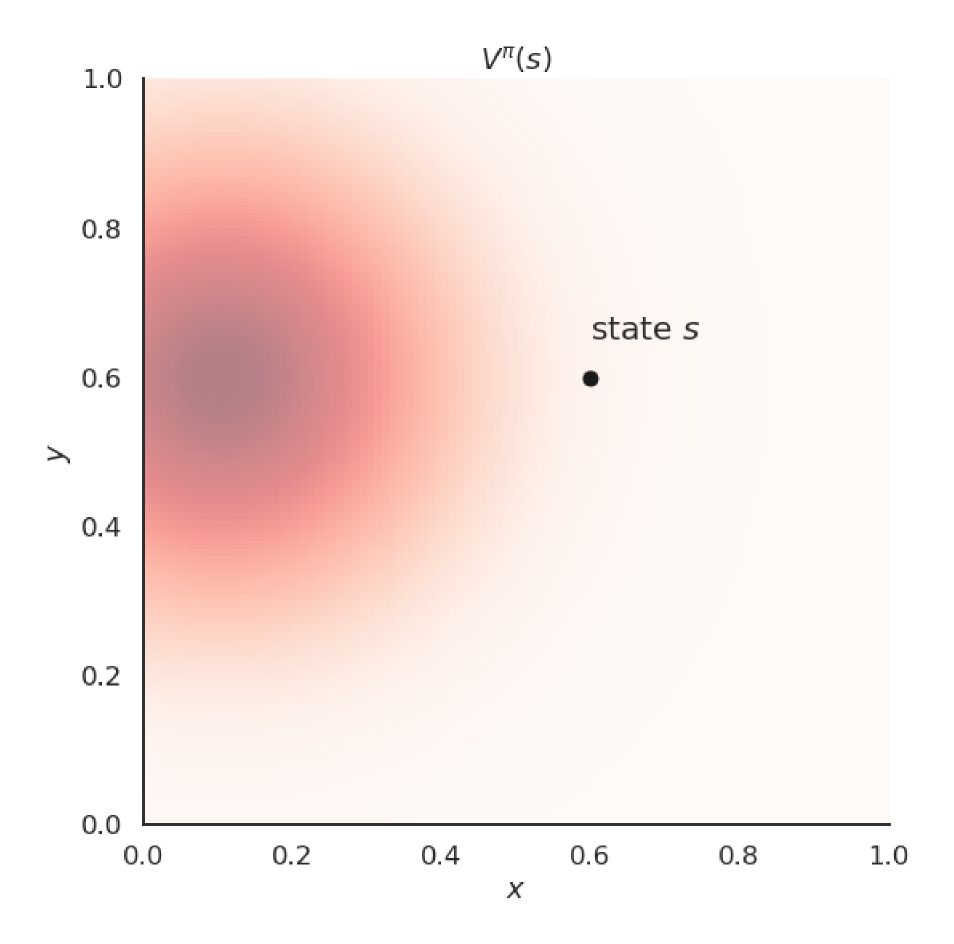
- x is the position, heta the angle, \dot{x} and $\dot{ heta}$ their derivatives.
- Can we predict the value of a state linearly?

$$V_{arphi}(s) = \sum_{i=1}^d w_i \, \phi_i(s) = \mathbf{w}^T imes \phi(s)$$

- No, a high angular velocity $\dot{\theta}$ is good when the pole is horizontal (going up) but bad if the pole is vertical (will not stop).
- ullet The value would depend linearly on something like $\dot{ heta} \sin heta$, which is a non-linear combination of features.

