

Reinforcement Learning and Optimal Control

A Selective Overview

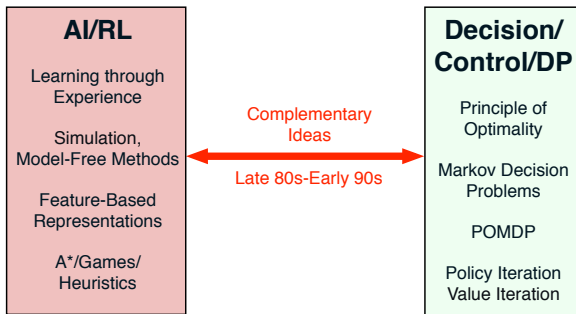
Dimitri P. Bertsekas

Laboratory for Information and Decision Systems
Massachusetts Institute of Technology

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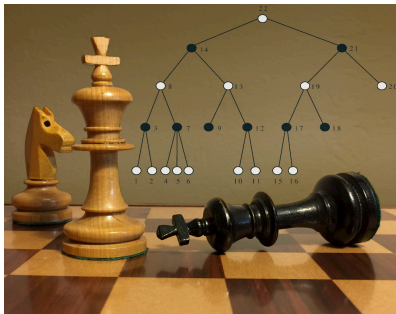
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Reinforcement Learning (RL): A Happy Union of AI and Decision/Control Ideas



Historical highlights

- Exact DP, optimal control (Bellman, Shannon, 1950s ...)
- First major successes: Backgammon programs (Tesauro, 1992, 1996)
- Algorithmic progress, analysis, applications, first books (mid 90s ...)
- Machine Learning, BIG Data, Robotics, Deep Neural Networks (mid 2000s ...)
- AlphaGo and Alphazero (DeepMind, 2016, 2017)



AlphaZero

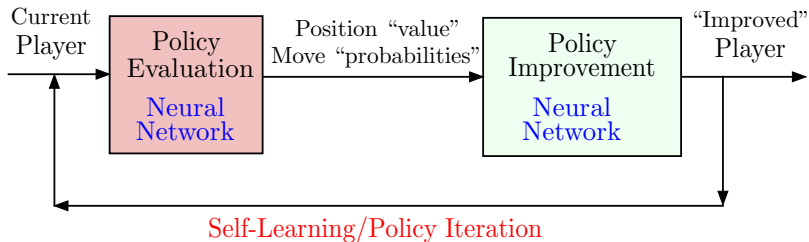
Plays much better than all chess programs

Plays different!

Learned from scratch ... with 4 hours of training!

Same algorithm learned multiple games (Go, Shogi)

AlphaZero was Trained Using Self-Generated Data



- The "current" player plays games that are used to "train" an "improved" player
- At a given position, the "move probabilities" and the "value" of a position are approximated by a deep neural net (NN)
- Successive NNs are trained using self-generated data and a form of regression
- A form of randomized policy improvement **Monte-Carlo Tree Search** (MCTS) generates move probabilities
- AlphaZero bears similarity to earlier works, e.g., **TD-Gammon** (Tesauro, 1992), but is more complicated because of the MCTS and the deep NN
- The success of AlphaZero is due to a skillful implementation/integration of known ideas, and awesome computational power

Approximate DP/RL Methodology is now Ambitious and Universal

Exact DP applies (in principle) to a very broad range of optimization problems

- Deterministic \longleftrightarrow Stochastic
- Combinatorial optimization \longleftrightarrow Optimal control w/ infinite state/control spaces
- One decision maker \longleftrightarrow Two player games
- ... BUT is plagued by the **curse of dimensionality** and **need for a math model**

Approximate DP/RL overcomes the difficulties of exact DP by:

- **Approximation** (use neural nets and other architectures to reduce dimension)
- **Simulation** (use a computer model in place of a math model)

State of the art:

- **Broadly applicable methodology**: Can address broad range of challenging problems. Deterministic-stochastic-dynamic, discrete-continuous, games, etc
- There are **no methods that are guaranteed to work** for all or even most problems
- There are **enough methods to try with a reasonable chance of success** for most types of optimization problems
- **Role of the theory**: Guide the art, delineate the sound ideas

