Learning Sparse Representations in Reinforcement Learning with Sparse Coding

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Abstract

A variety of representation learning approaches have been investigated for reinforcement learning; much less attention, however, has been given to investigating the utility of sparse coding. Outside of reinforcement learning, sparse coding representations have been widely used, with non-convex objectives that result in discriminative representations. In this work, we develop a supervised sparse coding objective for policy evaluation. Despite the non-convexity of this objective, we prove that all local minima are global minima, making the approach amenable to simple optimization strategies. We empirically show that it is key to use a supervised objective, rather than the more straightforward unsupervised sparse coding approach. We compare the learned representations to a canonical fixed sparse representation, called tile-coding, demonstrating that the sparse coding representation outperforms a wide variety of tilecoding representations.

1 Introduction

For tasks with large state or action spaces, where tabular representations are not feasible, reinforcement learning algorithms typically rely on function approximation. Whether they are learning the value function, policy or models, the success of function approximation techniques hinges on the quality of the representation. Typically, representations are hand-crafted, with some common representations including tile-coding, radial basis functions, polynomial basis functions and Fourier basis functions [Sutton, 1996; Konidaris *et al.*, 2011]. Automating feature discovery, however, alleviates this burden and has the potential to significantly improve learning.

Representation learning techniques in reinforcement learning have typically drawn on the large literature in unsupervised and supervised learning. Common approaches include feature selection, including ℓ_1 regularization on the value function parameters [Loth *et al.*, 2007; Kolter and Ng, 2009; Nguyen *et al.*, 2013] and matching pursuit [Parr *et al.*, 2008; Painter-Wakefield and Parr, 2012]; basis-function adaptation approaches [Menache *et al.*, 2005; Whiteson *et al.*, 2007];

instance-based approaches, such as locally weighted regression [Atkeson and Morimoto, 2003], sparse distributed memories [Ratitch and Precup, 2004], proto-value functions [Mahadevan and Maggioni, 2007] and manifold learning techniques [Mahadevan, 2009]; and neural network approaches, including more standard feedforward neural networks [Coulom, 2002; Riedmiller, 2005; Mnih *et al.*, 2015] as well as random representations [Sutton and Whitehead, 1993], linear threshold unit search [Sutton and Barto, 2013], and evolutionary algorithms like NEAT [Stanley and Miikkulainen, 2002].

Surprisingly, however, there has been little investigation into using sparse coding for reinforcement learning. Sparse coding approaches have been developed to learn MDP models for transfer learning [Ammar *et al.*, 2012]; outside this work, however, little has been explored. Nonetheless, such sparse coding representations have several advantages, including that they naturally enable local models, are computationally efficient to use, are much simpler to train than more complicated models such as neural networks and are biologically motivated by the observed representation in the mammalian cortex [Olshausen and Field, 1997].

In this work, we develop a principled sparse coding objective for policy evaluation. In particular, we formulate a joint optimization over the basis and the value function parameters, to provide a supervised sparse coding objective where the basis is informed by its utility for prediction. We highlight the importance of using the Bellman error or mean-squared return error for this objective, and discuss how the projected Bellman error is not suitable. We then show that, despite being a nonconvex objective, all local minima are global minima, under minimal conditions. We avoid the need for careful initialization strategies needed for previous optimality results for sparse coding [Agarwal et al., 2014; Arora et al., 2015], using recent results for more general dictionary learning settings [Haeffele and Vidal, 2015; Le and White, 2017], particularly by extending beyond smooth regularizers using Γ -convergence. Using this insight, we provide a simple alternating proximal gradient algorithm and demonstrate the utility of learning supervised sparse coding representations versus unsupervised sparse coding and a variety of tile-coding representations.

2 Background

In reinforcement learning, an agent interacts with its environment, receiving observations and selecting actions to maxi-

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mize a scalar reward signal provided by the environment. This interaction is usually modeled by a Markov decision process (MDP). An MDP consists of (S, A, P, R) where S is the set of states; A is a finite set of actions; $P : S \times A \times S \rightarrow [0, 1]$, the transition function, which describes the probability of reaching a state s' from a given state and action (s, a); and finally the reward function $R : S \times A \times S \rightarrow \mathbb{R}$, which returns a scalar value for transitioning from state-action (s, a)to state s'. The state of the environment is said to be *Markov* if $Pr(s_{t+1}|s_t, a_t) = Pr(s_{t+1}|s_t, a_t, \ldots, s_0, a_0)$.

