Q-functionals for Value-Based Continuous Control

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Abstract

We present Q-functionals, an alternative architecture for continuous control deep reinforcement learning. Instead of returning a single value for a state-action pair, our network transforms a state into a function that can be rapidly evaluated in parallel for many actions, allowing us to efficiently choose high-value actions through sampling. This contrasts with the typical architecture of off-policy continuous control, where a policy network is trained for the sole purpose of selecting actions from the Q-function. We represent our action-dependent Q-function as a weighted sum of basis functions (Fourier, Polynomial, etc) over the action space, where the weights are state-dependent and output by the Q-functional network. Fast sampling makes practical a variety of techniques that require Monte-Carlo integration over Q-functions, and enables action-selection strategies besides simple value-maximization. We characterize our framework, describe various implementations of Q-functionals, and demonstrate strong performance on a suite of continuous control tasks.

Introduction

The ultimate product of a successful reinforcement learning system is a policy that performs well in interacting with the environment, as measured by expected cumulative discounted reward. A dominant paradigm in reinforcement learning is to derive policies from a Q-function (Watkins and Dayan 1992) that estimates the expected cumulative reward for taking a given action. When the number of actions available is finite, a strong policy can be derived by enumerating all action-values for a given state and choosing the one with the highest value. However, enumeration is impossible when there are large or infinite possible actions, for example when actions are drawn from a continuous vector space. This is the natural description of, for example, robotic locomotion and control; significant effort has therefore gone into alternative approaches for action-selection in these domains (Gu et al. 2016; Asadi et al. 2021; Lillicrap et al. 2015). A common framework for so-called continuous control problems is to train a separate policy network that selects actions according to some criteria of the Q-values (Fujimoto, van Hoof, and Meger 2018; Haarnoja et al. 2018a).

The gradient update for a policy network comes completely from the Q-function, and as such, it essentially summarizes information about the Q-function for a given state so as to efficiently produce high-value actions. For example, standard practice is to train a policy network to estimate the maximum-valued action for any given state (Lillicrap et al. 2015). The quantities policy networks estimate are difficult to calculate on a per-state basis, so policy networks can be thought of as amortizing the expensive computation of finding desirable actions: the training procedure is expensive, but it allows actions to be identified with a single pass of a neural network. There are two potential pitfalls of this framework. First, since the policy network is not reinitialized after every Q-function update, at any given timestep most of the policy training has been done on old versions of the Q-function. The policy network is therefore maximizing a stale estimate of action-value. Second, policies are generally either deterministic or sampled from a tight distribution around the single highest valuable action. Therefore, it is unlikely for the policy to sample two high-value actions if they are too far apart from each other (Tessler, Tennenholtz, and Mannor 2019). This may limit exploration as well as the robustness of the value function. Perhaps most importantly, we show later that even a well-trained policy network is less effective than previously thought at its assigned task of maximizing Q-values.

We take a different perspective on this problem: instead of training a network to amortize expensive action-value computations, we propose a network architecture that ensures these evaluations are cheap to do in parallel. This opens up a new avenue for action-selection: instead of using a policy network, we can simply evaluate many actions in parallel and choose from them one with high value. Since the policy is directly derived from Q-values at each timestep it always reflects the most current action-value estimates. In addition, random sampling can easily generate high-value actions from throughout the action-space.

We introduce a class of Q-functions we call Q-functionals that allow for efficient sampling. A functional is a function that returns another function; as such, a Q-functional transforms a state into a function over only the action space, such that Q-values for many actions can be evaluated in parallel...
and with little overhead. Specifically, we compute Q-values as the dot product between state-dependent coefficients and an expressive basis representation of an action; thus, evaluating multiple actions for a state reduces to a single matrix multiplication. We describe a variety of implementations of this architecture, investigate its speed and effectiveness at sampling and action-maximization, and demonstrate competitive performance on a suite of continuous control reinforcement learning tasks.

Background

This work examines sequential decision making problems represented as Markov Decision Processes (MDPs) denoted by \( \langle S, A, T, R, P_0, \gamma \rangle \), where \( S \), \( A \) and \( \gamma \) are the state space, action space, and the discount factor, respectively (Sutton and Barto 2018). The transition and reward functions are given by \( T(s, a) \) and \( R(s, a) \) respectively, and \( P_0(s) \) is the initial state distribution. We seek to learn an action-selection strategy, or policy \( \pi : S \rightarrow A \), resulting in high cumulative discounted reward.

For a given policy \( \pi \), the state-action value function, or Q-function, is defined as:

\[
Q^{\pi}(s_t, a_t) = \mathbb{E}_{\tau}[R(s_t, a_t) + \gamma V^{\pi}(s_{t+1})],
\]

where \( V^{\pi}(s) \) is the expected value of following policy \( \pi \) from state \( s \):

\[
V^{\pi}(s) = \mathbb{E}_{a \sim \pi(s)}[Q^{\pi}(s, a)].
\]

We also investigate \( \tau \)-Entropy-Regularized RL (ER-RL), in which case the value of a state is regularized by the entropy of the current policy (Schulman, Chen, and Abbeel 2017; Haarnoja et al. 2018b):

\[
V^{\pi}_{\text{ENT}}(s) = \mathbb{E}_{a \sim \pi(s)}[Q^{\pi}(s, a) - \tau \log(\pi(a|s))],
\]

where \( \tau \) controls the degree of regularization.

