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# Coordinated Reinforcement Learning

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**Carlos Guestrin**

Computer Science Department, Stanford University, Stanford, CA 94305

**Michail Lagoudakis**

**Ronald Parr**

Department of Computer Science, Duke University, Durham, NC 27708

GUESTRIN@CS.STANFORD.EDU

MGL@CS.DUKE.EDU

PARR@CS.DUKE.EDU

## Abstract

We present several new algorithms for multiagent reinforcement learning. A common feature of these algorithms is a parameterized, structured representation of a policy or value function. This structure is leveraged in an approach we call *coordinated reinforcement learning*, by which agents coordinate both their action selection activities and their parameter updates. Within the limits of our parametric representations, the agents will determine a jointly optimal action without explicitly considering every possible action in their exponentially large joint action space. Our methods differ from many previous reinforcement learning approaches to multiagent coordination in that structured communication and coordination between agents appears at the core of both the learning algorithm and the execution architecture. Our experimental results, comparing our approach to other RL methods, illustrate both the quality of the policies obtained and the additional benefits of coordination.

## 1. Introduction

Consider a system where multiple agents, each with its own set of possible actions and its own observations, must coordinate in order to achieve a common goal. We want to find a mechanism for coordinating the agents' actions so as to maximize their joint utility. One obvious approach to this problem is to represent the system as a Markov Decision Process (MDP), where the "action" is a joint action for all of the agents and the reward is the total reward for all of the agents. The immediate difficulty with this approach is that the action space is quite large: If there are  $g$  agents, each of which can take  $a$  actions, then the action space is  $a^g$ .

One natural approach to reducing the complexity of this problem is to restrict the amount of information that is available to each agent and hope to maximize global welfare by solving local optimization problems for each agent [13]. In some cases, it is possible to manipulate the presentation of information to the agents in a manner that forces local optimizations to imply global optimizations [16]. In general, however, the problem of finding a globally optimal solution for agents with partial information is known to be intractable [2].

Following [8], we present an approach that combines value function approximation with a message passing scheme by

which the agents efficiently determine the jointly optimal action with respect to an approximate value function. Our approach is based on approximating the joint value function as a linear combination of local value functions, each of which relates only to the parts of the system controlled by a small number of agents. We show how such factored value functions allow the agents to find a globally optimal joint action using a very natural message passing scheme. This scheme can be implemented as a negotiation procedure for selecting actions at run time. Alternatively, if the agents share a common observation vector, each agent can efficiently determine the actions that will be taken by all of the collaborating agents without any additional communication.

Given an action selection mechanism, the remaining task is to develop a reinforcement learning algorithm that is capable of producing value functions of the appropriate form. An algorithm for computing such value functions is presented in [8] for the case where the model is known and represented as a factored MDP. This is the first application of these techniques in the context of reinforcement learning, where we no longer require a factored model or even a discrete state space.

We begin by presenting two methods of computing an appropriate value function through reinforcement learning: a variant of  $Q$ -learning and a variant of Least Squares Policy Iteration (LSPI) [11]. We also demonstrate how parameterized value functions of the form acquired by our reinforcement learning variants can be combined in a very natural way with direct policy search methods such as [12, 1, 14, 9]. The same communication and coordination structures used in the value function approximation phase are used in the policy search phase to sample from and update a factored stochastic policy function.

We call our approach *Coordinated Reinforcement Learning*, because structured coordination between agents is used in the core of our learning algorithms and in our execution architectures. Our initial experimental results with LSPI indicate that the message passing action selection mechanism and value function approximation can be combined to produce effective policies and that additional benefits are obtained with agent coordination.

## 2. Cooperative Action Selection

We begin by considering the simpler problem of having a group of agents select a globally optimal joint action to maximize the sum of their individual utility functions. Suppose we have a collection of agents, where each agent  $j$  must choose

an action  $a_j$  from a finite set of possible actions  $\text{Dom}(A_j)$ . We use  $\mathbf{A}$  to denote  $\{A_1, \dots, A_g\}$ . The agents are acting in a space described by a set of state variables,  $\mathbf{X} = \{X_1 \dots X_n\}$ . A state  $\mathbf{x}$  defines a setting  $x_j$  for each variable  $X_j$  and an action  $\mathbf{a}$  defines an action  $a_j \in \text{Dom}(A_j)$  for each agent. The agents must choose the joint action  $\mathbf{a}$  that maximizes the total utility.

In general, the total utility  $Q$  will depend on all state variables  $\mathbf{X}$  and on the actions of all agents  $\mathbf{A}$ . However, in many practical problems, it is possible to approximate the total utility  $Q$  by the sum of local sub-utilities  $Q_j$ , one for each agent. Now, the total utility becomes  $Q = \sum_j Q_j$ . For example, consider the decision process of a section manager in a warehouse. Her local utility  $Q_j$  may depend on the state of the inventory of her section, on her decision of which products to stock up and on the decision of the sales manager over pricing and special offers. On the other hand, it may not depend directly on the actions of the customer support team. However, the decisions of the customer support team will be indirectly relevant, as they may affect the actions of the sales manager.

Computing the action that maximizes  $Q = \sum_j Q_j$  seems intractable *a priori*, as it would require the enumeration of the joint action space of all agents. Fortunately, by exploiting the local structure in the  $Q_j$ -functions through a *coordination graph* we can compute the optimal action very efficiently, with limited communication between agents and limited observability, as proposed in [8]. We repeat the construction here as it will be important throughout this paper.