One important goal in reinforcement learning is policy evaluation: learning the *value function* for a policy. A value function $\mathbf{V}^{\pi} : S \to \mathbb{R}$ approximates the expected return. The return G_t from a state s_t is the total discounted future reward, discounted by $\gamma \in [0, 1)$, for following policy $\pi : S \times A \to [0, 1]$

$$G_t = \sum_{i=0}^{\infty} \gamma^i R_{t+1+i} = R_{t+1} + \gamma G_{t+1}$$

where $\mathbf{V}^{\pi}(s_t)$ is the expectation of this return from state s_t . This value function can also be thought of as a vector of values $\mathbf{V}^{\pi} \in \mathbb{R}^{|S|}$ satisfying the Bellman equation

$$\mathbf{V}^{\pi} = \mathbf{r}^{\pi} + \gamma \mathbf{P}^{\pi} \mathbf{V}^{\pi} \tag{1}$$

where
$$\mathbf{P}^{\pi}(s, s') = \sum_{a \in \mathcal{A}} \pi(s, a) P(s, a, s')$$

 $\mathbf{r}^{\pi}(s) = \sum_{a \in \mathcal{A}} \pi(s, a) \sum_{s' \in \mathcal{S}} P(s, a, s') R(s, a, s')$

Given the reward function and transition probabilities, the solution can be analytically obtained: $\mathbf{V}^{\pi} = (\mathbf{I} - \gamma \mathbf{P}^{\pi})^{-1} \mathbf{r}^{\pi}$.

In practice, however, we likely have a prohibitively large state space. The typical strategy in this setting is to use function approximation to learn $\mathbf{V}^{\pi}(s)$ from a trajectory of samples: a sequence of states, actions, and rewards s_0 , a_0 , r_0 , s_1 , a_1 , r_1 , s_2 , r_2 , a_2 ..., where s_0 is drawn from the start-state distribution, $s_{t+1} \sim P(\cdot|s_t, a_t)$ and $a_t \sim \pi(\cdot|s_t)$. Commonly, a linear function is assumed, $\mathbf{V}^{\pi}(s) \approx \phi(s)^{\top} \mathbf{w}$ for $\mathbf{w} \in \mathbb{R}^k$ a parameter vector and $\phi : S \to \mathbb{R}^k$ a feature function describing states. With this approximation, however, typically we can no longer satisfy the Bellman equation in (1), because there may not exist a \mathbf{w} such that $\Phi \mathbf{w}$ equals $\mathbf{r}^{\pi} + \gamma \mathbf{P}^{\pi} \Phi \mathbf{w}$ for $\Phi \in \mathbb{R}^{|S| \times k}$. Instead, we focus on minimizing the error to the true value function.

Reinforcement learning algorithms, such as temporal difference learning and residual gradient, therefore focus on finding an approximate solution to the Bellman equation, despite this representation issue. The quality of the representation is critical to accurately approximating \mathbf{V}^{π} with $\boldsymbol{\Phi}\mathbf{w}$, but also balancing compactness of the representation and speed of learning. Sparse coding, and sparse representations, have proven successful in machine learning and in reinforcement learning, particularly as fixed bases, such as tile coding, radial basis functions and other kernel representations. A natural goal, therefore, and the one we explore in this work, is to investigate learning these sparse representations automatically.

3 Sparse Coding for Reinforcement Learning

In this section, we formalize sparse coding for reinforcement learning as a joint optimization over the value function parameters and the representation. We introduce the true objective over all states, and then move to the sampled objective for the algorithm in the next section.

We begin by formalizing the representation learning component. Many unsupervised representation learning approaches consist of factorizing input observations¹ $\mathbf{X} \in \mathbb{R}^{|S| \times d}$ into a basis dictionary $\mathbf{B} \in \mathbb{R}^{k \times d}$ and new representation $\mathbf{\Phi} \in \mathbb{R}^{|S| \times k}$. The rows of **B** form a set of bases, with columns in $\mathbf{\Phi}$ weighting amongst those bases for each observation (column) in **X**. Though simple, this approach encompasses a broad range of models, including PCA, CCA, ISOMAP, locally linear embeddings and sparse coding [Singh and Gordon, 2008; Le and White, 2017]. The (unsupervised) sparse coding objective is [Aharon *et al.*, 2006]

$$\min_{\mathbf{\Phi}\in\mathbb{R}^{|\mathcal{S}|\times k},\mathbf{B}\in\mathbb{R}^{k\times d}} \|\mathbf{\Phi}\mathbf{B}-\mathbf{X}\|_{D}^{2} + \beta_{B}\|\mathbf{B}\|_{F}^{2} + \beta_{\phi}\|\mathbf{\Phi}\|_{D,1}$$