When we are interested in maximizing cumulative reward, we can derive a policy from a Q-function by choosing the maximum-valued action at every state:

\[
\pi^{Q}(s) = \arg \max_a Q(s, a).
\]

A common method for iteratively improving a Q function parameterized by \( \theta \) is through bootstrapping (Sutton 1988):

\[
\theta \leftarrow \theta + \alpha \delta Q(s, a; \theta)
\]

where \( \delta = r + \gamma V^{\pi}(s') - Q^{\pi}(s, a; \theta), \) (1)

where \( \alpha \) is the step-size and \( (s, a, r, s') \) tuples are drawn from experience. The fixed point of this update equation is \( Q^* \), the Q-function that describes the optimal policy \( \pi^* \):

\[
Q^*(s_t, a_t) = \mathbb{E}_{\pi^*}[R(s_t, a_t) + \gamma V^*(s_{t+1})].
\]

Central to this process is deriving a policy from a Q-function through maximization, or related techniques. When \( A \) is a set of discrete actions, this maximization can be done easily by comparing the Q-value for each action (Mnih et al. 2015). Alternatively, this work focuses on the problem of continuous control, where actions are drawn from a continuous vector space.

Related Work

Continuous-control is concerned with domains that have continuous action spaces; thus, instead of choosing from a finite set of actions, an agent must output a vector of continuous entries at every timestep. This difference necessitates characteristically different agent architectures from discrete-action agents such as DQN.

Policy-gradient (PG) methods are a dominant paradigm of continuous control due to their expressive policies and end-to-end training scheme (Sutton et al. 1999). Modern PG methods represent policies as deep neural networks, that are trained to output high-valued actions using a gradient signal produced by the Q-function. Deterministic PG methods output a single action, while stochastic PG methods generally parameterize a simple distribution (such as Gaussian) that can be sampled during action-selection (Haarnoja et al. 2018a). The output of a “perfectly” trained policy network would maximize the value (as calculated by the Q-function) at any given state. The chief observation motivating our work is that throughout training, this is far from the case.

Standard PG methods require a single evaluation of the policy network to produce an action, and a single evaluation of the Q-network to compute the value of that action. A fruitful line of research involves incorporating sampling methods, which compute the value of multiple actions, for either improved action-selection or better value-estimation. Prior work utilizes the cross-entropy method (CEM), a zero-order sample-based optimization method, to choose actions that have higher value than the original action output by the policy network (Simmons-Edler et al. 2019). The Expected Policy Gradient (EPG) framework reduces variance in the value computation through Monte Carlo integration over the policy’s outputs (Ciosek and Whiteson 2018). Evaluating actions from a distribution around the policy’s output has been found to reduce overfitting of a policy network to a Q-function (Fujimoto, van Hoof, and Meger 2018). All of these methods, however, are limited to evaluating small numbers of action-values in practice, because every computed value requires an additional evaluation of the Q-function’s neural network.

Our method does away with the policy network altogether, putting it in the class of value-function only continuous control. We list other examples from this class for completeness. “Continuous Action Q-Learning” (Millán, Posenato, and Dedieu 2002) uses mixed-integer programming to solve for maximal valued actions (though in practice runtime considerations stop this at approximately maximal) in piecewise-linear value functions (such as neural networks with rectified linear activation functions). “Normalized Advantage Functions” analytically select maximum-valued actions (though in practice run-time considerations stop this at approximately maximal) in piecewise-linear value functions (such as neural networks with rectified linear activation functions). “Normalized Advantage Functions” analytically select maximum-valued actions (though in practice run-time considerations stop this at approximately maximal) in piecewise-linear value functions (such as neural networks with rectified linear activation functions).
and describes a method to select approximately maximum-action values. The primary difference in our approach compared with past value-function only methods is the focus on sampling, which allows for a variety of action-selection strategies besides simple maximization, as well as use for variance-reducing Monte Carlo techniques.

**Q-Functionals**

Our method is based on a simple premise: many of continuous control’s subproblems described above (action selection/maximization, value-estimation, variance reduction) can be improved if we can rapidly compute the value of many actions.

Q-functionals are a way of expressing Q-functions so that many action-values can be evaluated in parallel for a given state. Consider a traditional deep Q-function, represented by a neural network:

\[ Q(s, a) : (S \times A) \rightarrow \mathbb{R}. \]

A standard design for such a Q-function in continuous control is to concatenate states and actions, and then pass the concatenated vectors through a neural network that outputs their estimated values. For this standard architecture, evaluating two action-values for the same state takes twice as many operations as evaluating one. A Q-functional breaks the computation of action-values into two parts: first, a state is transformed into parameters that define a function over the action space. Then, this function is evaluated for the action(s) in question:

\[ Q_{\text{FUNC}}(s, a) : S \rightarrow (A \rightarrow \mathbb{R}). \quad (2) \]

We design these functions such that while the per-state computation may be expensive, the per-action computation is relatively cheap. One simple way to represent these functions is by learning state-dependent coefficients of basis functions over the action space. As a brief example, a one-dimensional polynomial basis function represents a scalar action \( a \) by the vector \([1, a, a^2, \ldots]\). An appropriate choice of basis function also adds an inductive bias to learning action-values: though we may expect values to have complex dependence on state (especially when the state is high-dimensional such as in image representations), in general we expect values to be relatively smooth with respect to the action dimensions. In Figure 1, we visualize the difference between the two architectures as differing places where the actions enter the computation.