Approximation in Value Space

Central Idea: Lookahead with an approximate cost

- Compute an approximation \tilde{J} to the optimal cost function J^*
- At current state, apply control that attains the minimum in

$$\text{Current Stage Cost} + \tilde{J}(\text{Next State})$$

Multistep lookahead extension

- At current state solve an ℓ -step DP problem using terminal cost \tilde{J}
- Apply the first control in the optimal policy for the ℓ -step problem

Example approaches to compute \tilde{J} :

- **Problem approximation:** Use as \tilde{J} the optimal cost function of a simpler problem
- **Rollout and model predictive control:** Use a single policy iteration, with cost evaluated on-line by simulation or limited optimization
- **Self-learning/approximate policy iteration (API):** Use as \tilde{J} an approximation to the cost function of the final policy obtained through a policy iteration process
- **Role of neural networks:** "Learn" the cost functions of policies in the context of API; "learn" policies obtained by value space approximation

The purpose of this talk

To selectively review some of the methods, and bring out some of the AI-DP connections

References

- Quite a few Exact DP books (1950s-present starting with Bellman; my latest book "Abstract DP" came out earlier this year)
- Quite a few DP/Approximate DP/RL/Neural Nets books (1996-Present)
 - ▶ Bertsekas and Tsitsiklis, Neuro-Dynamic Programming, 1996
 - ▶ Sutton and Barto, 1998, Reinforcement Learning (new edition 2019, Draft on-line)
 - ▶ **NEW DRAFT BOOK**: Bertsekas, Reinforcement Learning and Optimal Control, 2019, on-line
- Many surveys on all aspects of the subject; Tesauro's papers on computer backgammon, and Silver, et al., papers on AlphaZero

RL uses Max/Value, DP uses Min/Cost

- **Reward of a stage** = (Opposite of) Cost of a stage.
- **State value** = (Opposite of) State cost.
- **Value (or state-value) function** = (Opposite of) Cost function.

Controlled system terminology

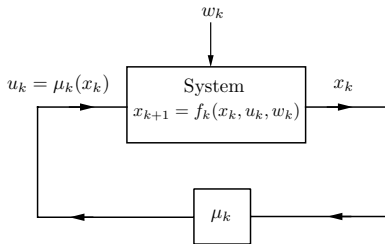
- **Agent** = Decision maker or controller.
- **Action** = Control.
- **Environment** = Dynamic system.

Methods terminology

- **Learning** = Solving a DP-related problem using simulation.
- **Self-learning (or self-play in the context of games)** = Solving a DP problem using simulation-based policy iteration.
- **Planning vs Learning distinction** = Solving a DP problem with model-based vs model-free simulation.

- 1 Approximation in Value Space
- 2 Problem Approximation
- 3 Rollout and Model Predictive Control
- 4 Parametric Approximation - Neural Networks
- 5 Neural Networks and Approximation in Value Space
- 6 Model-free DP in Terms of Q -Factors
- 7 Policy Iteration - Self-Learning

Finite Horizon Problem - Exact DP



- System

$$x_{k+1} = f_k(x_k, u_k, w_k), \quad k = 0, 1, \dots, N-1$$

where x_k : State, u_k : Control, w_k : Random disturbance

- Cost function:

$$E \left\{ g_N(x_N) + \sum_{k=0}^{N-1} g_k(x_k, u_k, w_k) \right\}$$

- Perfect state information: u_k is applied with (exact) knowledge of x_k
- **Optimization over feedback policies** $\{\mu_0, \dots, \mu_{N-1}\}$: Rules that specify the control $\mu_k(x_k)$ to apply at each possible state x_k that can occur

The DP Algorithm and Approximation in Value Space

Go backwards, $k = N - 1, \dots, 0$, using

$$J_N(x_N) = g_N(x_N)$$

$$J_k(x_k) = \min_{u_k} E_{w_k} \left\{ g_k(x_k, u_k, w_k) + J_{k+1}(f_k(x_k, u_k, w_k)) \right\}$$