where $\|\mathbf{Y}\|_{F}^{2} = \sum_{ij} \mathbf{Y}_{ij}^{2}$ is the squared Frobenius norm; $\mathbf{B} \in \mathbb{R}^{k \times d}$ is a learned basis dictionary; $\beta_{B}, \beta_{\phi} > 0$ determine the magnitudes of the regularizers; $\mathbf{D} \in [0, 1]^{|\mathcal{S}| \times |\mathcal{S}|}$ is a diagonal matrix giving a distribution over states, corresponding to the stationary distribution of the policy $d^{\pi} : \mathcal{S} \to [0, 1]$; and $||\mathbf{z}||_{D}^{2} = \mathbf{z}^{\top} \mathbf{D} \mathbf{z}$ is a weighted norm. The reconstruction error

$$\|\mathbf{\Phi}\mathbf{B} - \mathbf{X}\|_D^2 = \sum_{s \in S} d^{\pi}(s) \|\mathbf{\Phi}(s,:)\mathbf{B} - \mathbf{X}(s,:)\|_2^2$$

is weighted by the stationary distribution d^{π} because states are observed with frequency indicated by d^{π} . The weighted ℓ_1

$$\|\mathbf{\Phi}\|_{D,1} = \sum_{s \in \mathcal{S}} d^{\pi}(s) \sum_{j=1}^{k} |\mathbf{\Phi}(s,j)|$$

promotes sparsity on the entries of Φ , preferring entries in Φ to be entirely pushed to zero rather than spreading magnitude across all of Φ . The Frobenius norm regularizer on **B** ensures that **B** does not become too large. Without this regularizer, all magnitude can be shifted to **B**, producing the same Φ **B**, but pushing $\|\Phi\|_{D,1}$ to zero and nullifying the utility of its regularizer. Optimizing this sparse coding objective would select a sparse representation ϕ for each observation x such that ϕ **B** approximately reconstructs x.

Further, however, we would like to learn a new representation that is also optimized towards approximating the value function. Towards this aim, we need to jointly learn Φ and w, where Φw provides the approximate value function. In this way, the optimization must balance between accurately recreating X and approximating the value function Φw . For this, we must choose an objective for learning w.

We consider two types of objectives: fixed-point objectives and squared-error objectives. Two common fixed-point objectives are the mean-squared Bellman error (MSBE), also called the Bellman residual [Baird, 1995]

$$\|\mathbf{\Phi}\mathbf{w} - (\mathbf{r}^{\pi} + \gamma \mathbf{P}^{\pi} \mathbf{\Phi}\mathbf{w})\|_{D}^{2}$$

and mean-squared projected BE (MSPBE) [Sutton et al., 2009]

$$\mathbf{\Phi}\mathbf{w} - \Pi(\mathbf{r}^{\pi} + \gamma \mathbf{P}^{\pi} \mathbf{\Phi} \mathbf{w}) \|_{D}^{2}$$

¹This variable \mathbf{X} can also be a base set of features, on which the agent can improve or which the agent can sparsify.

where $\mathbf{D} \in [0,1]^{|\mathcal{S}| \times |\mathcal{S}|}$ is a diagonal matrix giving a distribution over states, corresponding to the stationary distribution of the policy; $||\mathbf{z}||_D^2 = \mathbf{z}^\top \mathbf{D} \mathbf{z}$ is a weighted norm; and the projection matrix for linear value functions is $\Pi =$ $\mathbf{\Phi}(\mathbf{\Phi}^{\top}\mathbf{D}\mathbf{\Phi})^{-1}\mathbf{\Phi}^{\top}\mathbf{D}$. The family of TD algorithms converge to the minimum of the MSPBE, whereas residual gradient algorithms typically use the MSBE (see [Sun and Bagnell, 2015] for an overview). Both have useful properties [Scherrer, 2010], though arguably the MSPBE is more widely used.

There are also two alternative squared-error objectives, that do not correspond to fixed-point equations: the mean-squared return error (MSRE) and the Bellman error (BE). For a trajectory of samples $\{(\mathbf{x}_i, r_{i+1}, \mathbf{x}_{i+1})\}_{i=0}^{t-1}$, BE is defined as

$$\sum_{i=0}^{t-1} \|r_{i+1} + \gamma \boldsymbol{\phi}_{i+1}^{\top} \mathbf{w} - \boldsymbol{\phi}_{i}^{\top} \mathbf{w}\|_{2}^{2}$$

and the MSRE as $\sum_{i=0}^{t-1} \|g_{i+1} - \boldsymbol{\phi}_i^{ op} \mathbf{w}\|_2^2$

where $g_{i+1} = \sum_{j=i}^{t-1} \gamma^{j-i} r_{j+1}$ is a sample return. In expectation, these objectives are, respectively

$$\sum_{s \in \mathcal{S}} d^{\pi}(s) \mathbb{E} \left[\left(r(S_t, A_t, S_{t+1}) + \gamma \boldsymbol{\phi}(S_{t+1})^{\top} \mathbf{w} - \boldsymbol{\phi}(S_t)^{\top} \mathbf{w} \right)^2 | S_t = s \right]$$
$$\sum_{s \in \mathcal{S}} d^{\pi}(s) \mathbb{E} \left[\left(\sum_{i=0}^{\infty} \gamma^i r(S_{t+i}, A_{t+i}, S_{t+1+i}) - \boldsymbol{\phi}(s)^{\top} \mathbf{w} \right)^2 | S_t = s \right]$$

where the expectation is w.r.t. the transition probabilities and taking actions according to policy π .