**Q-Functional Implementations**

We describe three implementations of Q-functionals using different basis functions. First, a Q-functional calculated using the Fourier basis is shown below in equation 3. The Fourier basis representation of states had great success when used for Q-learning in small state spaces (Konidaris, Osentoski, and Thomas 2011); we rely on the same inductive bias over the moderately-sized action spaces typical of continuous control. \( L \) represents the difference between the max and min action space value allowed, \( x_i(s), y_i(s) \) are the state-dependent coefficients output by the Q-functional network. \( C \) is a matrix of frequencies for the Fourier basis. For a rank \( R \) Q-functional using the Fourier basis and operating in an environment with action dimension \( d \), the full set of frequencies is represented by the Cartesian product \( \{0, 1, 2, \ldots, R\}^d \). In practice however, we find that limiting the set of frequencies to those where the sum of all elements in the frequency is less than or equal to the rank provides sufficient generalization. Formally, \( C = \{C_i = \{0, 1, 2, \ldots, R\}^d \mid \sum_{j=0}^{d} C_{ij} \leq R \} \).

\[ Q_{\text{FUNC}}(s, a) = \sum_{C_i \in C} x_i(s) \sin\left(\frac{\pi}{L} a \cdot C_i\right) + y_i(s) \cos\left(\frac{\pi}{L} a \cdot C_i\right). \quad (3) \]

Crucially, the learned coefficients \( x_i(s) \) and \( y_i(s) \) are computed by the Q-functional neural network using only the state as input. Each action-evaluation for that state then reuses these same coefficients. Thus, Q-functionals can compute many action-values for the same state with only a single neural network evaluation. As demonstrated later, this allows for significantly faster sampling of Q-values.

We can similarly define a second basis over polynomials; we represent our action-value function such that the total polynomial-order is less than some \( R \):

\[ Q_{\text{FUNC}}(s, a) = \sum_{C_i \in C} x_i(s) \prod_{j=0}^{d} a_j^{C_{ij}}. \quad (4) \]

A third implementation of Q-functionals is the Legendre basis. To encourage feature-independence, we can orthogonalize the polynomial basis over \( A \) by extending the Legendre polynomials to multiple dimensions (see the Appendix for details on the Legendre basis). Let \( L_i(x) \) be the \( i \)-th order Legendre polynomial. Then, each frequency vector defines an orthogonal polynomial:

\[ \hat{L}_{C_i}(a) = \prod_{j=0}^{d} L_{C_{ij}}(a_j), \]

and our Q-function can be calculated from learned coefficients:

\[ Q_{\text{FUNC}}(s, a) = \sum_{C_i \in C} x_i(s) \hat{L}_{C_i}(a). \quad (5) \]

Previous work demonstrates that action-values can be analytically maximized for quadratic polynomials over actions

![Figure 1: Network architecture of traditional Q-function (left) and Q-functional (right)](image-url)
The Q-function is computed as the simple maximization.
late state-values with various methods that work better than as described in the next section, sampling allows us to calculate complex, higher-order polynomial Q-functions. In addition, the sampling framework can derive a strong policy from more functions over actions which can be overly limiting. Our ever, Q-functions defined as such are restricted to convex

\(Q(s, a) = \arg \max_{a \in A} Q(s, a)\)

\(V(s) = Q(s, \pi(s))\).

Instead of training a policy network, we can directly estimate the maximum through sampling \(k\) actions:

\(\pi(s) = \arg \max_{a_i \in A^k} Q(s, a_i)\) ;
\(V(s) = \max_{a_i \in A^k} Q(s, a_i)\).

(6)

This sampling scheme is directly analogous to how actions are selected in discrete-action domains. These functions are detailed in Algorithm 1, and can be used as drop-in replacements for backpropogation-trained policy networks in the learning process.

For ER-RL, the optimal policy is proportional to the Boltzmann distribution over Q-values, as this optimally trades off between policy entropy and expected return (Schulman, Chen, and Abbeel 2017). Thus, given a trained Q-function, the optimal policy and associated value are (Haarnoja et al. 2017):

\(\pi(s) = \Pr(a|s) \propto \exp\left\{Q(s, a)/\tau\right\}/\int_A \exp\left\{Q(s, a)/\tau\right\}\)

\(V(s) = \tau \log \left(\int_A \exp\left\{Q(s, a)/\tau\right\}\right)\).

(7)

Due to limitations imposed by common representations of stochastic policies (often constrained to state-dependent normal distributions over actions), this relation between policy and value cannot be achieved by standard policy-parameterizations (Haarnoja et al. 2017). By contrast, we can consistently estimate these qualities through Monte Carlo sampling:

\(\pi(s) = \Pr(a|s) = \frac{1}{k} \sum_{a_i \in A^k} \exp\left\{Q(s, a)/\tau\right\}\)

\(V(s) = \tau \log \left(\frac{1}{k} \sum_{a_i \in A^k} \exp\left\{Q(s, a)/\tau\right\}\right)\).