$J_k(x_k)$: **Optimal cost-to-go** starting from state x_k

Approximate DP is motivated by the **ENORMOUS** computational demands of exact DP

Approximation in value space: Use an approximate cost-to-go function \tilde{J}_{k+1}

$$\tilde{\mu}_k(x_k) \in \arg \min_{u_k} E_{w_k} \left\{ g_k(x_k, u_k, w_k) + \tilde{J}_{k+1}(f_k(x_k, u_k, w_k)) \right\}$$

There is also a **multistep lookahead** version

At state x_k solve an ℓ -step DP problem with terminal cost function approximation $\tilde{J}_{k+\ell}$.
Use the first control in the optimal ℓ -step sequence.

Approximation in Value Space Methods

One-step case at state x_k :

Approximate minimization

$$\min_{u_k} E \left\{ g_k(x_k, u_k, w_k) + \tilde{J}_{k+1}(x_{k+1}) \right\}$$

Diagram description: The equation is centered. A red double-headed arrow labeled "First Step" spans from $g_k(x_k, u_k, w_k)$ to $\tilde{J}_{k+1}(x_{k+1})$. A blue double-headed arrow labeled "Future" spans from $\tilde{J}_{k+1}(x_{k+1})$ to the right. An arrow points from "Approximate minimization" to the \min_{u_k} operator. Another arrow points from "Computation of \tilde{J}_{k+1} :" to the $\tilde{J}_{k+1}(x_{k+1})$ term.

Approximations:

Simplify $E\{\cdot\}$
(certainty equivalence)
Adaptive simulation

Computation of \tilde{J}_{k+1} :

Problem approximation
Rollout
Model Predictive Control
Parametric approximation
Aggregation

Multistep case at state x_k :

DP minimization

$$\min_{u_k, \mu_{k+1}, \dots, \mu_{k+\ell-1}} E \left\{ g_k(x_k, u_k, w_k) + \sum_{m=k+1}^{k+\ell-1} g_k(x_m, \mu_m(x_m), w_m) + \tilde{J}_{k+\ell}(x_{k+\ell}) \right\}$$

Diagram description: The equation is centered. A red double-headed arrow labeled "First ℓ Steps" spans from $g_k(x_k, u_k, w_k)$ to $\sum_{m=k+1}^{k+\ell-1} g_k(x_m, \mu_m(x_m), w_m)$. A blue double-headed arrow labeled "Future" spans from $\tilde{J}_{k+\ell}(x_{k+\ell})$ to the right. An arrow points from "DP minimization" to the \min operator.

Lookahead Minimization

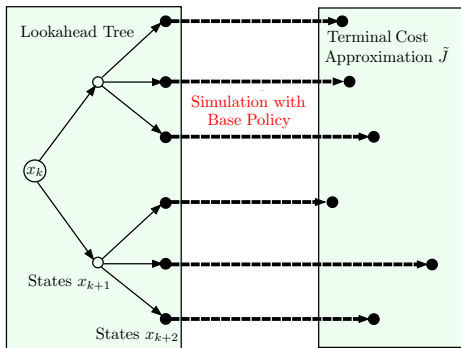
Cost-to-go
Approximation

Use as cost-to-go approximation \tilde{J}_{k+1} the **exact cost-to-go of a simpler problem**

Many problem-dependent possibilities:

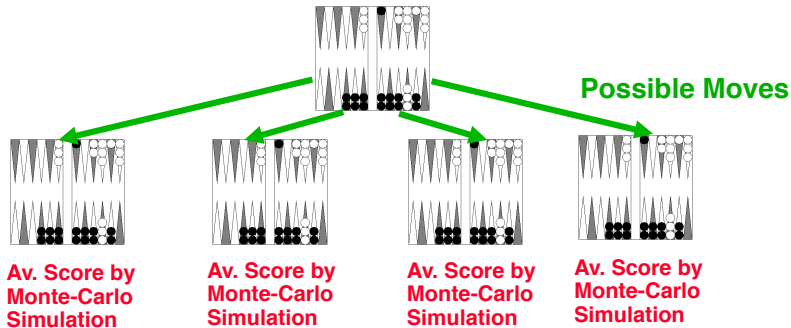
- **Probabilistic approximation**
 - ▶ Certainty equivalence: Replace stochastic quantities by deterministic ones (makes the lookahead minimization deterministic)
 - ▶ Approximate expected values by limited simulation
 - ▶ Partial versions of certainty equivalence
- **Enforced decomposition of coupled subsystems**
 - ▶ One-subsystem-at-a-time optimization
 - ▶ Constraint decomposition
 - ▶ Lagrangian relaxation
- **Aggregation**: Group states together and view the groups as aggregate states
 - ▶ Hard aggregation: \tilde{J}_{k+1} is a piecewise constant approximation to J_{k+1}
 - ▶ Feature-based aggregation: The aggregate states are defined by "features" of the original states
 - ▶ Biased hard aggregation: \tilde{J}_{k+1} is a piecewise constant local correction to some other approximation \hat{J}_{k+1} , e.g., one provided by a neural net

Rollout: On-Line Simulation-Based Approximation in Value Space



- The base policy can be **any suboptimal policy** (obtained by another method)
- **One-step or multistep lookahead**; exact minimization or a "randomized form of lookahead" that involves "adaptive" simulation and Monte Carlo tree search
- **With or without terminal cost approximation** (obtained by another method)
- Some forms of **model predictive control** can be viewed as special cases (base policy is a short-term deterministic optimization)
- Important theoretical fact: **With exact lookahead and no terminal cost approximation, the rollout policy improves over the base policy**

Example of Rollout: Backgammon (Tesauro, 1996)



- Base policy was a backgammon player developed by a different RL method [TD(λ) trained with a neural network]; was also used for terminal cost approximation
- The best backgammon players are based on rollout ... but are too slow for real-time play (MC simulation takes too long)

AlphaGo has similar structure to backgammon

The base policy and terminal cost approximation are obtained with a deep neural net. In AlphaZero the rollout-with-base-policy part was dropped (long lookahead suffices)

Parametric Approximation in Value Space

Lookahead Minimization **Cost-to-go Approximation**

First ℓ Steps “Future”

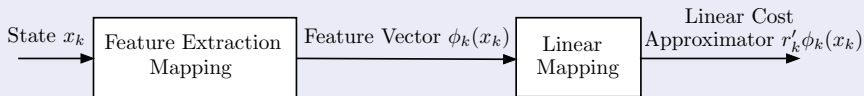
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$$\min_{u_k, \mu_{k+1}, \dots, \mu_{k+\ell-1}} E \left\{ g_k(x_k, u_k, w_k) + \sum_{m=k+1}^{k+\ell-1} g_k(x_m, \mu_m(x_m), w_m) + \tilde{J}_{k+\ell}(x_{k+\ell}) \right\}$$

↑
Parametric approximation

\tilde{J}_k comes from a class of functions $\tilde{J}_k(x_k, r_k)$, where r_k is a tunable parameter vector

Feature-based architectures: The linear case



This is just DP with intermediate approximation at each step

- Start with $\tilde{J}_N = g_N$ and **sequentially train going backwards**, until $k = 0$
- Given \tilde{J}_{k+1} , we **construct a number of samples** (x_k^s, β_k^s) , $s = 1, \dots, q$,

$$\beta_k^s = \min_u E \left\{ g(x_k^s, u, w_k) + \tilde{J}_{k+1}(f_k(x_k^s, u, w_k), r_{k+1}) \right\}, \quad s = 1, \dots, q$$

- We “train” \tilde{J}_k on the set of samples (x_k^s, β_k^s) , $s = 1, \dots, q$

