Policy Tree: Adaptive Representation for Policy Gradient

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Abstract

Much of the focus on finding good representations in reinforcement learning has been on learning complex non-linear predictors of value. Policy gradient algorithms, which directly represent the policy, often need fewer parameters to learn good policies. However, they typically employ a fixed parametric representation that may not be sufficient for complex domains. This paper introduces the Policy Tree algorithm, which can learn an adaptive representation of policy in the form of a decision tree over different instantiations of a base policy. Policy gradient is used both to optimize the parameters and to grow the tree by choosing splits that enable the maximum local increase in the expected return of the policy. Experiments show that this algorithm can choose genuinely helpful splits and significantly improve upon the commonly used linear Gibbs softmax policy, which we choose as our base policy.

1 Introduction

The search for a good state representation in reinforcement learning is a challenging problem. Value function based algorithms are not guaranteed to work well in the presence of partial observability (Singh, Jaakkola, and Jordan 1994) or when function approximation is used (Boyan and Moore 1995), and so learning an accurate representation of state is important.

One way to get an adaptive representation of state is to learn a decision tree over the history of observations. The U-Tree algorithm (McCallum 1996) is an example of this. It starts with a single node to represent state, and recursively performs statistical tests to check if there exists a split for which the child nodes have significantly different value. The resulting tree represents a piecewise constant estimation of value.

Such an algorithm would grow the tree whenever doing so improves the value estimate. It is possible for an entire branch of the tree to contain multiple splits over fine distinctions of value, even if the optimum action from each of the nodes in the branch is the same. For example, Figure 1 shows a simple grid world with a single terminal state, indicated by a circle, transitioning to which gives a reward of

Figure 1: A simple world, and a corresponding policy tree

1. The agent is equipped with four sensors to detect whether there is a wall in each direction. Assuming a discount factor of 0.5, the values of the states are shown in each grid cell. To represent an accurate value function, a decision tree would need to split on the current as well as past observations, and eventually distinguish between all the states. In complex domains, one would end up learning a very large tree. In contrast, if we were to represent the policy itself as a function over the observations, we could get a simple and optimal policy in the form of the decision tree on the right.

Policy gradient algorithms are an alternate approach to control in reinforcement learning, which directly optimize a parameterized policy, instead of indirectly optimizing the policy via a value function. Some commonly used algorithms of this type are REINFORCE (Williams 1992), GPOMDP (Baxter and Bartlett 2000) and Natural Actor Critic (Peters and Schaal 2008). Unlike value-based methods, they are guaranteed to converge even when complete information about the state is unavailable. As the previous example demonstrates, the policy function is often simpler and requires fewer parameters than the value function. However, state of the art policy gradient algorithms use a fixed parameterization, and less work has focused on how the policy representation could be learned or improved.

One notable work on adaptive representation for policy gradient includes the NPPG algorithm (Kersting and Driessens 2008). In each iteration of the algorithm, a regression model is learned over a batch of data, with the functional gradient as its target. The final policy is a weighted sum of these models. The regression model can be any complex function of the data, including decision trees. A disadvantage of this method is that each gradient step adds a new model to the policy, increasing the computational cost of action selection, and degrading the generalization ability.

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The reinforcement learning problem we consider involves an agent which interacts with its environment at discrete time steps \( t \in \{1, 2, 3, \ldots \} \). At each time step, the agent perceives its environment via a vector of features \( \phi \), selects an action \( a \in \mathcal{A} \), and receives a scalar reward \( r \in \mathbb{R} \).

The agent chooses an action \( a \) at each time step by sampling a probability distribution \( \pi_\theta(a|\phi) \), where \( \theta \) represents an internal parameterization of its policy. In episodic tasks, the return of an episode is usually defined as the cumulative sum of rewards obtained. In continuing tasks, this sum is unbounded, and the return is defined to be either the average reward per time step or a discounted sum of rewards.

The goal of a learning agent is to find the policy parameters which maximize the expected return from its interaction with the environment, denoted as \( \eta(\theta) \).

Policy gradient works by applying gradient ascent to find a locally optimal solution to this problem. Note that computing the gradient \( \nabla_\theta \eta(\theta) \) would involve an expectation over observed rewards, with the underlying probability distribution being a function of both the policy and the model of the environment. The model is unknown to the agent, but a sample estimate of the gradient can be obtained by observing trajectories of observations and rewards, while acting on-policy. This is the principle underlying all policy gradient algorithms, like REINFORCE, GPOMDP or Natural Actor Critic. Some such algorithms additionally maintain a value function estimate, which reduces the variance of the gradient estimate (Sutton et al. 2000).

The actual policy can be any valid probability distribution over actions that is differentiable with respect to all of its parameters. That is, \( \nabla_\theta \pi_\theta(a|\phi) \) should exist. A commonly used parameterization is the linear Gibbs softmax policy:

\[
\pi_\theta(a|\phi) = \frac{\exp(\theta_a^T \phi)}{\sum_{i=1}^{|\mathcal{A}|} \exp(\theta_i^T \phi)},
\]

where \( \theta_a \) is a \( D \)-dimensional vector of parameters corresponding to action \( a \).