These differ from the fixed-point objectives because of the placement of the expectation. To see why, consider the MSBE and BE. The expected value of the BE is the expected squared error between the prediction from this state and the reward plus the value from a possible next state. The MSBE, on the other hand, is the squared error between the prediction from this state and the expected reward plus the expected value for the next state. Though the MSPBE and MSBE constitute the most common objectives chosen for reinforcement learning, these squared-error objectives have also been shown to be useful particularly for learning online [Sun and Bagnell, 2015].

For sparse coding, however, the MSPBE is not a suitable choice-compared to the MSBE. BE and MSRE-for two reasons. First, the MSBE, BE and MSRE are all convex in Φ , whereas the MSPBE is not. Second, because of the projection onto the space spanned by the features, the MSPBE can be solved with zero error for any features Φ . Therefore, because it does not inform the choice of Φ , the MSPBE produces a two stage approach², where features are learned in a completely unsupervised way and prediction performance does not influence Φ .

The final objective for loss $L(\Phi, w)$ set to either MSBE, BE or MSRE is

$$\min_{\mathbf{w}\in\mathbb{R}^{k},\mathbf{\Phi}\in\mathbb{R}^{|\mathcal{S}|\times k},\mathbf{B}\in\mathbb{R}^{k\times d}} L(\mathbf{\Phi},\mathbf{w}) + \|\mathbf{\Phi}\mathbf{B}-\mathbf{X}\|_{D}^{2}$$
(2)

 $+ \beta_w \|\mathbf{w}\|_2^2 + \beta_B \|\mathbf{B}\|_F^2 + \beta_\phi \|\mathbf{\Phi}\|_{D,1}$

Algorithm for Sparse Coding 4

We now derive the algorithm for sparse coding for policy evaluation: SCoPE. We generically consider either the BE or MSRE. For a trajectory of samples $\{(\mathbf{x}_i, r_{i+1}, \mathbf{x}_{i+1})\}_{i=0}^{t-1}$, the objective is

$$\min_{\mathbf{w}\in\mathbb{R}^{k}, \boldsymbol{\Phi}\in\mathbb{R}^{t+1\times k}, \mathbf{B}\in\mathbb{R}^{k\times d}} \frac{1}{t} \sum_{i=0}^{t-1} (y_{i} + \bar{\gamma}\boldsymbol{\phi}_{i+1}^{\top}\mathbf{w} - \boldsymbol{\phi}_{i}^{\top}\mathbf{w})^{2}$$
(3)

$$+ \frac{1}{t} \sum_{i=0}^{t} \|\phi_i \mathbf{B} - \mathbf{x}_i\|_2^2 + \beta_B \|\mathbf{B}\|_F^2 + \beta_w \|\mathbf{w}\|_2^2 + \frac{\beta_\phi}{t} \sum_{i=0}^{t} \|\phi_i\|_1^p$$

for BE, $y_i = r_{i+1}$ and $\bar{\gamma} = \gamma$ and for MSRE, $y_i =$ $\sum_{j=i}^{t} \gamma^{j-i} r_{j+1}$ and $\bar{\gamma} = 0$. We consider two possible powers for the ℓ_1 norm p = 1 or 2, where the theory relies on using p = 2, but in practice we find they perform equivalently and p = 1 provides a slightly simpler optimization. The loss is averaged by t, to obtain a sample average, which in the limit converges to the expected value under d^{π} . This averaged loss is also more scale-invariant-in terms of the numbers of samples-to the choice of regularization parameters.

SCoPE consists of alternating amongst these three variables, \mathbf{B}, \mathbf{w} and $\boldsymbol{\Phi}$, with a proximal gradient update for the nondifferentiable ℓ_1 norm. The loss in terms of **B** and **w** is differentiable; to solve for \mathbf{B} (or \mathbf{w}) with the other variables fixed, we can simply used gradient descent. To solve for Φ with the B and w fixed, however, we cannot use a standard gradient descent update because the ℓ_1 regularizer is nondifferentiable. The proximal update consists of stepping in the direction of the gradient for the smooth component of the objective—which is differentiable—and then projecting back to a sparse solution using the proximal operator: a soft thresholding operator. The convergence of this alternating minimization follows from results on block coordinate descent for non-smooth regularizers [Xu and Yin, 2013].