Training by least squares/regression

- We minimize over r_k

$$\sum_{s=1}^q (\tilde{J}_k(x_k^s, r_k) - \beta_k^s)^2 + \gamma \|r_k - \bar{r}\|^2$$

where \bar{r} is an initial guess for r_k and $\gamma > 0$ is a regularization parameter

Major fact about neural networks

They **automatically construct features** to be used in a linear architecture

- Neural nets are approximation architectures of the form

$$\tilde{J}(x, v, r) = \sum_{i=1}^m r_i \phi_i(x, v) = r' \phi(x, v)$$

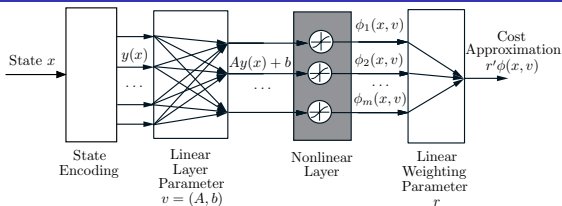
involving two parameter vectors r and v with different roles

- **View $\phi(x, v)$ as a feature vector**
- **View r as a vector of linear weights** for $\phi(x, v)$
- By training v jointly with r , we obtain automatically generated features!

Neural nets can be used in the fitted value iteration scheme

Train the stage k neural net (i.e., compute \tilde{J}_k) using a training set generated with the stage $k + 1$ neural net (which defines \tilde{J}_{k+1})

Neural Network with a Single Nonlinear Layer

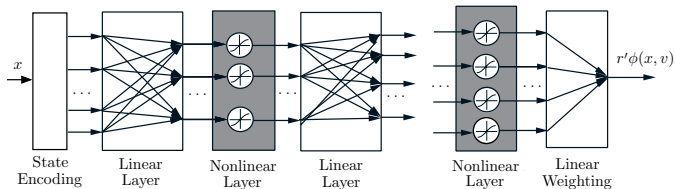


- State encoding (could be the identity, could include special features of the state)
- Linear layer $Ay(x) + b$ [parameters to be determined: $v = (A, b)$]
- Nonlinear layer produces m outputs $\phi_i(x, v) = \sigma((Ay(x) + b)_i)$, $i = 1, \dots, m$
- σ is a scalar nonlinear differentiable function; several types have been used (hyperbolic tangent, logistic, rectified linear unit)
- Training problem is to use the training set (x^s, β^s) , $s = 1, \dots, q$, for

$$\min_{v, r} \sum_{s=1}^q \left(\sum_{i=1}^m r_i \phi_i(x^s, v) - \beta^s \right)^2 + (\text{Regularization Term})$$

- Solved often with incremental gradient methods (known as **backpropagation**)
- **Universal approximation theorem:** With sufficiently large number of parameters, "arbitrarily" complex functions can be closely approximated

Deep Neural Networks



- More complex NNs are formed by **concatenation of multiple layers**
- The outputs of each nonlinear layer become the inputs of the next linear layer
- **A hierarchy of features**
- Considerable success has been achieved in major contexts

Possible reasons for the success

- With **more complex features**, the number of parameters in the linear layers may be drastically decreased
- We may **use matrices A with a special structure** that encodes special linear operations such as convolution

- The Q -factor of a state-control pair (x_k, u_k) at time k is defined by

$$Q_k(x_k, u_k) = E\{g_k(x_k, u_k, w_k) + J_{k+1}(x_{k+1})\}$$

where J_{k+1} is the optimal cost-to-go function for stage $k + 1$

- Note that

$$J_k(x_k) = \min_{u \in U_k(x_k)} Q_k(x_k, u)$$

so the DP algorithm is written in terms of Q_k

$$Q_k(x_k, u_k) = E\left\{g_k(x_k, u_k, w_k) + \min_{u \in U_{k+1}(x_{k+1})} Q_{k+1}(x_{k+1}, u)\right\}$$

- We can approximate Q -factors instead of costs

- Consider fitted value iteration of Q-factor parametric approximations

$$\tilde{Q}_k(x_k, u_k, r_k) \approx E \left\{ g_k(x_k, u_k, w_k) + \min_{u \in U_{k+1}(x_{k+1})} \tilde{Q}_{k+1}(x_{k+1}, u, r_{k+1}) \right\}$$