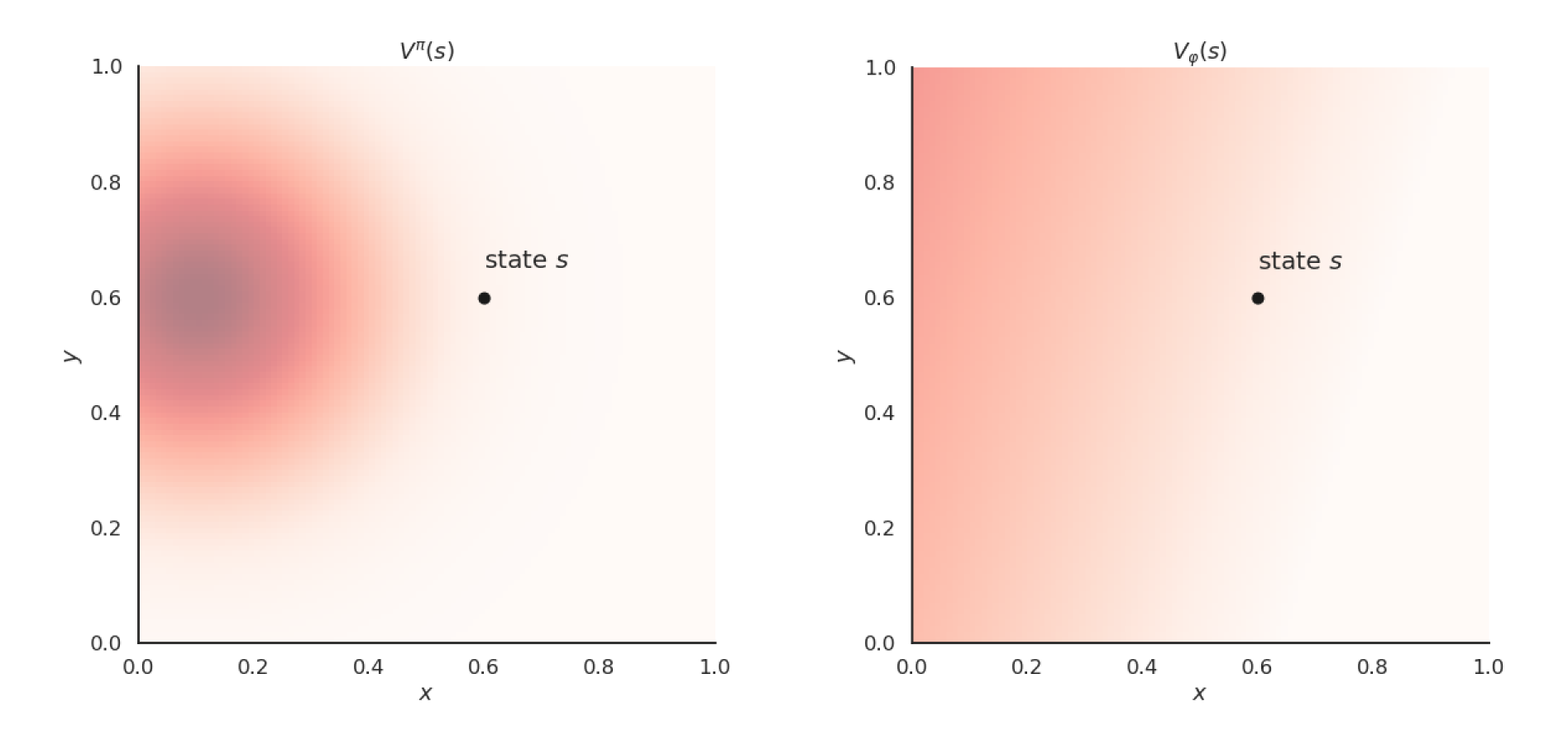
Feature coding

- Let's suppose we have a simple problem where the state s is represented by two continuous variables x and y.
- The true value function $V^\pi(s)$ is a non-linear function of x and y.



Linear approximation

- If we apply linear FA directly on the feature vector [x,y], we catch the tendency of $V^{\pi}(s)$ but we make a lot of bad predictions:
 - high bias (underfitting).



Polynomial features

- To introduce non-linear relationships between continuous variables, a simple method is to construct the feature with **polynomials** of the variables.
- Example with polynomials of order 2:

$$\phi(s) = egin{bmatrix} 1 & x & y & xy & x^2 & y^2 \end{bmatrix}^T$$

- We transform the two input variables x and y into a vector with 6 elements. The 1 (order 0) is there to learn the offset.
- Example with polynomials of order 3:

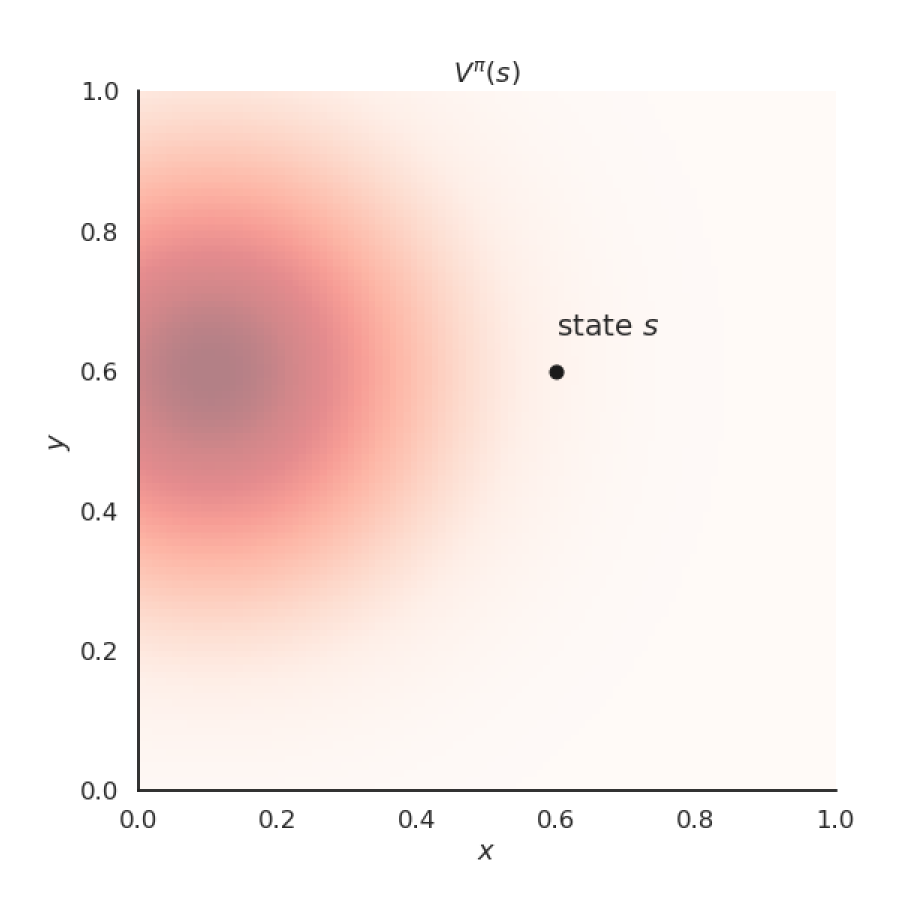
$$\phi(s) = egin{bmatrix} 1 & x & y & xy & x^2 & y^2 & x^2y & xy^2 & x^3 & y^3 \end{bmatrix}^T$$

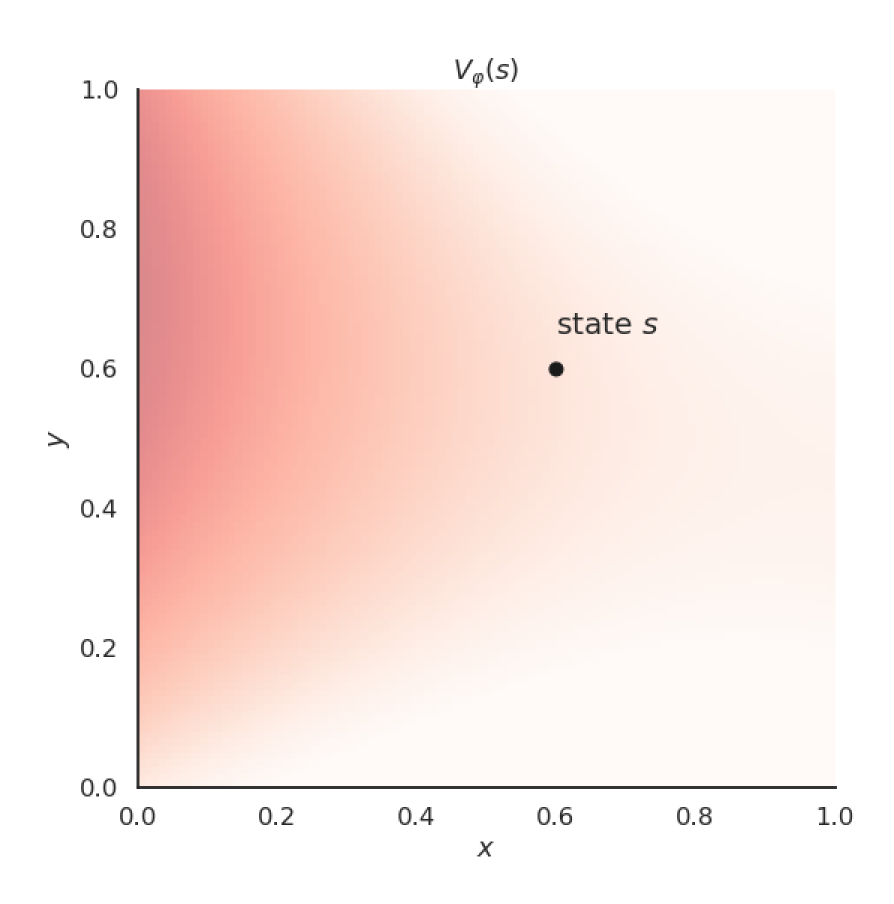
• And so on. We then just need to apply linear FA on these feature vectors (polynomial regression).