(7)

In our experiments, we derive a robust policy \(\pi(s)\) and value function \(V(s)\) through a “top-n of k” approach: we evaluate \(k\) random actions, and represent our policy and value by sampling from the best \(n\) actions during interaction with the environment and averaging these \(n\) values during bootstrapping updates for stability.

\(\pi(s) \sim \text{top-n}\left\{Q(s, a_1), ...Q(s, a_k)\right\}\)

\(V(s) = \text{mean}\left[\text{top-n}\left\{Q(s, a_1), ...Q(s, a_k)\right\}\right]\).

(8)

Figure 2: Quality of actions selected by trained policy network, and random sampling, on the LunarLanderContinuous task.
This is a similar strategy to “target policy smoothing” (Fujimoto, van Hoof, and Meger 2018), which computes $V(s)$ as a single-sample mean of the Q-values surrounding the policy-networks output. We find that averaging over many values, instead of simply choosing one, leads to decreased noisiness of $V(s)$ for the Bellman target in the update, and superior performance at maximizing reward.

**Comparisons**

The core claim of this paper is that by structuring our Q-functions as functionals we gain access to superior action-selection strategies that standard architectures cannot efficiently implement: that policy networks do not effectively maximize the Q-function they are trained on, and that Q-functionals allow for significantly larger sampling with the same computational budget. All experiments are done on tasks from the OpenAI Gym continuous control suite (Brockman et al. 2016). Reproducing code can be found at the linked repository 1.

**Evaluating Policy Networks**

We begin by investigating the implicit assumption that policies trained through gradient descent are effective at maximizing the output of the Q-function they are trained on. We can quantify this concept by comparing the Q-value of the policy network’s output with the distribution of Q-values of randomly sampled actions. A well-trained policy network should consistently output actions with higher value than the large majority of random actions. We train three instances of a DDPG agent on the “LunarLanderContinuous-v2” task, and for each one collect a dataset of 10,000 states from on-policy rollouts. For each state, we compute the Q-value of the policy’s output using the agent’s Q-function. We then compare this to the value of 10,000 randomly sampled actions, and determine the fraction of these for which the policy’s output has lower value (Figure 2). The horizontal red line indicates the median action’s value at each state. This shows that in 15% of states, on average the policy network selects an action with lower value than one chosen at random. Furthermore, the policy’s output is only in the top decile of action-values in 5% of states. This enormous gap between a policy network’s implied function and its empirical behavior suggests that alternative action-selection strategies can prove useful.

By contrast, the orange line represents the Q-functional policy from Equation 6, that selects the highest-valued action from a pool of 1,000 random samples. The output of this policy is better than a single random action 99.9% of the time. We include similar figures for a variety of environments in the Appendix.

**Sampling Q-functions Versus Q-functionals**

As demonstrated in the previous section, the standard Q-function architecture can naively evaluate many actions by simply passing the same state to its input many times. However, Figure 3 demonstrates the speedup of using the Q-functional architecture for multiple evaluations. For a single batch of 1024 states, we evaluate an increasing number of actions on the Hopper task (action dimension of 3) for 100 iterations. We find that a rank 3 Legendre Q-functional evaluates actions roughly 3.5 times faster on a single Nvidia 2080-ti GPU than a neural network that takes in both states and action as inputs. The output of this policy is better than a single random action 99.9% of the time. We include similar figures for a variety of environments in the Appendix.

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1Code available at https://github.com/samlobel/q_functionals
action-evaluation is decoupled from the complexity of the basis. Using this procedure, we can evaluate actions using a rank-3 Legendre Q-functional more than 20 times as quickly, and with less than one tenth the memory-footprint, as a standard Q-function. This procedure has minimal to no effect on reinforcement learning behavior, as confirmed in the Appendix, and we use precomputed representations through the remainder of the experiments.

**Experimental Results**

In this section, we favorably compare our method to existing off-policy policy-gradient methods, on both the standard and the entropy-regularized RL objective.

**Benchmarking**

We compare Q-functionals to two popular and strong off-policy policy-gradient baselines:

**Deep Deterministic Policy Gradient (DDPG)** is a deterministic policy gradient method which relies on a deterministic policy for action-selection and bootstrapping (Lillicrap et al. 2015). The policy network is trained to maximize the Q-function using batch gradient descent, and the value-function is calculated from the Q-function using the policy network’s output.

**Twin Delayed DDPG (TD3)** is an extension of DDPG that addresses issues of overestimation bias and overfitting present in the original algorithm (Fujimoto, van Hoof, and Meger 2018). They do this with the addition of two components. To address overestimation bias, the Q-function is parameterized by two networks, and a state’s value is computed as the minimum of both, evaluated on the policy’s output. To address overfitting, TD3 introduces target policy smoothing, which adds Gaussian noise to the policy’s output for use in bootstrapping.

We construct two versions of our architecture for fair comparison, analogous to the two policy gradient methods above. The first computes Q-values using a single network, and chooses the maximum-valued action (Equation 6) to compute values for bootstrapping, analogously to DDPG. Like TD3, the second version computes Q-values using the minimum of two networks to address overestimation bias. To address overfitting, in this version we calculate value using the “top-n of k” strategy detailed in Equation 8.
In Figure 4, we compare these four methods on the OpenAI Gym continuous control suite (Brockman et al. 2016; Todorov, Erez, and Tassa 2012). Tasks have action-dimension ranging from 1 (Pendulum) to 17 (Humanoid). For all benchmark experiments, we use the Legendre basis with rank 3, and use 1,000 samples for action-selection both in bootstrapping and interaction. Details on environments and architectural choices can be found in the Appendix. We find that Q-functionals compare favorably to their policy-gradient analogues across a variety of tasks. Somewhat surprisingly, Q-functionals perform best on Ant and Humanoid, the two tasks with the highest action-dimension. This is contrary to popular wisdom that sampling methods are unsuitable for large vector spaces, and perhaps points to policy gradient methods sharing the same flaw.