(Note a mathematical magic: **The order of $E\{\cdot\}$ and \min have been reversed.**)

- We obtain $\tilde{Q}_k(x_k, u_k, r_k)$ by training with many pairs $((x_k^s, u_k^s), \beta_k^s)$, where β_k^s is a sample of the approximate Q-factor of (x_k^s, u_k^s) . **No need to compute $E\{\cdot\}$**
- No need for a model to obtain β_k^s .** Sufficient to have a simulator that generates random samples of state-control-cost-next state

$$((x_k, u_k), (g_k(x_k, u_k, w_k), x_{k+1}))$$

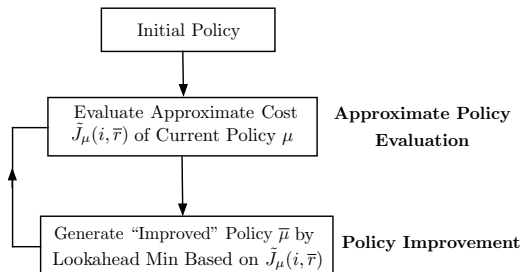
- Having computed r_k , the one-step lookahead control is obtained on-line as

$$\bar{\mu}_k(x_k) = \arg \min_{u \in U_k(x_k)} \tilde{Q}_k(x_k, u, r_k)$$

without the need of a model or expected value calculations

- Also the **on-line calculation of the control is simplified**

A Few Remarks on Infinite Horizon Problems



- **Most popular setting:** Stationary finite-state system, stationary policies, discounting or termination state
- Policy iteration (PI) method generates a sequence of policies
 - ▶ The **current policy μ is evaluated** using a parametric architecture: $\tilde{J}_\mu(x, \bar{r})$
 - ▶ An **"improved" policy $\bar{\mu}$ is obtained** by one-step lookahead using $\tilde{J}_\mu(x, \bar{r})$
- The architecture is trained using simulation data with μ
- Thus the system "observes itself" under μ and uses the data to "learn" the improved policy $\bar{\mu}$ - **"self-learning"**
- Exact PI converges to an optimal policy; **approximate PI "converges" to within an "error zone" of the optimal**, then oscillates
- **TD-Gammon, AlphaGo, and AlphaZero, all use forms of approximate PI for training**

A Few Topics we did not Cover in this Talk

- **Infinite horizon extensions**: Approximate value and policy iteration methods, error bounds, model-based and model-free methods
- **Temporal difference methods**: A class of methods for policy evaluation in infinite horizon problems with a rich theory, issues of variance-bias tradeoff
- **Sampling for exploration**, in the context of policy iteration
- **Monte Carlo tree search**, and related methods
- **Aggregation methods**, synergism with other approximate DP methods
- **Approximation in policy space**, actor-critic methods, policy gradient and cross-entropy methods
- **Special aspects of imperfect state information problems**, connections with traditional control schemes
- **Infinite spaces optimal control**, connections with aggregation schemes
- **Special aspects of deterministic problems**: Shortest paths and their use in approximate DP
- **A broad view of using simulation for large-scale computations**: Methods for large systems of equations and linear programs, connection to proximal algorithms

Concluding Remarks

Some words of caution

- There are challenging implementation issues in all approaches, and **no fool-proof methods**
- Problem approximation and feature selection require **domain-specific knowledge**
- **Training algorithms are not as reliable** as you might think by reading the literature
- Approximate PI involves **oscillations**
- **Recognizing success or failure** can be a challenge!
- The RL successes in game contexts are spectacular, but they have benefited from **perfectly known and stable models** and **small number of controls** (per state)
- **Problems with partial state observation** remain a big challenge

On the positive side

- **Massive computational power** together with distributed computation are a source of hope
- **Silver lining**: We can begin to address practical problems of unimaginable difficulty!
- There is **an exciting journey ahead!**

Thank you!