In our framework, each agent  $j$  has a local utility function  $Q_j$ . An agent's local  $Q$ -function might be influenced by a subset of the state variables, the agent's action and actions of some other agents; we define  $\text{Scope}[Q_j] \subset \mathbf{X} \cup \mathbf{A}$  to be the set of state variables and action variables that influence  $Q_j$ . (We use  $Q_j(\mathbf{x}, \mathbf{a})$  to denote the value of  $Q_j$  applied to the instantiation of the variables in  $\text{Scope}[Q_j]$  within  $\mathbf{x}, \mathbf{a}$ .) The scope of  $Q_j$  can be further divided into two parts: the observable state variables:

$$\text{Observable}[Q_j] = \{X_i \in \mathbf{X} \mid X_i \in \text{Scope}[Q_j]\};$$

and the relevant agent decision variables:

$$\text{Relevant}[Q_j] = \{A_i \in \mathbf{A} \mid A_i \in \text{Scope}[Q_j]\}.$$

This distinction will allow us to characterize the observations each agent needs to make and the type of communication needed to obtain the jointly optimal action, i.e., the joint action choice that maximizes the total utility  $Q = \sum_j Q_j$ . We note that each  $Q_j$  may be further decomposed as a linear combination of functions that involve fewer variables; in this case, the complexity of the algorithm may be further reduced.

Recall that our task is to find a coordination strategy for the agents to maximize  $\sum_j Q_j$  at state  $\mathbf{x}$ . We assume that the agents have full observability of the relevant state variables, i.e., agent  $j$  can observe  $\text{Observable}[Q_j]$ . Given a particular state  $\mathbf{x} = \{x_1, \dots, x_n\}$ , agent  $j$  can instantiate the part of  $Q_j$  that depends on the state  $\mathbf{x}$ , i.e., condition  $Q_j$  on state  $\mathbf{x}$ . Note that each agent only needs to observe the variables in

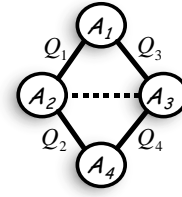


Figure 1. Coordination graph for a 4-agent problem.

$\text{Observable}[Q_j]$ , thereby decreasing considerably the amount of information each agent needs to gather.

After conditioning on the current state, each  $Q_j$  will only depend on the agent's action choice  $A_j$ . Our task is now to select a joint action  $\mathbf{a}$  that maximizes  $\sum_j Q_j(\mathbf{a})$ . The fact that the  $Q_j$ 's depend on the actions of multiple agents forces the agents to coordinate their action choices. We can represent the coordination requirements of the system using a *coordination graph*, where there is a node for each agent and an edge between two agents if they must directly coordinate their actions to optimize some particular  $Q_j$ . Fig. 1 shows the coordination graph for an example where

$$Q = Q_1(a_1, a_2) + Q_2(a_2, a_4) + Q_3(a_1, a_3) + Q_4(a_3, a_4).$$

A graph structure suggests the use of a *cost network* [6], which can be solved using *non-serial dynamic programming* [3] or a variable elimination algorithm which is virtually identical to variable elimination in a Bayesian network. We review this construction here, as it is a key component.

The idea is that, rather than summing all functions and then maximizing, we maximize over variables one at a time. When maximizing over  $a_l$ , only summands involving  $a_l$  participate in the maximization. In our example, we wish to compute:

$$\max_{a_1, a_2, a_3, a_4} Q_1(a_1, a_2) + Q_2(a_2, a_4) + Q_3(a_1, a_3) + Q_4(a_3, a_4).$$

Let us begin our optimization with agent 4. To optimize  $A_4$ , functions  $Q_1$  and  $Q_3$  are irrelevant. Hence, we obtain:

$$\max_{a_1, a_2, a_3} Q_1(a_1, a_2) + Q_3(a_1, a_3) + \max_{a_4} [Q_2(a_2, a_4) + Q_4(a_3, a_4)].$$

We see that to choose  $A_4$  optimally, the agent must know the values of  $A_2$  and  $A_3$ . In effect, it is computing a conditional strategy, with a (possibly) different action choice for each action choice of agents 2 and 3. Agent 4 can summarize the value that it brings to the system in the different circumstances using a new function  $f_4(A_2, A_3)$  whose value at the point  $a_2, a_3$  is the value of the internal max expression. This new function is a new joint value function for agents 2 and 3 (indicated with a dashed line in Fig. 1), summarizing their joint contribution to the total reward under the assumption that agent 4 will act optimally with respect to their choices.

Our problem now reduces to computing

$$\max_{a_1, a_2, a_3} Q_1(a_1, a_2) + Q_3(a_1, a_3) + f_4(a_2, a_3),$$

having one fewer agent. Next, agent 3 makes its decision:

$$\max_{a_1, a_2} Q_1(a_1, a_2) + f_3(a_1, a_2),$$

where  $f_3(a_1, a_2) = \max_{a_3} [Q_3(a_1, a_3) + f_4(a_2, a_3)]$ .

Agent 2 now makes its decision, giving

$$f_2(a_1) = \max_{a_2} Q_1(a_1, a_2) + f_3(a_1, a_2),$$

and agent 1 can now simply choose the action  $a_1$  that maximizes  $f_1 = \max_{a_1} f_2(a_1)$ . The result at this point is a number, which is the desired maximum over  $a_1, a_2, a_3$ , and  $a_4$ .

We can recover the maximizing set of actions by performing the entire process in reverse: the maximizing choice for  $f_1$  selects the action  $a_1^*$  for agent 1. To fulfill its commitment to agent 1, agent 2 must choose the value  $a_2^*$  which maximizes  $f_2(a_1^*)$ . This, in turn, forces agent 3 and then agent 4 to select their actions appropriately.

In general, the algorithm maintains a set  $\mathcal{F}$  of functions, which initially contains  $\{Q_1, \dots, Q_g\}$ . The algorithm then repeats the following steps:

1. Select an uneliminated agent  $A_l$ ;
2. Take all  $f_1, \dots, f_L \in \mathcal{F}$  whose scope contains  $A_l$ .
3. Define a new function  $f = \max_{a_l} \sum_j f_j$  and introduce it into  $\mathcal{F}$ . The scope of  $f$  is  $\cup_{j=1}^L \text{Scope}[f_j] - \{A_l\}$ .

As above, the maximizing action choices are recovered by sending messages in the reverse direction.