**Maximizing Regularized Objectives**

Sampling-based value estimation allows for additional flexibility in representing policies over the action space. Instead of being restricted to policies that are either deterministic (Lillicrap et al. 2015), or parameterize a simple state-dependent distribution (Haarnoja et al. 2018a), Q-functionals can sample actions from arbitrary functions over their Q-values. We demonstrate the utility of this approach to ER-RL, under which the Boltzmann distribution over Q-values $Q(s, a)$ maximizes a state’s value $V(s)$ (Schulman, Chen, and Abbeel 2017).

We compare a rank 3 Q-functional using the Fourier basis to Soft Actor Critic (SAC), a popular continuous control algorithm for solving ER-RL problems (Haarnoja et al. 2018a), on a simple bandit task (Slivkins et al. 2019). The reward for a given action is the sum of two Gaussian functions centered around $-0.5$ and $0.5$. Figure 5 shows the learned policies of both algorithms. The optimal policy under ER-RL, shown in yellow, gives equal weight to each peak of the bimodal distribution. Because the SAC agent represents its policy as a Gaussian distribution squashed to the valid action space, it cannot represent the bimodal nature of the optimal policy. By contrast, using Equation 7 to derive a policy allows the Q-functional agent to fit the general shape of the optimal policy.

**Performance with Increased Sampling**

As the number of samples used to select actions increases, so does the average Q-value of the returned action. In Figure 6, we investigate whether increased sampling also leads to better performance on the Humanoid task. We train Q-functional agents using the configurations from our benchmark experiments, and evaluate their performance as we vary the number of samples used in action-selection. On this task, we see drastic improvement as sampling increases to 1,000, and then continued diminishing returns as it continues to increase.

**Conclusion**

The translation between a Q-function and a policy is a central research question in continuous-control reinforcement learning. In this work, we highlight the substantial room for improvement in this question over the currently accepted method of policy-gradients. We introduce Q-functionals as a general framework for Q-functions that can evaluate many actions in parallel. Q-functionals allow for far greater flexibility in how one derives a policy from a Q-function, and achieve impressive results even on tasks with extremely high action dimensions. In addition, under the Q-functional framework, a variety of sample-based techniques become much more practical. We view the flexibility of this framework as well as its strong empirical performance as an attractive alternative to policy-gradient methods in continuous control.

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References


Technical Appendix: Q-functionals for Value-Based Continuous Control

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Basis Representations

We provide detailed examples of each basis function presented in our paper, for rank two in a two-dimensional action space. For all examples, \( a = [a_1, a_2] \), and each action dimension is bounded by \([-L/2, L/2]\). The six “frequency vectors” for a rank-two basis in a two-dimensional action space are as follows:

\[
\begin{align*}
C_1 &= [0, 0] ; \\
C_2 &= [1, 0] ; \\
C_3 &= [0, 1] ; \\
C_4 &= [2, 0] ; \\
C_5 &= [1, 1] ; \\
C_6 &= [0, 2]
\end{align*}
\]

Fourier Basis

We construct a rank-three Fourier basis for a two-dimensional action space. Recall that for the Fourier basis, we compute Q-values as:

\[
Q_{\text{FUNC}}(s, a) = \sum_{C_i \in \mathcal{C}} x_i(s) \sin \left( \frac{\pi}{L} a \cdot C_i \right) + y_i(s) \cos \left( \frac{\pi}{L} a \cdot C_i \right).
\]

A Fourier basis of rank 2 includes all frequency terms with sum less than or equal to 2. Thus, given the twelve state-dependent, learned, coefficients \( x_i(s) \) and \( y_i(s) \) for \( 1 \leq i \leq 6 \), we can compute the Q-values of any state and action as follows:

\[
Q(s, a) = x_1(s) \sin(0) + y_1(s) \cos(0) \\
+ x_2(s) \sin \left( \frac{\pi}{L} (a_1) \right) + y_2(s) \cos \left( \frac{\pi}{L} (a_1) \right) \\
+ x_3(s) \sin \left( \frac{\pi}{L} (a_2) \right) + y_3(s) \cos \left( \frac{\pi}{L} (a_2) \right) \\
+ x_4(s) \sin \left( \frac{\pi}{L} (2a_1) \right) + y_4(s) \cos \left( \frac{\pi}{L} (2a_1) \right) \\
+ x_5(s) \sin \left( \frac{\pi}{L} (a_1 + a_2) \right) + y_5(s) \cos \left( \frac{\pi}{L} (a_1 + a_2) \right) \\
+ x_6(s) \sin \left( \frac{\pi}{L} (2a_2) \right) + y_6(s) \cos \left( \frac{\pi}{L} (2a_2) \right).
\]