$$V_{arphi}(s) = w_0 + w_1\,x + w_2\,y + w_3\,x\,y + w_4\,x^2 + w_5\,y^2 + \dots$$

Polynomials

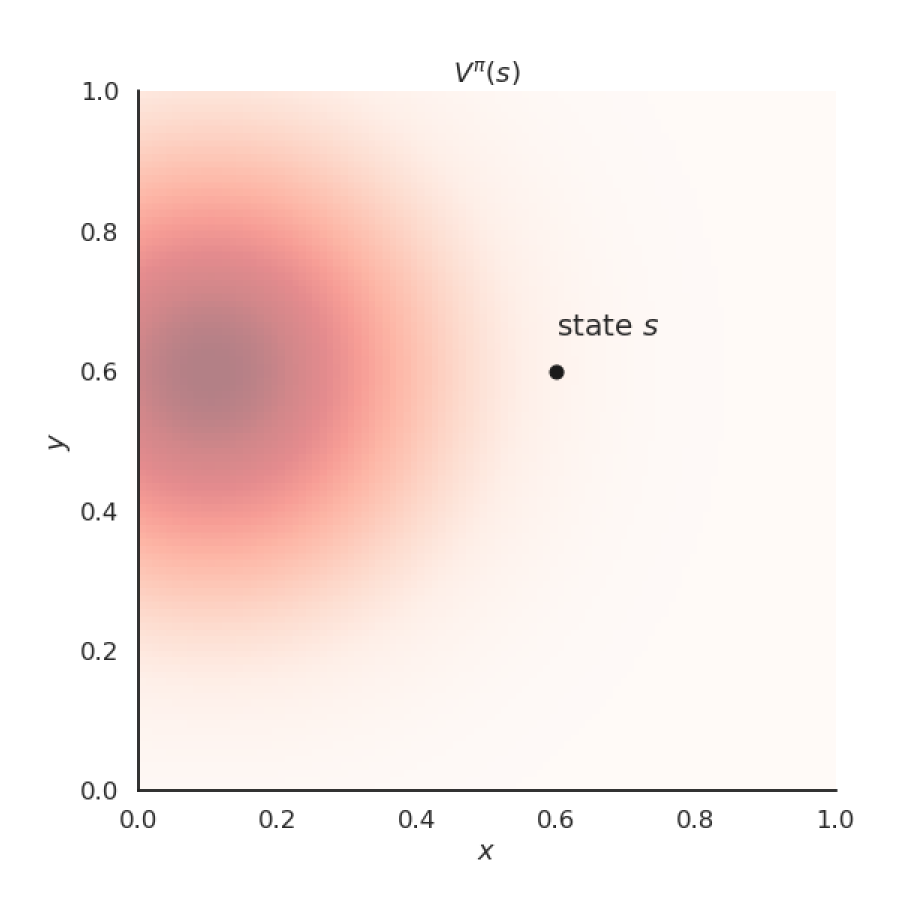
• Polynomials of order 2 already allow to get a better approximation.