To apply the standard proximal operator for the ℓ_1 regularizer, we need to compute an upper bound on the Lipschitz constant for this objective. The upper bound is $2(1+\bar{\gamma}^2) \|\mathbf{w}\|_2^2 + 2\|\mathbf{B}\|_{sp}^2$, computed by finding the maximum

²This problem seems to have been overlooked in two approaches for basis adaptation based on the MSPBE: adaptive bases algorithm for the projected Bellman error (ABPBE) [Di Castro and Mannor, 2010, Algorithm 9] and mirror descent $Q(\lambda)$ with basis adaptation [Mahadevan et al., 2013]. For example, for ABPBE, it is not immediately obvious this would be a problem, because a stochastic approximation approach is taken. However, if written as a minimization over the basis parameters and the weights, one would obtain a

minimum error solution (i.e., error zero) immediately for any basis parameters. The basis parameters are considered to change on a slow timescale, and the weights on a fast timescale, which is a reflection of this type of separate minimization. Menache et al. [2005] avoided this problem by explicitly using a two-stage approach, using MSPBE approaches for learning the parameters and using other score function, such as the squared Bellman error, to update the bases. This basis learning approach, however, is unsupervised.

Representation learning strategies for the MSPBE have been developed, by using local projections [Yu and Bertsekas, 2009; Bhatnagar et al., 2009]. These strategies, however, do not incorporate sparse coding.

singular value of the Hessian of the objective w.r.t. ϕ_i for each *i*. We will provide additional details for this calculation, and implementation details, in a supplement.

4.1 Local Minima Are Global Minima

In this section, we show that despite nonconvexity, the objective for SCoPE has the nice property that all local minima are in fact global minima. Consequently, though there may be many different local minima, they are in fact equivalent in terms of the objective. This result justifies a simple alternating minimization scheme, where convergence to local minima ensures an optimal solution is obtained.

We need the following technical assumption. It is guaranteed to be true for a sufficiently large $k \le t$ (see Haeffele and Vidal [2015]; Le and White [2017]).

Assumption 1 For the given $k \ge d$, the following function is convex in $\mathbf{Z} \in \mathbb{R}^{t+1 \times d+1}$

$$\min_{\boldsymbol{\Phi}, \mathbf{B}, \mathbf{w}, \mathbf{Z} = \boldsymbol{\Phi}[\mathbf{B} \mathbf{w}]} \beta_B \|\mathbf{B}\|_F^2 + \beta_w \|\mathbf{w}\|_2^2 + \frac{\beta_\phi}{t} \sum_{i=0} \|\phi_i\|_1^2$$

Theorem 1 (Landscape of the SCoPE objective). For the objective in equation (3) with p = 2,

1. under Assumption 1, all full-rank local minima are global minima; and

2. *if a local minimum* $(\mathbf{\Phi}, \mathbf{B}, \mathbf{w})$ *has* $\mathbf{\Phi}_{:i} = \mathbf{0}$ *(i.e., a zero column) and* $\mathbf{w}_{i} = 0$, $\mathbf{B}_{i:} = \mathbf{0}$ *(i.e., a zero row) for some* $1 \leq i \leq k$, then it is a global minimum.

Proof. For the first statement, we construct a limit of twicedifferentiable functions f_n that Γ -converge to the SCoPE objective f. With this, we can then show that all minimizers of the sequence converge to minimizers of f, and vice-versa [Braides, 2013]. Because all local minimizers of the twicedifferentiable functions f_n are global minimizers from [Le and White, 2017, Theorem 10], we can conclude that all corresponding minimizers of f are global minimizers.

We use the pseudo-Huber loss [Fountoulakis and Gondzio, 2013], which is twice-differentiable approximation to the absolute value: $|x|_{\mu} = \sqrt{\mu^2 + x^2} - \mu$. Let $\boldsymbol{\theta} = (\boldsymbol{\Phi}, \mathbf{B}, \mathbf{w})$. The sequence of functions f_n are defined with $\mu_n = 1/n$, as

$$f_n(\boldsymbol{\theta}) = L(\boldsymbol{\theta}) + \frac{\beta_\phi}{t} \sum_{ij} \sqrt{\mu_n^2 + \Phi_{ij}^2} - \mu_n$$

where $L(\theta)$ equals the equation in (3), but without the ℓ_1^2 regularizer on Φ .

Part 1: All local minima of f_n for all n are global minima. To show this, we show each f_n satisfies the conditions of [Le and White, 2017, Theorem 10 and Proposition 11].