Polynomial Basis

A polynomial basis of rank 2 includes all polynomials of the form \( a_1^i a_2^j \) such that \( i + j \leq 2 \). Thus, given the six state-dependent coefficients \( x_i(s) \) for \( 1 \leq i \leq 6 \), we can compute Q-values as follows:

\[
Q(s, a) = x_1(s) \\
+ x_2(s)a_1 \\
+ x_3(s)a_2 \\
+ x_4(s)a_1^2 \\
+ x_5(s)a_1a_2 \\
+ x_6(s)a_2^2
\]

Legendre Basis

The Legendre polynomials are a set of 1-dimensional polynomials that are orthogonal with respect to the inner product

\[
\langle L_1, L_2 \rangle = \int_{-1}^{1} L_1(x)L_2(x)dx.
\]

The set of Legendre polynomials can be recursively computed through the Gram-Schmidt process (with \( L_0(x) = 1 \)):

\[
L_i = x^i - \sum_{j=0}^{i-1} \frac{\langle x^i, L_j(x) \rangle}{\langle L_j(x), L_j(x) \rangle} L_j(x)
\]

The first few Legendre polynomials are:
\[ \mathcal{L}_0(x) = 1 \]
\[ \mathcal{L}_1(x) = x \]
\[ \mathcal{L}_2(x) = \frac{3}{2}x^2 - \frac{1}{2} \]
\[ \mathcal{L}_3(x) = \frac{5}{2}x^3 - \frac{3}{2}x \]

We use a version of Legendre polynomials scaled to have a maximum absolute value of 1 over the interval \([-1, 1]\). The coefficients for these polynomials are available through the numpy command

```
np.polynomial.legendre.Legendre.basis(order, [-1, 1]).convert(kind=numpy.polynomial.Polynomial).coeffs
```

We extend the Legendre polynomials to multiple dimensions. Coefficient vector \( C \) associates each action dimension with a Legendre polynomial. For example, coefficient vector \([1, 3]\) produces the polynomial

\[ \hat{\mathcal{L}}_{1,3} = \mathcal{L}_1(a_1) \cdot \mathcal{L}_3(a_2) \]

Each element of the extended Legendre polynomials is orthogonal to the rest. If \( \mathbf{a} \) is an \( n \)-dimensional vector, then:

\[
< \hat{\mathcal{L}}_{C_i}, \hat{\mathcal{L}}_{C_j} > = \int \int \hat{\mathcal{L}}_{C_i}(\mathbf{x})\hat{\mathcal{L}}_{C_j}(\mathbf{x})d\mathbf{x}_1 \ldots d\mathbf{x}_n = \prod_{k \leq n} \int \mathcal{L}_{C_{i,k}}(x_k)\mathcal{L}_{C_{j,k}}(x_k)dx_k.
\]

At least one term in the product is 0 unless \( C_{i,k} = C_{j,k} \) for all \( k \), or in other words \( C_i = C_j \). A Legendre basis of rank 2 includes all extended Legendre polynomials \( \mathcal{L}_i(a_1)\mathcal{L}_j(a_2) \) such that \( i + j \leq 2 \). Thus, given the six state-dependent coefficients \( x_i(s) \) for \( 1 \leq i \leq 6 \), we can compute Q-values as follows:

\[
Q(s, a) = x_1(s)\mathcal{L}_0(a_1)\mathcal{L}_0(a_2) + x_2(s)\mathcal{L}_1(a_1)\mathcal{L}_0(a_2) + x_3(s)\mathcal{L}_0(a_1)\mathcal{L}_1(a_2) + x_4(s)\mathcal{L}_2(a_1)\mathcal{L}_0(a_2) + x_5(s)\mathcal{L}_1(a_1)\mathcal{L}_1(a_2) + x_6(s)\mathcal{L}_0(a_1)\mathcal{L}_2(a_2)
\]
Experimental Details and Hyperparameters

We test our method on 8 tasks from the OpenAI Gym continuous control benchmark suite (Brockman et al. 2016)(Todorov, Erez, and Tassa 2012). All benchmark experiments were run on a computing cluster, with each job having access to 1 NVIDIA RTX 3090ti GPU, 4 cores, and 36GB RAM. Table 1 contains the state and action dimension of each.

| Name                               | |S| | |A| |
|------------------------------------|-----|-----|-----|
| Pendulum-v1                        | 3   | 1   |
| Reacher-v2                         | 11  | 2   |
| LunarLanderContinuous-v2           | 8   | 2   |
| BipedalWalker-v3                   | 24  | 4   |
| Hopper-v3                          | 11  | 3   |
| Walker2d-v2                        | 17  | 6   |
| Ant-v3                             | 111  | 8   |
| Humanoid-v2                        | 376  | 17  |

Table 1: Environment Details for Continuous Control Tasks

Q-Functionals

We parameterize our Q-functionals using neural networks, with input dimension |S| and output dimension equal to the number of learned coefficients needed. For Polynomial and Legendre basis functions, rank $r$ and action-dimension $d$ require $\binom{r+d}{d}$ learned coefficients. Fourier basis requires twice this, due to having coefficients for both the sine and cosine terms. All shared hyperparameters for Q-functional methods are listed in Table 2.

Learning rates were chosen from $\{1e^{-3}, 3e^{-4}, 1e^{-4}\}$ by running a subset of tasks for 2 seeds, and choosing the learning rate that performed best on aggregate. The same learning rate is used in all experiments.