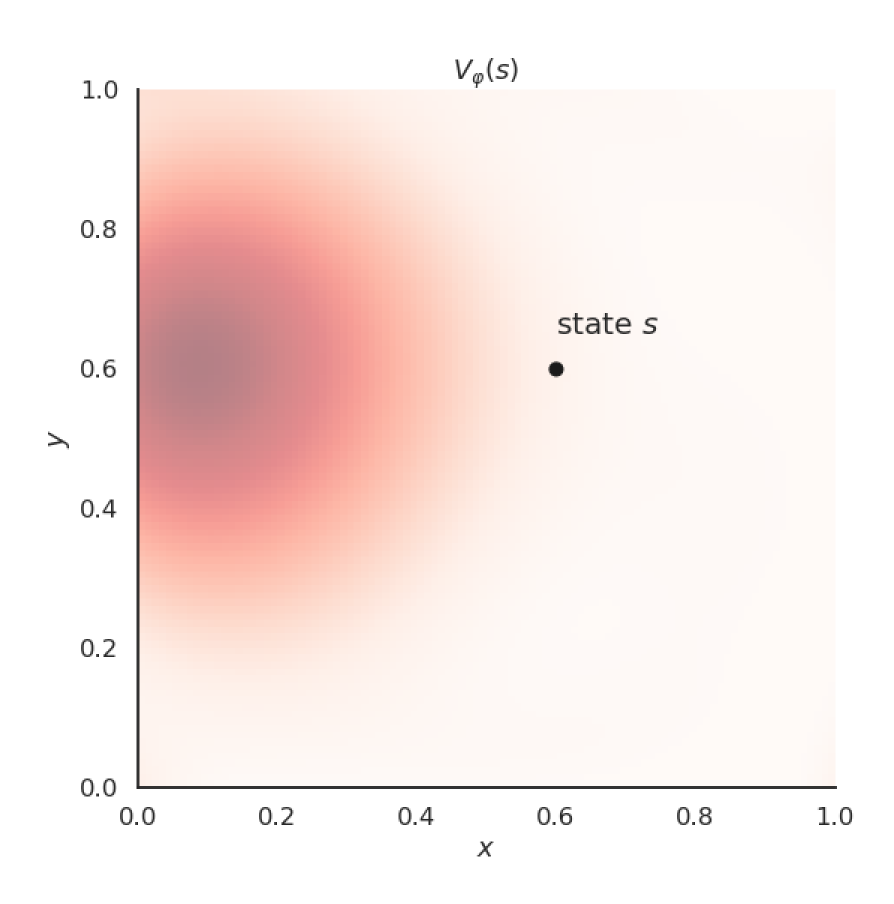




Polynomials

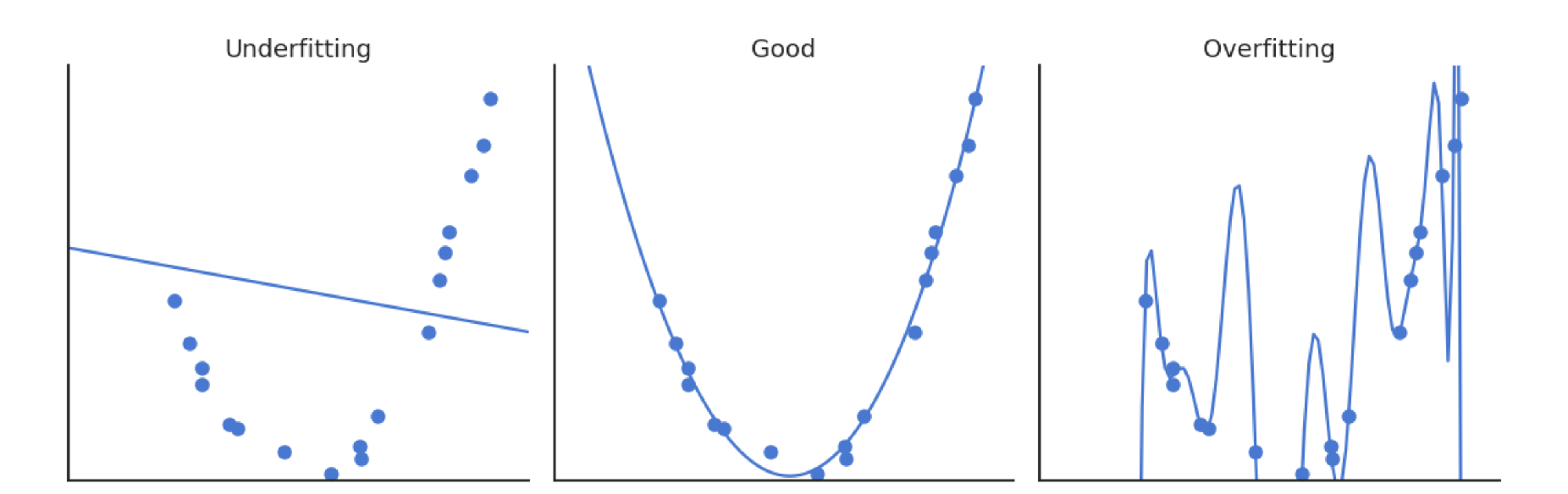
• Polynomials of order 6 are an even better fit for our problem.