The computational cost of this algorithm is linear in the number of new “function values” introduced in the elimination process. More precisely, consider the computation of a new function  $e$  whose scope is  $\mathbf{Z}$ . To compute this function, we need to compute  $|\text{Dom}[\mathbf{Z}]|$  different values. The cost of the algorithm is linear in the overall number of these values, introduced throughout the algorithm. As shown in [6], this cost is exponential in the induced width of the coordination graph for the problem. The algorithm is *distributed* in the sense that the only communication required is between agents that participate in the interior maximizations described above. There is no need for a direct exchange of information between other agents. Thus, the induced tree width has a natural interpretation in this context; it is the maximum number of agents who will need to directly collaborate on the action choice.

The order in which the variables are eliminated will have an important impact on the efficiency of this algorithm. We assume that this is determined *a priori* and known to all agents. It is also possible to devise a simple communication protocol which relaxes the need for a fixed elimination order. In this case, the maximization is performed asynchronously: while an agent is performing its local maximization, its neighbors are “locked”; any agent which is “unlocked” and which has “unlocked” neighbors can start its local maximization step. This process is guaranteed to achieve the same optimal result, although the tree width of the resulting elimination ordering may be larger than that of an ordering which is carefully chosen *a priori*.

At this point, we have shown that if the global utility function  $Q$  is approximated by the sum of local utilities  $Q_j$ , then it is possible to use the coordination graph to compute the maximizing joint action efficiently. In the remainder of this paper, we will show how we can learn these local utilities efficiently.

### 3. Markov Decision Processes

The mechanism described above can be used to maximize not just immediate value, but long term cumulative rewards by using  $Q$ -functions that are derived from the value function of an MDP. The extent to which such a scheme will be successful will be determined by our ability to represent the value function in a form that is usable by our action selection mechanism. Before addressing this question, we first review the MDP framework.

An MDP is defined as a 4-tuple  $(\mathbf{X}, \mathcal{A}, P, R)$  where:  $\mathbf{X}$  is a finite set of states;  $\mathcal{A}$  is a finite set of actions;  $P$  is a *Markovian transition model* where  $P(\mathbf{x}, a, \mathbf{x}')$  represents the probability of going from state  $\mathbf{x}$  to state  $\mathbf{x}'$  with action  $a$ ; and  $R$  is a *reward function*  $R: \mathbf{X} \times \mathcal{A} \times \mathbf{X} \mapsto \mathbb{R}$ , such that  $R(\mathbf{x}, a, \mathbf{x}')$  represents the reward obtained when taking action  $a$  in state  $\mathbf{x}$  and ending up in state  $\mathbf{x}'$ . For convenience, we will sometimes use  $\mathcal{R}(\mathbf{x}, a) = \sum_{\mathbf{x}'} P(\mathbf{x}, a, \mathbf{x}')R(\mathbf{x}, a, \mathbf{x}')$ .

We will be assuming that the MDP has an infinite horizon and that future rewards are discounted exponentially with a discount factor  $\gamma \in [0, 1)$ . A stationary policy  $\pi$  for an MDP is a mapping  $\pi: \mathbf{X} \mapsto \mathcal{A}$ , where  $\pi(\mathbf{x})$  is the action the agent takes at state  $\mathbf{x}$ . The optimal value function  $\mathcal{V}^*$  is defined so that the value of a state must be the maximal value achievable by any action at that state. More precisely, we define:

$$Q_{\mathcal{V}}(\mathbf{x}, a) = R(\mathbf{x}, a) + \gamma \sum_{\mathbf{x}'} P(\mathbf{x}' | \mathbf{x}, a) \mathcal{V}(\mathbf{x}');$$

and the *Bellman operator*  $\mathcal{T}^*$  to be:

$$\mathcal{T}^* \mathcal{V}(\mathbf{x}) = \max_a Q_{\mathcal{V}}(\mathbf{x}, a).$$

The optimal value function  $\mathcal{V}^*$  is the fixed point  $\mathcal{V}^* = \mathcal{T}^* \mathcal{V}^*$ .

For any value function  $\mathcal{V}$ , we can define the policy obtained by acting greedily relative to  $\mathcal{V}$ . In other words, at each state, we take the action that maximizes the one-step utility, assuming that  $\mathcal{V}$  represents our long-term utility achieved at the next state. More precisely, we define  $\text{Greedy}(\mathcal{V})(\mathbf{x}) = \arg \max_a Q_{\mathcal{V}}(\mathbf{x}, a)$ . The greedy policy relative to the optimal value function  $\mathcal{V}^*$  is the optimal policy  $\pi^* = \text{Greedy}(\mathcal{V}^*)$ . It follows immediately from the definition of  $\pi^*$  and is a basic property of MDPs that an agent following  $\pi^*$  maximizes its long term return in an environment. Thus, an agent with knowledge of  $Q_{\mathcal{V}^*}$  can maximize with respect to  $Q_{\mathcal{V}^*}$  (as in Section 2) and achieve optimal long-term return.

Since  $Q_{\mathcal{V}^*}$  is (exponentially) difficult to compute exactly for large MDPs, we use approximation to find local utilities,  $Q_j$ , such that  $\sum_j Q_j$  is a good approximation of  $Q_{\mathcal{V}^*}$ . Recall that we are interested in computing local utilities  $Q_j$  for each agent that will represent an approximation to the global utility  $Q$ . In [8], we presented an algorithm for computing such value functions for the case where the model is known and represented as a factored MDP. In this paper, we consider the case of unknown reward and transition models, i.e., the reinforcement learning case. In the next three sections, we will present alternative, and complementary, approaches for coordinating agents in order to learn the local  $Q_j$ -functions.