Part 1.1 We can rewrite the loss in terms of $\Phi[\mathbf{B} \mathbf{w}]$

$$\frac{1}{t} \sum_{i=0}^{t-1} \|y_i + \bar{\gamma} \boldsymbol{\phi}_{i+1}^{\top} \mathbf{w} - \boldsymbol{\phi}_i^{\top} \mathbf{w}\|_2^2 + \frac{1}{t} \sum_{i=0}^{t} \|\boldsymbol{\phi}_i \mathbf{B} - \mathbf{x}_i\|_2^2$$
$$= \frac{1}{t} \|\mathbf{X} - \boldsymbol{\Phi} \mathbf{B}\|_2^2 + \frac{1}{t} \|\mathbf{y} - (\mathbf{I}_{0:t-1} - \bar{\gamma} \mathbf{I}_{1:t}) \boldsymbol{\Phi} \mathbf{w}\|_2^2$$
(4)

where $\mathbf{I}_{1:t} \in \mathbb{R}^{t \times t+1}$ a diagonal matrix of all ones with the first diagonal entry set to zero, and $\mathbf{I}_{0:t-1}$ with the last diagonal entry set to zero. This loss is convex in the joint variable $\Phi[\mathbf{B} \mathbf{w}]$ because equation (4) is the composition of a convex

function (squared norm) and an affine function (multiplication by $\bar{\gamma} \mathbf{I}_{1:t} - \mathbf{I}_{0:t-1}$ and addition of \mathbf{y}).

Part 1.2 The regularizer on $[\mathbf{B} \mathbf{w}]$ must be a weighted Frobenius norm, with weightings on each column; here, we have weighting using regularization parameters β_B for the first *d* columns (corresponding to **B**) and regularization parameter β_w for the last column (corresponding to **w**).

Part 1.3 The inner dimension k > d, which is true by assumption and the common setting for sparse coding.

Part 1.4 The pseudo-Huber loss, on the columns of Φ , is convex, centered and twice-differentiable.

Part 2: The sequence f_n converges uniformly to f. To see why, recall the definition of uniform convergence. A sequence of functions $\{f_n\}$ is uniformly convergent with limit f if for every $\epsilon > 0$, there exists $N \in \mathbb{N}$ such that for all $\theta \in \Theta$ all $n \ge N$, $|f_n(\theta) - f(\theta)| < \epsilon$. Further recall that for any complete metric space, if f_n is uniformly Cauchy, then it is uniformly convergent. The sequence is uniformly Cauchy if for all $n, m \ge N$, $|f_n(\theta) - f_m(\theta)| < \epsilon$. Take any $\epsilon > 0$ and let $N = \lceil \frac{4k(t+1)\beta_{\phi}}{t\epsilon} \rceil$. Then

$$\begin{aligned} &|f_n(\boldsymbol{\theta}) - f_m(\boldsymbol{\theta})| \\ &= \frac{\beta_\phi}{t} \left\| \left(\sum_{ij} \sqrt{\mu_n^2 + \boldsymbol{\Phi}_{ij}^2} - \mu_n \right) - \left(\sum_{ij} \sqrt{\mu_m^2 + \boldsymbol{\Phi}_{ij}^2} - \mu_m \right) \right| \\ &\leq \frac{\beta_\phi}{t} \sum_{ij} \left(\left| \sqrt{\mu_n^2 + \boldsymbol{\Phi}_{ij}^2} - \sqrt{\mu_m^2 + \boldsymbol{\Phi}_{ij}^2} \right| + |\mu_n - \mu_m| \right) \end{aligned}$$

The upper bound of the first component is maximized when $\Phi_{ij} = 0$, and so we get

$$\begin{aligned} |f_n(\boldsymbol{\theta}) - f_m(\boldsymbol{\theta})| &\leq \frac{2k(t+1)\beta_{\phi}}{t} |\mu_n - \mu_m| \leq \frac{2k(t+1)\beta_{\phi}}{t} \left| \frac{1}{n} - \frac{1}{m} \right| \\ &\leq \frac{4k(t+1)\beta_{\phi}}{tN} \leq \epsilon. \end{aligned}$$

Part 3: Asymptotic equivalence of minimizers of f_n and f. Because f is continuous, and so lower semi-continuous, and f_n uniformly converges to f, we know that $f_n \Gamma$ -converges to $f: f_n \xrightarrow{\Gamma} f$ Braides [2013].