Additionally, the Q-functional TD3-analogue implemented the “top-n of k” sampling approach for bootstrapping only, with $n = 0.01k$. For interaction, we use the maximum-valued action for both versions of our method. $n$ was chosen from $\{10, 100, 200\}$ by running a subset of tasks for 2 seeds, and choosing the $n$ that performed best on aggregate. The same $n$ is used in all experiments.

Policy Gradient Methods

We use the implementations of SAC (Haarnoja et al. 2018), TD3 (Fujimoto, van Hoof, and Meger 2018), and DDPG (Lillicrap et al. 2015) from the “Stable Baselines 3” Python library (Raffin et al. 2019). All shared hyperparameters for TD3 and DDPG are listed in Table 3.

Learning rates were chosen from $\{1e^{-3}, 3e^{-4}, 1e^{-4}\}$ by running a subset of tasks for 2 seeds, and choosing the learning rate that performed best on aggregate. The same learning rate is used in all experiments.

For TD3, we use the “target policy smoothing” recommended in the original paper of $\mathcal{N}(0, 0.2)$ clipped to lie in range $[-0.5, 0.5]$ along each dimension.

<table>
<thead>
<tr>
<th>Learning Rate</th>
<th>$1e^{-4}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hidden Layer sizes</td>
<td>[256, 256]</td>
</tr>
<tr>
<td>Learning Rate</td>
<td>$1e^{-4}$</td>
</tr>
<tr>
<td>Optimizer</td>
<td>Adam</td>
</tr>
<tr>
<td>Target Update Rate</td>
<td>$5e^{-3}$</td>
</tr>
<tr>
<td>Batch Size</td>
<td>512</td>
</tr>
<tr>
<td>Iterations per timestep</td>
<td>1</td>
</tr>
<tr>
<td>Discount factor</td>
<td>0.99</td>
</tr>
<tr>
<td>Exploration Policy</td>
<td>$\mathcal{N}(0, 0.05)$</td>
</tr>
<tr>
<td>Samples used for action-selection</td>
<td>1000</td>
</tr>
<tr>
<td>Samples used for bootstrapping</td>
<td>1000</td>
</tr>
<tr>
<td>Replay Buffer Size</td>
<td>500000</td>
</tr>
<tr>
<td>Activation Function</td>
<td>Tanh</td>
</tr>
</tbody>
</table>

Table 2: Hyperparameters for Q-Functional Benchmark Experiments

<table>
<thead>
<tr>
<th>Learning Rate</th>
<th>$1e^{-4}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hidden Layer sizes (Policy)</td>
<td>[256, 256]</td>
</tr>
<tr>
<td>Learning Rate</td>
<td>$1e^{-3}$</td>
</tr>
<tr>
<td>Optimizer</td>
<td>Adam</td>
</tr>
<tr>
<td>Target Update Rate</td>
<td>$5e^{-3}$</td>
</tr>
<tr>
<td>Batch Size</td>
<td>512</td>
</tr>
<tr>
<td>Iterations per timestep</td>
<td>1</td>
</tr>
<tr>
<td>Discount factor</td>
<td>0.99</td>
</tr>
<tr>
<td>Exploration Policy</td>
<td>$\mathcal{N}(0, 0.1)$</td>
</tr>
<tr>
<td>Replay Buffer Size</td>
<td>500000</td>
</tr>
<tr>
<td>Activation Function</td>
<td>ReLU</td>
</tr>
</tbody>
</table>

Table 3: Hyperparameters for Policy Gradient Methods
Proofs

We prove using methods from the Calculus of Variations (Gelfand, Silverman et al. 2000) that in 1D bounded action spaces, the Boltzmann distribution (softmax) over Q-values is the policy that maximizes the entropy-regularized value. To the best of our knowledge this is a new proof method for this known fact.

Let \( Q(s, a) : S \times A \to \mathbb{R} \) be the Q-value at every action for the given state, with \( 0 \leq a \leq 1 \). Let \( \pi(s, a) = \Pr(a \mid s) \) be the probability of a given action. We seek a policy \( \pi \) that maximizes entropy-regularized value,

\[
V_\pi^{\text{ENT}}(s) = \int_0^1 \pi(s, a)(Q(s, a) - \tau \log \pi(s, a))da.
\]

We define \( \Pi(s, a) = \int_0^a \pi(s, x)dx \) to be the CDF of \( \pi \), with \( \Pi(0) = 0 \) and \( \Pi(1) = 1 \), noting that \( \frac{d}{da}\Pi(s, a) = \pi(a) \).