#### 4. Coordination Structure in $Q$ -learning

$Q$ -learning is a standard approach to solving an MDP through reinforcement learning. In  $Q$ -learning the agent directly learns the values of state-action pairs from observations of quadruples of the form (state, action, reward, next-state), which we will henceforth refer to as  $(\mathbf{x}, a, r, \mathbf{x}')$ . For each such quadruple,  $Q$ -learning performs the following update:

$$Q(\mathbf{x}, a) \leftarrow Q(\mathbf{x}, a) + \alpha[r + \gamma V(\mathbf{x}') - Q(\mathbf{x}, a)],$$

where  $\alpha$ , is the ‘‘learning rate,’’ or step size parameter and  $V(\mathbf{x}') = \max_a Q(\mathbf{x}', a)$ . With a suitable decay schedule for the learning rate, a policy that ensures that every state-action pair is experienced infinitely often and a representation for  $Q(\mathbf{x}, a)$  which can assign an independent value to every state-action pair,  $Q$ -learning will converge to estimates for  $Q(\mathbf{x}, a)$  which reflect the expected, discounted value of taking action  $a$  in state  $\mathbf{x}$  and proceeding optimally thereafter.

In practice the formal convergence requirements for  $Q$ -learning almost never hold because the state space is too large to permit an independent representation of the value of every state. Typically, a parametric function approximator such as a neural network is used to represent the  $Q$  function for each action. The following gradient-based update scheme is used:

$$\mathbf{w} \leftarrow Q(\mathbf{x}, a, \mathbf{w}) + \alpha[r + \gamma V(\mathbf{x}') - Q(\mathbf{x}, a, \mathbf{w})] \nabla_{\mathbf{w}} Q(\mathbf{x}, a, \mathbf{w}), \quad (1)$$

where  $\mathbf{w}$  is a weight vector for our function approximation architecture and, again, the value  $V(\mathbf{x}')$  of the next state is:

$$V(\mathbf{x}') = \max_a Q(\mathbf{x}', a, \mathbf{w}). \quad (2)$$

The  $Q$ -learning update mechanism is completely generic and requires only that the approximation architecture is differentiable. We are free to choose an architecture that is compatible with our action selection mechanism. Therefore, we can assume that every agent maintains a local  $Q_j$ -function defined over some subset  $\mathbf{x}_j$  of the state variables (which can be the entire state  $\mathbf{x}$ ) and a subset  $\mathbf{a}_j$  of the action variables  $\mathbf{a}$  (its own actions and possibly actions of some other agents). The global  $Q$ -function is now a function of the global state  $\mathbf{x}$  and the joint action vector  $\mathbf{a}$ :

$$Q(\mathbf{x}, \mathbf{a}, \mathbf{w}) = \sum_{j=1}^g Q_j(\mathbf{x}_j, \mathbf{a}_j, \mathbf{w}_j).$$

The  $Q_j$ 's can be maintained locally by each agent as an arbitrary, differentiable function of a set of local weights  $\mathbf{w}_j$ .

There are some somewhat subtle consequences of this representation. The first is that determining  $V(\mathbf{x}')$  in Eq. (2) seems intractable, because it requires a maximization over an exponentially large action space. Fortunately, the  $Q$ -function is factored as a linear combination of local  $Q_j$ -functions, where each  $Q_j$  depends on a subset  $\mathbf{a}_j$  of  $\mathbf{a}$ . Thus, we can apply the coordination graph procedure from Sec. 2 to obtain the maximum value  $V(\mathbf{x}')$  at any given state  $\mathbf{x}'$ .

Once we have defined the local  $Q_j$ -functions, we must compute the weight update in Eq. (1). Each agent must know

$$\Delta(\mathbf{x}, \mathbf{a}, r, \mathbf{x}', \mathbf{w}) = [r + \gamma V(\mathbf{x}') - Q(\mathbf{x}, \mathbf{a}, \mathbf{w})], \quad (3)$$

the difference between the current  $Q$ -value and the discounted value of the next state. We have just shown that it is possible to apply the coordination graph from Sec. 2 to compute  $V(\mathbf{x}')$ . The other unknown terms in Eq. (3) are the reward  $r$  and the previous  $Q$ -value  $Q(\mathbf{x}, \mathbf{a}, \mathbf{w})$ . The reward is observed and the  $Q$ -value is computed by a simple message passing scheme similar to the one in the coordination graph by fixing the action of every agent to the one assigned in  $\mathbf{a}$ .

Therefore, after the coordination step, each agent will have access to the value of  $\Delta(\mathbf{x}, \mathbf{a}, r, \mathbf{x}', \mathbf{w})$ . At this point, the weight update equation is entirely local:

$$\mathbf{w}_i \leftarrow Q(\mathbf{x}, \mathbf{a}, \mathbf{w}) + \alpha \Delta(\mathbf{x}, \mathbf{a}, r, \mathbf{x}', \mathbf{w}) \nabla_{\mathbf{w}_i} Q_i(\mathbf{x}_j, \mathbf{a}_j, \mathbf{w}_i),$$

The reason is simply that the gradient decomposes linearly: once an action is selected, there are no cross-terms involving any  $\mathbf{w}_i$  and  $\mathbf{w}_j$ . The locality of the weight updates in this formulation of  $Q$ -learning make it very attractive for a distributed implementation. Each agent can maintain an entirely local  $Q$ -function and does not need to know anything about the structure of the neighboring agents'  $Q$ -functions. Different agents can even use different architectures, e.g., one might use a neural network and another might use a CMAC. The only requirement is that the joint  $Q$ -function be expressed as a sum of these individual  $Q$ -functions.

The only negative aspect of this  $Q$ -learning formulation is that, like almost all forms of  $Q$ -learning with function approximation, it is difficult to provide any kind of formal convergence guarantees.

#### 5. Multiagent LSPI

Least Squares policy iteration (LSPI) [11] is a new reinforcement learning method that performs policy iteration by using a stored corpus of samples instead of a model. LSPI represents  $Q$ -functions using as a linear combination of basis functions. Given a policy,  $\pi_i$ , LSPI computes a  $Q$ -function,  $Q^{\pi_i}$  (in the space spanned by the basis functions) which is a fixed point for  $\pi_i$  with respect to the samples. The new  $Q^{\pi_i}$  then implicitly defines policy  $\pi_{i+1}$  and the process is repeated until some form of convergence is achieved.