By the fundamental theorem of Γ -convergence, if the $\{f_n\}$ is an equi-coercive family of functions, then the minimizers of f_n converge to minimizers of f. A sequence of functions $\{f_n\}$ is equi-coercive iff there exists a lower semi-continuous coercive function $\psi : \Theta \to \mathbb{R} \cup \{-\infty, \infty\}$ such that $f_n \ge \psi$ on Θ for every $n \in \mathbb{N}$ [Dal Maso, 2012, Proposition 7.7]. A function is coercive if $\psi(\theta) \to \infty$ as $\|\theta\| \to \infty$. For $\psi(\theta) =$ $L(\theta)$, it is clear that ψ is coercive, as well as lower semicontinuous (since it is continuous). Further, $f_n(\theta) \ge L(\theta) =$ $\psi(\theta)$, because the regularizer on Φ is non-negative. Therefore, the family $\{f_n\}$ is equi-coercive, and so the minimizers of f_n converge to minimizers of f.

For the other direction, if a local minimum θ of f is an isolated local minimum, then there exists a sequence $\theta_n \to \theta$ with θ_n a local minimizer of f_n for μ_n sufficiently small [Braides, 2013, Theorem 5.1]. Because we have Frobenius norm regularizers on **B**, **w**, which are strongly convex, the objective is strictly convex with respect to **B**, **w**. Further, because **X** is full rank, $\|\Phi \mathbf{B} - \mathbf{X}\|_F^2$ is a strictly convex function



Figure 1: Learning curves for SCoPE versus a variety of tile coding representations in three domains. The graphs depict early learning; the numbers in brackets correspond to final error, after 5000 samples. Because errors are sampled every 50 samples, and because MSRE is used for optimization, the lines have an irregular pattern. The differences are nonetheless statistically significant, with an average over 50 runs, and so the small standard error bars are omitted. SCoPE outperforms the best of the TC representations in Mountain Car and Acrobot using a more compact sparse representation; in Puddle World, it performs more poorly, which we discuss further in the text. The larger TC representations likely perform poorly due to hashing.

with respect to Φ . Therefore, locally the objective is strictly convex with respect to θ . We therefore know that local minima of f are isolated, and so there exists an N such that for all n > N, θ_n are local minimizers of f_n . Since these local minimizers are global minimizers, and they converge to θ , this means θ is a global minimum of f.

For the second statement, we use [Haeffele and Vidal, 2015, Theorem 15]. Because we already showed above that our loss can be cast as factorization, it is clear our loss and regularizers are positively homogenous, of order 2. A minimum is guaranteed to exist for our objective, because the loss function is continuous, bounded below (by zero) and goes to infinity as the parameters go to $\pm \infty$.

5 Experimental Results

We aim to address the question: can we learn useful representations using SCoPE? We therefore tackle the setting where the representation is first learned, and then used, to avoid conflating incremental estimation and the utility of the representation. We particularly aim to evaluate estimation accuracy, as well as qualitatively understanding the types of sparse representations learned by SCoPE.

Domains. We conducted experiments in three benchmark RL domains - Mountain Car, Puddle World and Acrobot [Sutton, 1996]. All domains are episodic, with discount set to 1 until termination. The data in Mountain Car is generated using the standard energy-pumping policy policy with 10% randomness. The data in Puddle World is generated by a policy that chooses to go North with 50% probability, and East with 50% probability on each step, with the starting position in the lower-left corner of the grid, and the goal in the top-right corner. The data in Acrobot is generated by a near-optimal policy.

Evaluation. We measure value function estimation accuracy using mean absolute percentage value error (MAPVE), with rollouts to compute the true value estimates. MAPVE = $\frac{1}{t_{test}} \sum_{s \in X_{test}} \frac{|\hat{V}(s) - V^*(s)|}{|V^*(s)|}$, where X_{test} is the set of test states, $t_{test} = 5000$ is the number of samples in the test set,

 $\hat{V}(s)$ is the estimated value of state s and $V^*(s)$ is the true value of state s computed using extensive rollouts. Errors are averaged over 50 runs.

Algorithms. We compare to using several fixed tile-coding (TC) representations. TC uses overlapping grids on the observation space. It is a sparse representation that is well known to perform well for Mountain Car. Puddle World, and Acrobot. We varied the granularity of the grid-size N and number of tilings D, where D is the number of active features for each observation. The grid is either N×N for Mountain Car and Puddle World or N^4 for Acrobot. We explore (D=4, N=4), (D=4,N=8), (D=16,N=4), (D=16,N=8), (D=32,N=4), (D=32,N=8); a grid size of 16 performed poorly, and so is omitted. For Mountain Car and Puddle World the number of features respectively are 64, 256, 256, 1024, 512, 2048, then hashed to 1024 dimensions; for Acrobot, the number of features are 1024, 16384, 4096, 65536, 8192, 131072, then hashed to 4096. Both of these hashed sizes are much larger than our chosen k = 100.