We now make use of the Euler-Lagrange equation from the Calculus of Variations:

**Statement of the Euler-Lagrange Theorem:** The action functional \( S \) is defined as

\[
S[q] = \int_a^b F(x, q(x), q'(x))dx.
\]

Let \( q(x) \) be a function satisfying boundary conditions \( q(a) = b_1 \) and \( q(y) = b_2 \). Then \( q(x) \) is a stationary point of \( S \) iff:

\[
\frac{\partial F}{\partial q}(x, q(x), q'(x)) - \frac{d}{dx}\frac{\partial F}{\partial q'} = 0.
\]

We now write down the minimized function, \( V_\pi^{\text{ENT}} \), in terms of the CDF \( \Pi \):

\[
S[\Pi] = \int_0^1 F(a, \Pi(s, a), \Pi'(s, a))da = \int_0^1 \Pi'(s, a)(Q(s, a) - \tau \log \Pi'(s, a))da
\]

We note that here, \( F \) does not explicitly depend on \( \Pi \) (only \( \Pi' \)). Therefore,

\[
\frac{\partial F}{\partial \Pi} = 0.
\]

Additionally,

\[
\frac{\partial F}{\partial \Pi'} = Q(s, a) - \tau \log \Pi'(s, a) - \tau.
\]

Evaluating the Euler-Lagrange equation, we are left with

\[
0 = \frac{\partial F}{\partial q}(x, q(x), q'(x)) - \frac{d}{dx}\frac{\partial F}{\partial q'} = \frac{d}{dx}\left[ Q(s, a) - \tau \log \Pi'(s, a) - \tau \right]
\]

\[
\Rightarrow C = Q(s, a) - \tau \log \Pi'(s, a) - \tau
\]

\[
\Rightarrow \log \pi(s, a) = \frac{Q(s, a)}{\tau} - C/\tau - 1
\]

\[
\Rightarrow \pi(s, a) = D\exp\{Q(s, a)/\tau\}
\]

where \( C \) is an integration constant and \( D = \exp\{-C/\tau - 1\} \). Therefore, optimal policy \( \pi^* \) is proportional to the Boltzmann distribution with temperature \( \tau \), with a normalizing constant equal to \( \text{NORM} = \int_0^1 \exp\{Q(s, a)/\tau\}da \).

Intuitively, the above proof goes as follows: since \( \pi \) is constrained to have total probability 1, changes to \( \pi \) can be made by moving probability from one action to another. A given \( \pi \) is a stationary point only when gains cannot be made by moving probability from one action to another, or in other words, when the regularized value of all actions are equal. This condition only holds when \( \pi(s, a) \propto \exp\{Q(s, a)\} \).

We can now additionally simplify \( V_\pi^{\text{ENT}} \):
\[ V_{\text{ENT}}^\pi(s) = \int_0^1 \pi(s, a)(Q(s, a) - \tau \log \pi(s, a))da \]

\[ = \int_0^1 \frac{1}{\text{NORM}} \exp\{Q(s, a)/\tau\}\{Q(s, a) - \tau(Q(s, a)/\tau) + \log \text{NORM}\}da \]

\[ = \int_0^1 \frac{1}{\text{NORM}} \exp\{Q(s, a)/\tau\}\{\log \text{NORM}\}da \]

\[ = (\log \text{NORM}) \int_0^1 \frac{1}{\text{NORM}} \exp\{Q(s, a)/\tau\}da \]

\[ = (\log \text{NORM}) \int_0^1 \frac{1}{\text{NORM}} \exp\{Q(s, a)/\tau\}da \]

\[ = (\log \text{NORM}) \times 1 \]

\[ = \log \int_0^1 \exp\{Q(s, a)/\tau\}da \]

For an alternative proof of above claims, please consult Appendix 1.A of (Haarnoja et al. 2018).
Policy Quality Graphs

We include plots similar to that of Figure 2 in the main paper, for all benchmark tasks. These plots depict how frequently (x axis) each policy-gradient method selects actions better than some fraction of random actions (y axis). In general, TD3 outperforms DDPG at the task of choosing high-value actions, including near-optimal action selection on the Reacher task. This is perhaps due to its smoother Q-value function or more stable training. However, on a large number of tasks both methods frequently perform poorly at action selection. Figure 1 details action-selection performance for DDPG, and Figure 2 details action-selection performance for TD3.

Figure 1: Action-selection quality by DDPG, on all tasks.
Figure 2: Action-selection quality by TD3, on all tasks.
Supplementary Experiments

Precomputing Basis Representations

In Figure 3 we compare performance between using precomputed action representations, and re-computing action representations for each sample. We find generally similar performance between both approaches, although the Q-functional DDPG-analogue performs significantly worse on one domain with precomputed representations.

Figure 3: Comparison of performance for precomputing-vs-recomputing basis representations. The shaded region represents the standard error over 8 runs. One iteration corresponds to 10,000 environment steps.

We do however find a significant difference in speed, motivating our use of precomputed representations in our benchmarking experiments.

Figure 4: Comparison of wallclock runtime for precomputing-vs-recomputing basis representations. The shaded region represents the standard error over 8 runs. One iteration corresponds to 10,000 environment steps.
Rank Sweep

In Figure 5, we compare performance using different ranks on 3 tasks, using the Legendre basis, and both the DDPG and TD3 Q-functional analogues. We find that rank 1 is frequently not enough to solve these tasks, while higher ranks generally perform similarly.

Figure 5: Performance of Q-functionals on 3 tasks from the OpenAI continuous control suite using different ranks. The shaded region represents the standard error over 8 runs. One iteration corresponds to 10,000 environment steps.
Basis Sweep

In Figure 6, we compare performance using different basis functions on 3 tasks, for both the DDPG and the TD3 analogue Q-functionals. On these tasks, the Polynomial basis performs marginally better than the Legendre basis, and both of these frequently outperform the Fourier basis.

Figure 6: Performance of Q-functionals on 3 tasks from the OpenAI continuous control suite using different basis functions. The shaded region represents the standard error over 8 runs. One iteration corresponds to 10,000 environment steps.

References