We briefly review the mathematical operations required for LSPI. We assume that our  $Q$ -functions will be approximated by a linear combination of basis functions (features),

$$Q^\pi(\mathbf{x}, a, w) = \sum_{i=1}^k \phi_i(\mathbf{x}, a) w_i = \phi(\mathbf{x}, a)^\top w,$$

For convenience we express our basis in matrix form:

$$\Phi = \begin{pmatrix} \phi(\mathbf{x}_1, a_1)^\top \\ \vdots \\ \phi(\mathbf{x}, a)^\top \\ \vdots \\ \phi(\mathbf{x}_{|\mathbf{X}|}, a_{|\mathbf{A}|})^\top \end{pmatrix}.$$

where  $\Phi$  is  $(|\mathbf{X}||\mathbf{A}| \times k)$ . If we knew the transition matrix,  $\mathbf{P}^\pi$ , for the current policy and knew the reward function we could, in principle, compute the fixed point by defining and solving the system  $\mathbf{A}w^\pi = b$ , where  $\mathbf{A} = \Phi^\top(\Phi - \gamma\mathbf{P}^\pi\Phi)$  and  $b = \Phi^\top\mathcal{R}$ .

In reinforcement learning, we sample experiences from the environment in place of  $\mathcal{R}$  and  $\mathbf{P}^\pi$ . Given a set of samples,  $D = \{s_{d_i}, a_{d_i}, s'_{d_i}, r_{d_i} \mid i = 1, 2, \dots, L\}$ , we can construct approximate versions of  $\Phi$ ,  $\mathbf{P}^\pi \Phi$ , and  $\mathcal{R}$  as follows:

$$\widehat{\Phi} = \begin{pmatrix} \phi(s_{d_1}, a_{d_1})^\top \\ \dots \\ \phi(s_{d_i}, a_{d_i})^\top \\ \dots \\ \phi(s_{d_L}, a_{d_L})^\top \end{pmatrix} \quad \widehat{\mathbf{P}^\pi \Phi} = \begin{pmatrix} \phi(s'_{d_1}, \pi(s'_{d_1}))^\top \\ \dots \\ \phi(s'_{d_i}, \pi(s'_{d_i}))^\top \\ \dots \\ \phi(s'_{d_L}, \pi(s'_{d_L}))^\top \end{pmatrix}$$

$$\widehat{\mathcal{R}} = \begin{pmatrix} r_{d_1} \\ \dots \\ r_{d_i} \\ \dots \\ r_{d_L} \end{pmatrix}$$

Then, we can construct  $\widehat{\mathbf{A}} = \widehat{\Phi}^\top (\widehat{\Phi} - \gamma \widehat{\mathbf{P}^\pi \Phi})$  and  $\widehat{b} = \widehat{\Phi}^\top \widehat{\mathcal{R}}$ . Approximations derived from different sets of samples can be combined additively and this leads to an incremental update rule for  $\widehat{\mathbf{A}}$  and  $\widehat{b}$ . Assume that initially  $\widehat{\mathbf{A}} = 0$  and  $\widehat{b} = 0$ . For a fixed policy, a new sample  $(\mathbf{x}, a, r, \mathbf{x}')$  contributes to the approximation according to the following update equations:

$$\widehat{\mathbf{A}} \leftarrow \widehat{\mathbf{A}} + \phi(\mathbf{x}, a) \left( \phi(\mathbf{x}, a) - \gamma \phi(\mathbf{x}', \pi(\mathbf{x}')) \right)^\top$$

$$\widehat{b} \leftarrow \widehat{b} + \phi(\mathbf{x}, a) r.$$

The solution  $\widehat{w}^\pi = \widehat{\mathbf{A}}^{-1} \widehat{b}$  approximates the true solution  $w^\pi$ .

We note that this approach is very similar to the LSTD algorithm [5]. Unlike LSTD, which defines a system of equations relating state values to state values, LSPI is defined over  $Q$ -values. Each iteration of LSPI yields the  $Q$ -values for the current policy. Thus, each solution implicitly defines the next policy for policy iteration.

An important feature of LSPI is that it is able to reuse the same set of samples even as the policy changes. For example, suppose the corpus contains a transition from state  $\mathbf{x}$  to state  $\mathbf{x}'$  under action  $a_1$  and  $\pi_i(\mathbf{x}') = a_2$ . This is entered into the  $\widehat{\mathbf{A}}$  matrix as if a transition were made from  $(\mathbf{x}, a_1)$  to  $(\mathbf{x}', a_2)$ . If  $\pi_{i+1}(\mathbf{x}')$  changes the action for  $\mathbf{x}'$  from  $a_2$  to  $a_3$ , then the next iteration of LSPI enters a transition from  $(\mathbf{x}, a_1)$  to  $(\mathbf{x}', a_3)$  into the  $\widehat{\mathbf{A}}$  matrix. The sample can be reused because the dynamics for state  $\mathbf{x}$  under action  $a_1$  have not changed.

The application of collaborative action selection to the LSPI framework is surprisingly straightforward. We first note that any set of  $Q$ -functions produced by LSPI will, by construction, be of the right form for collaborative action selection. Each agent is assigned a local set of basis functions which define its local  $Q$ -function. These basis functions can be defined over the agent's own actions as well as the actions of a small number of other agents. As with ordinary LSPI, the current policy  $\pi_i$  is defined implicitly by the current set of  $Q$ -functions,  $Q^{\pi_i}$ . However, in the multiagent case, we cannot enumerate each possible action to determine the policy at some given state because this set of actions is exponential in the number of agents. Fortunately, we can again exploit the structure of the coordination graph to determine the optimal actions relative to  $Q^{\pi_i}$ : for each transition from state  $\mathbf{x}$  to

state  $\mathbf{x}'$  under joint action  $\mathbf{a}$  the coordination graph is used to determine the optimal joint action  $\mathbf{a}'$  for  $\mathbf{x}'$ . The transition is added to the  $\widehat{\mathbf{A}}$  matrix as a transition from  $(\mathbf{x}, \mathbf{a})$  to  $(\mathbf{x}', \mathbf{a}')$ .