For consistency, once the SCoPE representation is learned, we use the same batch gradient descent update on the MSRE for all the algorithms, with line search to select step-sizes. The regularization weights β_B are chosen from $\{1^{-5}, \ldots, 1^{-1}, 0\}$, based on lowest cumulative error. For convenience, β_w is fixed to be the same as β_B . For learning the SCoPE representations, regularization parameters were chosen using 5-fold cross-validation on 5000 training samples, with $\beta_{\phi} = 0.1$ fixed to give a reasonable level of sparsity. This data is only used to learn the representation; for the learning curves, the weights are learned from scratch in the same way they are learned for TC. The dimension k = 100 is set to be smaller than for tile coding, to investigate if SCoPE can learn a more compact sparse representation. We tested unsupervised sparse coding, but the error was poor (approximately $10 \times$ worse). We discuss the differences between the representations learned by supervised and unsupervised sparse coding below.

Learning curves. We first demonstrate learning with increasing number of samples, in Figure 1. The weights are recom-



Figure 2: Learned representations Φ for 5000 samples and k = 100, respectively for SCoPE, unsupervised sparse coding, and non-negative unsupervised sparse coding, in Puddle World. The representations learned for Mountain Car and Acrobot have similar structure. The non-negative sparse coding has the additional constraint that each entry in Φ is non-negative. The goal for this addition was to determine if further constraints could improve prediction accuracy for unsupervised sparse coding; though the representation qualitatively looks more reasonable, prediction performance remained poor.

puted using the entire batch up to the given number of samples.

Across domains, SCoPE results in faster learning and, in Mountain Car and Acrobot, obtains lowest final error. Matching the performance of TC is meaningful, as TC is wellunderstood and optimized for these domains. For Acrobot, it's clear a larger TC is needed resulting in relatively poor performance, whereas SCoPE can still perform well with a compact, learned sparse representation. These learning curves provide some insight that we can learn effective sparse representations with SCoPE, but also raise some questions. One issue is that SCoPE is not as effective in Puddle World as some of the TC representations, namely 4-4 and 16-4. The reason for this appears to be that we optimize MSRE to obtain the representation, which is a surrogate for the MAPVE. When measuring MSRE instead of MAPVE on the test data, SCoPE consistently outperforms TC. Optimizing both the representation and weights according to MSRE may have overfitting issues; extensions to MSBE or BE, or improvements in selecting regularization parameters, may alleviate this issue.

Learned representations. We also examine the learned representations, both for unsupervised sparse coding and SCoPE, shown in Figure 2. We draw two conclusions from these results: the structure in the observations is not sufficient for unsupervised sparse coding, and the combination of supervised and unsupervised losses sufficiently constrain the space to obtain discriminative representations. For these two-dimensional and four-dimensional observations, it is relatively easy to reconstruct the observations by using only a small subset of dictionary atoms (row vectors of \mathbf{B} in equation (2)). The unsupervised representations, even with additional non-negativity constraints to narrow the search space, are less distributed, with darker and thicker blocks, and more frequently pick less features. For the supervised sparse coding representation, however, the sparsity pattern is smoother and more distributed: more features are selected by at least one sample, but the level of sparsity is similar. We further verified the utility of supervised sparse coding, by only optimizing the supervised loss (MSRE), without including the unsupervised loss; the resulting representations looked similar to the purely unsupervised representations. The combination of the two losses, therefore, much more effectively constrains or regularizes the space of

feasible representations and improves discriminative power.

The learning demonstrated for SCoPE here is under ideal conditions. This was intentionally chosen to focus on the question: can we learn effective sparse representations using the SCoPE objective? With the promising results here, future work needs to investigate the utility of jointly estimating the representation and learning the value function, as well as providing incremental algorithms for learning the representations and setting the regularization parameters.

6 Conclusion

In this work, we investigated sparse coding for policy evaluation in reinforcement learning. We proposed a supervised sparse coding objective, for joint estimation of the dictionary, sparse representation and value function weights. We provided a simple algorithm that uses alternating minimization on these variables, and proved that this simple and easy-touse approach is principled. We finally demonstrate results on three benchmark domains, Mountain Car, Puddle World and Acrobot, against a variety of configurations for tile coding.

This paper provides a new view of using dictionary learning techniques from machine learning in reinforcement learning. It lays a theoretical and empirical foundation for further investigating sparse coding, and other dictionary learning approaches, for policy evaluation and suggests that they show some promise. Formalizing representation learning as a dictionary learning problem facilitates extending recent and upcoming advances in unsupervised learning to the reinforcement learning setting. For example, though we considered a batch gradient descent approach for this first investigation, the sparse coding objective is amenable to incremental estimation, with several works investigating effective stochastic gradient descent algorithms [Mairal et al., 2009, 2010; Le and White, 2017]. The generality of the approach and easy to understand optimization make it a promising direction for representation learning in reinforcement learning.

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