Polynomials

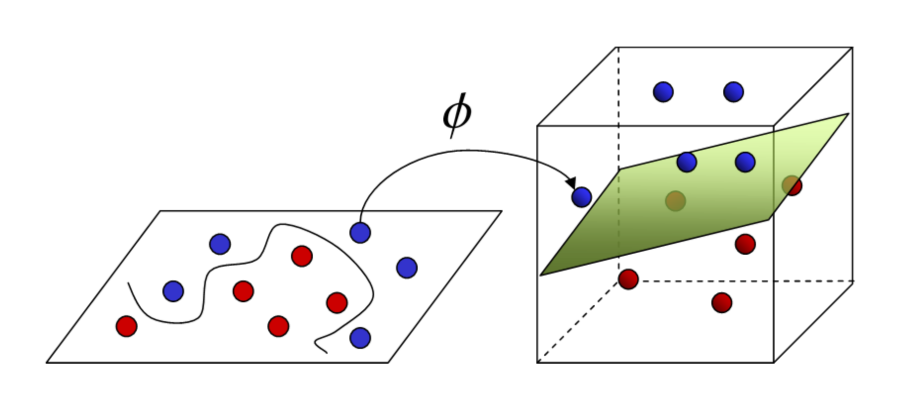
- The higher the degree of the polynomial, the better the fit, but the number of features grows exponentially.
 - Computational complexity.
 - Overfitting: if we only sample some states, high-order polynomials will not interpolate correctly.



Feature spaces

Input Space

- In machine learning (ML), the oldest trick in the book is the use of a **feature space** allowing to project data into a higher-dimensional and non-linear space, so that the problem becomes linearly separable / predictable.
- We can do the same in RL, using any kind of **feature extraction** methods:

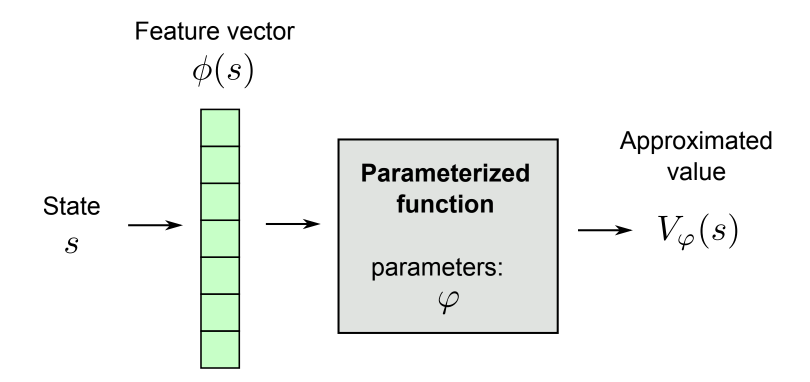


- Polynomial features
- Gaussian (RBF) features
- Fourier transforms
- Tile coding
- Deep neural networks

• If the right features heve been extracted, linear methods can be applied.

Feature Space

Summary of function approximation



- In FA, we project the state information into a **feature space** to get a better representation.
- We then apply a linear approximation algorithm to estimate the value function:

$$V_{arphi}(s) = \mathbf{w}^T \, \phi(s)$$

• The linear FA is trained using some variant of gradient decent:

$$\Delta \mathbf{w} = \eta \left(V^{\pi}(s) - V_{arphi}(s)
ight) \phi(s)$$

- Deep neural networks are the most powerful function approximators in supervised learning.
- Do they also work with RL?