An advantage of the LSPI approach to collaborative action selection is that it computes a value function for each successive policy which has a coherent interpretation as a projection into the linear space spanned by the individual agent's local  $Q$ -functions. Thus, there is reason to believe that if this space is expressive enough to approximate closely the true value function, coordinated action selection will make action choices similar to those of the greedy policy with respect to the true value function.

A disadvantage of LSPI is that it is not currently amenable to a distributed implementation during the learning phase: Construction of the  $\mathbf{A}$  matrix requires knowledge of the evaluation of each agent's basis functions for every state in the corpus, not only for every action that is actually taken, but for every action recommended by every policy considered.

## 6. Coordination in Direct Policy Search

Value function based reinforcement learning methods have recently come under some criticism as being unstable and difficult to use in practice. A function approximation architecture that is not well-suited to the problem can diverge or produce poor results with little meaningful feedback that is directly useful for modifying the function approximator to achieve better performance.

LSPI was designed to address some of the concerns with  $Q$ -learning based value function approximation. It is more stable than  $Q$ -learning and is more transparent, thus easier to debug. However, LSPI is still an approximate policy iteration procedure and can be quite sensitive to small errors in the estimated  $Q$ -values for policies [4]. In practice, LSPI can take large, coarse steps in policy space.

The shortcomings of value function based methods have led to a surge of interest in direct policy search methods [12, 1, 14, 9]. These methods use gradient ascent to search a space of parameterized stochastic policies. As with all gradient methods, local optima can be problematic. Defining a relatively smooth but expressive policy space and finding reasonable starting points within this space are all important elements of any successful application of gradient ascent.

We now show how to seed a gradient ascent procedure with a multiagent policy generated by  $Q$ -learning or LSPI as described above. To guarantee that the gradient exists, policy search methods require stochastic policies. Our first task is to convert the deterministic policy implied by our value  $Q$ -functions into a stochastic, policy  $\mu(\mathbf{a}|\mathbf{x})$ , i.e., a distribution over actions given the state. A natural way to do this, which also turns out to be compatible with most policy search methods, is to create a softmax over the  $Q$ -values:

$$\mu(\mathbf{a} | \mathbf{x}) = \frac{e^{\frac{1}{T} \sum_j Q_j(\mathbf{x}, \mathbf{a})}}{\sum_{\mathbf{b}} e^{\frac{1}{T} \sum_k Q_k(\mathbf{x}, \mathbf{b})}}; \quad (4)$$

where  $T$  is a temperature parameter indicating how stochastic we want to make the initial policy. For simplicity of presen-

tation, we will use  $T = 1$ . To be able to apply policy search methods for such policy representation, we must present two additional steps. The first is a method of efficiently generating samples from our stochastic policy and the second is a method of efficiently differentiating our stochastic policy for gradient ascent purposes.

Sampling from our stochastic policy may appear problematic because of the size of the joint action space. For sampling purposes, we can ignore the denominator, since it is the same for all actions, and sample from the numerator directly as an unnormalized potential function. To do this sampling we again use variable elimination on a coordination graph with exactly the same structure as the one in Sec. 2. Conditioning on the current state  $\mathbf{x}$  is again easy: each agent needs to observe only the variables in  $Observable[Q_j]$  and instantiate  $Q_j$  appropriately. At this point, we need to generate a sample from  $Q_j$ -functions that depend only on the action choice.

Following our earlier example, our task is now to sample from the potential corresponding to the numerator of  $\mu(\mathbf{a} | \mathbf{x})$ . Suppose, for example, that the individual agent's  $Q$ -functions have the following form:

$$Q = Q_1(a_1, a_2) + Q_2(a_2, a_4) + Q_3(a_1, a_3) + Q_4(a_3, a_4).$$

and we wish to sample from the potential function for

$$e^{Q_1(a_1, a_2)} e^{Q_2(a_2, a_4)} e^{Q_3(a_1, a_3)} e^{Q_4(a_3, a_4)}.$$

To sample actions one at a time, we will follow a strategy of marginalizing out actions until we are left with a potential over a single action. We then sample from this potential and propagate the results backwards to sample actions for the remaining agents. Suppose we begin by eliminating  $a_4$ . Agent 4 can summarize its impact on the rest of the distribution by combining its potential function with that of agent 2 and defining a new potential:

$$f_4(A_2, A_3) = \sum_{A_4} e^{Q_2(a_2, a_4)} e^{Q_4(a_3, a_4)}.$$

The problem now reduces to sampling from

$$e^{Q_1(a_1, a_2)} e^{Q_3(a_1, a_3)} f_4(a_2, a_3),$$

having one fewer agent. Next, agent 3 communicates its contribution giving:

$$f_3(a_1, a_2) = \sum_{a_3} e^{Q_3(a_1, a_3)} f_4(a_2, a_3).$$

Agent 2 now communicates its contribution, giving

$$f_2(a_1) = \sum_{a_2} e^{Q_1(a_1, a_2)} f_3(a_1, a_2),$$

and agent 1 can now sample actions from the potential  $P(a_1) \sim f_2(a_1)$ .

We can now sample actions for the remaining agents by reversing the direction of the messages and sampling from the distribution for each agent, conditioned on the choices of the previous agents. For example, when agent 2 is informed of the action selected by agent 1, agent 2 can sample actions from the distribution:

$$P(a_2 | a_1) = \frac{P(a_2, a_1)}{P(a_1)} = \frac{e^{Q_1(a_1, a_2)} f_3(a_1, a_2)}{f_2(a_1)}.$$

The general algorithm has the same message passing topology as the original action selection mechanism. The only difference is the content of the messages: the forward pass messages are probability potentials and the backward pass messages are used to compute conditional distributions from which actions are sampled.

The next key operation is the computation of the gradient of our stochastic policy function, a key operation in a REINFORCE style [15] policy search algorithm.<sup>1</sup> First, recall that the global  $Q$ -function is the sum of the local  $Q_j$ -functions:

$$Q(\mathbf{x}, \mathbf{a}, \mathbf{w}) = \sum_{j=1}^g Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j).$$

Our stochastic policy representation now becomes:

$$\mu(\mathbf{a} | \mathbf{x}) = \frac{e^{\sum_j Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j)}}{\sum_{\mathbf{b}} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)}}.$$

We can now compute the gradient of the log policy:

$$\begin{aligned} \nabla_{\mathbf{w}_j} \ln \mu(\mathbf{a} | \mathbf{x}) &= \\ &= \nabla_{\mathbf{w}_j} \ln \left( \frac{e^{\sum_j Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j)}}{\sum_{\mathbf{b}} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)}} \right); \\ &= \nabla_{\mathbf{w}_j} \ln e^{\sum_j Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j)} - \nabla_{\mathbf{w}_j} \ln \sum_{\mathbf{b}} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)}; \\ &= \nabla_{\mathbf{w}_j} \sum_j Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j) - \frac{\sum_{\mathbf{b}} \nabla_{\mathbf{w}_j} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)}}{\sum_{\mathbf{b}'} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}', \mathbf{w}_j)}}; \\ &= \nabla_{\mathbf{w}_j} Q_j(\mathbf{x}, \mathbf{a}, \mathbf{w}_j) - \frac{\sum_{\mathbf{b}} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)} \nabla_{\mathbf{w}_j} Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)}{\sum_{\mathbf{b}'} e^{\sum_j Q_j(\mathbf{x}, \mathbf{b}', \mathbf{w}_j)}}. \end{aligned}$$

We note that both the numerator and the denominator in the summation can be determined by a variable elimination procedure similar to the stochastic action selection procedure. Specifically, the denominator can be written as  $\sum_{\mathbf{b}'} \prod_j e^{Q_j(\mathbf{x}, \mathbf{b}', \mathbf{w}_j)}$ . The variable elimination procedure for computing this sum of products is exactly the same as the forward pass used for sampling from the policy. We can write the numerator as  $\sum_{\mathbf{b}} \left( \prod_j e^{Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)} \right) \nabla_{\mathbf{w}_j} Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)$ . The procedure for computing this value is exactly the same as the denominator, except that one extra potential is introduced, namely  $\nabla_{\mathbf{w}_j} Q_j(\mathbf{x}, \mathbf{b}, \mathbf{w}_j)$ .

These action selection and gradient computation mechanisms provide the basic functions required for essentially any policy search method. As in the case of  $Q$ -learning, a global error signal must be shared by the entire set of agents. Apart from this, the gradient computations and stochastic policy sampling procedure involve a message passing scheme with the same topology as the action selection mechanism. We believe that these methods can be incorporated into any of a number of policy search methods to fine tune a policy derived by  $Q$ -learning with linear  $Q$ -functions or by LSPI.

<sup>1</sup>Most policy search algorithms are of this style.

## 7. Experimental Results

We validated our coordinated RL approach on two domains: the multiagent SysAdmin [8] and the power grid [13]. In the SysAdmin problem, there is a network of computers; each is associated with an administrator agent. Each machine runs processes and receives a reward if a process terminates. Processes take longer to terminate on faulty machines and dead machines can send bad packets to neighbors, causing them to become faulty and eventually die. Each machine is associated with an agent  $A_j$ , which, at each step, decides whether or not to reboot the machine. Rebooting a machine makes its status good with probability 1, but running processes are lost. These agents have to coordinate their actions so as to maximize the total reward for the system, or in other words, maximize the total number of successfully completed jobs in the system. Thus, agents must coordinate to satisfy two, potentially conflicting, goals: running their own processes to termination and not sending bad packets to neighboring machines, as they can eventually cause the failure of the entire network. For a network of  $n$  machines, the number of states in this MDP is  $9^n$  (3 status levels  $\times$  3 load levels per computer) and the joint action space contains  $2^n$  actions.

We implemented our multiagent LSPI algorithm and tested it on a variety of network topologies, as defined in [7]. Fig. 2 shows the estimated value of the resulting policies for problems with increasing number of agents. For comparison, we also plot the results reported by Guestrin et al. [8] for three other methods: their LP-based (LP) approach; and Schneider et al.’s [13] Distributed Reward (DR) and Distributed Value Function (DVF) algorithms. We also plot the “utopic maximum value”, a loose upper bound on the value of the optimal policy [8]. Note, the LP-based approach is a planning algorithm, i.e., uses full knowledge of the (factored) MDP model. On the other hand, coordinated RL, DR and DVF are all model-free reinforcement learning approaches.

In our experiments, we created two sets of multiagent LSPI basis functions corresponding to the backprojections of the “single” or “pair” indicator functions from [8]. For  $n$  machines, we found that about  $600n$  samples are sufficient for multiagent LSPI to learn a good policy. Samples were collected by starting at the initial state (with all working machines) and following a purely random policy. To avoid biasing our samples too heavily by the stationary distribution of the random policy, each episode was truncated at 15 steps. Thus, samples were collected from  $40n$  episodes each one 15 steps long. The resulting policies were evaluated by averaging performance over 20 100-step long runs. The entire experiment was repeated 10 times with different sample sets and the results were averaged.

The results in all cases clearly indicate that multiagent LSPI learns very good policies comparable to the LP approach using the same basis functions, but *without* any use of the model. Note that these policies are near-optimal, as their values are very close to the upper bound on the value of the optimal policy. It is worth noting that the number of samples used in each case grows linearly in the number of agents, whereas the joint state-action space grows exponentially. For example,

a problem with 15 agents has over 205 trillion states and 32 thousand possible actions, but required only 9000 samples.

We also tested our multiagent LSPI approach on the power grid domain of Schneider et al. [13]. Here, the grid is composed of a set of nodes. Each node is either a *provider* (a fixed voltage source), a *customer* (with a desired voltage) or a *distributor* (where control takes place). Links between nodes are associated with resistances and no customer is connected directly to a provider. The distributors must set the resistances attached to them to meet the demand of the customers. If the demand of a particular customer is not met, then the grid incurs a cost equal to the demand minus the supply. The total cost is the sum of all costs for all customers. At every time step, each distributor decides whether to *double*, *halve*, or *maintain* (3 possible actions) the value of the resistance at each of its links (6 possible resistance levels). If two distributors are linked, they share the same resistance and their action choices may conflict. In such case, a simple conflict resolution schema is applied (see [13] for details).

Schneider et al. [13] applied a set of algorithms, including DR and DFV, to this problem. In their set up, each distributor is an agent that observes a set of state variables directly related to itself and all its neighbors, and makes a local decision for its links. Therefore, the complexity of each agent depends on the number of its neighbors. We applied our multiagent LSPI algorithm to the same problem with one agent for each endpoint of a link on a distributor node. Thus, we end up with more, but simpler and identical, agents. Two simple types of state-action basis functions were used: NOCOMM, which has indicators for each assignment of the state of the resistor and the action choice, giving a total of 9 indicator bases for each agent; and, PAIRCOMM, which has indicator bases for each assignment of the resistance level, action of agent  $i$ , and action of agent  $j$  for each pair  $(i, j)$  of directly connected agents, giving a total of 27 indicator bases for each pair of agents. Thus, our agents observe a much smaller part of the state space than those of Schneider et al. [13]. The average costs incurred by the resulting policies, along with results for a uniformly random policy, are shown in Table 1. Multiagent LSPI used only 10,000 training samples for each run (as opposed to 60,000 used in [13]) and the resulting policies were tested for 60,000 steps (same as in [13]). The multiagent LSPI results with the NOCOMM basis set are sub-optimal. With some exceptions, most policies were close to random and the resulting average cost was high (with large confidence intervals). However, the very simple pairwise coordination strategy obtained from the PAIRCOMM basis set yielded near-optimal policies. These agents incur a lower average cost than the DR and DVF agents for all grids using less training data and observing a smaller part of the state space.

## 8. Conclusions and Future Work

We proposed a new approach to reinforcement learning: *Coordinated RL*. In this approach, agents make coordinated decisions and share information to achieve a principled learning strategy. Our method successfully incorporates the cooperative action selection mechanism derived in [8] into the rein-

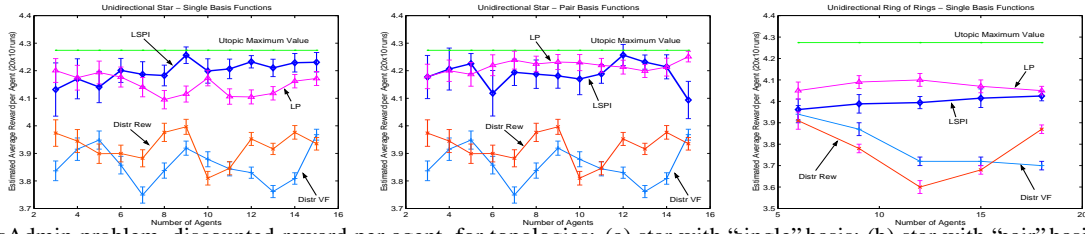


Figure 2. SysAdmin problem, discounted reward per agent, for topologies: (a) star with “single” basis; (b) star with ‘pair’ basis; (c) ring of rings with ‘single’ basis. LP, DR and DRF results as reported in [8].

Grid (see [13])	Random	DR	DVF	Multiagent LSPI	
				NOCOMM	PAIRCOMM
A	29.70 ± 0.13	41.00 ± 0.30	17.17 ± 5.87	28.19 ± 8.30	0.08 ± 0.01
B	52.00 ± 0.24	0.65 ± 0.57	0.32 ± 0.07	37.08 ± 21.85	0.13 ± 0.02
C	96.80 ± 0.31	90.00 ± 1.78	44.00 ± 8.75	83.32 ± 16.57	40.86 ± 1.14
D	44.14 ± 0.37	0.32 ± 0.19	0.17 ± 0.02	28.45 ± 17.83	0.11 ± 0.02

Table 1. Power grid problem: average cost over 10 runs of 60000 steps and 95% confidence intervals. DR and DVF results as reported in [13].

forcement learning framework to allow for structured communication between agents, each of which has only partial access to the state description. We believe the coordination mechanism can be applied to almost any reinforcement learning method. In this paper we applied the Coordinated RL approach to  $Q$ -learning, LSPI, and policy search. With  $Q$ -learning and policy search, the learning mechanism can be distributed. Agents communicate reinforcement signals, utility values, and conditional policies. In LSPI some centralized coordination is required to compute the projection of the value function. The resulting policies can always be executed in a distributed manner. A feature of our method is that the structure of the communication between agents is not fixed a priori, but derived directly from the value function or policy architecture. In our view, an algorithm such as LSPI can provide an offline estimate of the  $Q$ -functions. Subsequently,  $Q$ -learning or direct policy search can be applied online to refine this estimate. By using our Coordinated RL method, we can smoothly shift between these two phases.

Our method can be applied to maximize long term return for any MDP. As with all value function approximation methods, there is a tradeoff between accuracy and complexity. Our approach will be most advantageous when the true  $Q$ -function can be approximated reasonably by a linear combination of local  $Q$ -functions defined over subsets of the actions. In our experiments with this type of value function, we reliably learned policies that were comparable to the best policies achieved by other methods and close to the theoretical optimal achievable in our test domains. The amount of data required scaled linearly with the number of state and action variables even though the state and action spaces were growing exponentially. Furthermore, we demonstrated that coordination can significantly improve the quality of the policies obtained.

Our experiments involved discrete state spaces and were chosen primarily to compare learning performance with previous closed-form approximation methods. Our basis functions match closely the basis functions used in previous work. In future work, we plan to use continuous variables and basis functions. While our methods require discrete actions, they generalize immediately to continuous state variables.

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