When the actions belong to a continuous space, a common parameterization of the policy is in the form of a Gaussian distribution, with \( \theta_\mu \) and \( \theta_\sigma \) being \( D \)-dimensional parameter vectors specifying the mean and standard deviation of the distribution (Williams 1992):

\[
\pi_\theta(a|\phi) = \mathcal{N}(a|\mu(\phi), \sigma(\phi)),
\]

where,

\[
\mu(\phi) = \theta_\mu^T \phi,
\]

\[
\sigma(\phi) = \exp(\theta_\sigma^T \phi),
\]

\[
\mathcal{N}(a|\mu, \sigma) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left(-\frac{(a-\mu)^2}{2\sigma^2}\right).
\]

### 3 The Policy Tree Algorithm

The Policy Tree algorithm represents its policy using a base parametric policy and a binary decision tree. We assume that the features are \( D \)-dimensional binary vectors \( \phi \in \{0, 1\}^D \). There exist methods, such as Coarse Coding, Tile Coding or Kanerva Coding (Sutton and Barto 1998), which can convert a space of real valued parameters into such a binary vector space.

The internal nodes of the tree are decisions on an element of the feature vector. We call the index of this element the decision index of the node. Every leaf node \( l(\phi) \) in the tree, which is associated with a set of parameters denoted by \( \theta_{l(\phi)} \). The base policy at the leaf is denoted by \( \pi_{\theta_{l(\phi)}}(a|\phi) \). This could be the linear Gibbs softmax policy (Equation 1), or the Gaussian policy (Equation 2), or any other differentiable policy function. If the linear Gibbs softmax policy is chosen, \( \theta_{l(\phi)} \) contains \( |\mathcal{A}| \times |\mathcal{D}| \) parameters. A simple example of a policy tree is shown in Figure 2, in which, based on the feature vector \( \phi \), we use the base policy instantiation corresponding to one of three leaf nodes \( (L, L', L'') \) for action selection.

The high level procedure can be described as:

- Start with a single-node decision tree, with its root node containing a randomly initialized parameterization of the base policy. Repeat the following two stages alternatively:
  - **Parameter Optimization:** Optimize all leaf node parameters using policy gradient for a fixed number of episodes or time steps.
- **Tree Growth:** Keep the parameters fixed for a number of episodes or time steps, while the merit of each split is judged. Choose a leaf node and a decision index on which to split, according to our tree growth criterion. Create two new children of this node, which inherit the same policy parameters as their parent.

During both of the stages, the action at each time step is drawn from the policy function \( \pi_{\theta_t(a|\phi)} \). We describe in detail the two alternating stages of the algorithm in the following sections.

### 3.1 Parameter Optimization

During this phase, the tree structure is kept fixed. Note that \( \frac{\partial}{\partial \theta} \pi_{\theta_t(a|\phi)} \) is known to exist for \( \theta \in \Theta_t(\phi) \), and is trivially 0 for \( \theta \notin \Theta_t(\phi) \), as \( \pi_{\theta_t(a|\phi)} \) is not a function of any such parameter. So, the policy function is differentiable and the parameters can be optimized using any policy gradient technique. Convergence to a locally optimal solution is guaranteed.

The per-step computational complexity during this phase depends on the actual algorithm and parameterization used. Most policy gradient algorithms have a complexity that is linear in the number of parameters. In our case, the number of parameters is \( N_L N_P \), where \( N_L \) is the number of leaf nodes and \( N_P \) is the number of parameters in the base policy \( (N_P = D|A|) \) for the linear Gibbs softmax policy.

### 3.2 Tree Growth

In this phase, the structure of the tree is altered by splitting one of the leaf nodes, changing the underlying representation. In order to choose a good candidate split, we would ideally like to know the global effect of a split after optimizing the resulting tree. This would require making every candidate split and performing parameter optimization in each case, which is unrealistic and inefficient. However, if we suggest a candidate split, and keep its parameters fixed, the gradient of the expected return of this new policy function allows us to compute a first order approximation of the expected return. This approximation is valid within a small region of the parameter space, and can be used to measure the local effect of the split. This is the basis of our criterion to grow the tree. First, we describe a method to calculate the gradients corresponding to every possible split in the tree.

A valid addition to the policy tree involves a split on one of the leaf nodes, on a parameter \( k \in \{1, \ldots, D\} \), such that \( k \) is not a decision index on the path from the leaf node to the root. For every leaf node \( L \) in the tree, and for every valid index \( k \), a pair of fringe child nodes are created, denoted as \( F_{L,k} \) and \( F'_{L,k} \). They represent the child nodes of \( L \) which would be active when \( \phi_k = 1 \) and \( \phi_k = 0 \), respectively. Both of these nodes are associated with a parameter vector which is the same as that of the parent leaf node, that is, for all \( L \) and \( k \):

\[
\theta_{F_{L,k}} = \theta_{F'_{L,k}} = \theta_L. \tag{3}
\]

Let \( \psi_{L,k} \) denote the combination of all the parameters associated with the tree, when it is expanded to include the pair of fringe nodes \( F_{L,k} \) and \( F'_{L,k} \). This is a combination of the vectors \( \theta_{F_{L,k}}, \theta_{F'_{L,k}} \) and \( \theta_L \) for all \( L' \neq L \). Note that each \( \psi_{L,k} \) corresponds to a different policy function, which we denote by \( \pi_{\psi_{L,k}} \). Let \( \eta(\psi_{L,k}) \) denote the corresponding expected return.

Equation 3 ensures that \( \pi_{\psi_{(\cdot),k}}(a|\phi) = \pi_{\theta_t(a|\phi)}(a|\phi) \), which means that all these policies have the same distribution over actions as the one represented by the existing policy tree, even though the underlying representation has changed. This ensures that we can measure the correct sample estimate of the gradient \( \nabla_{\psi_{L,k}} \eta(\psi_{L,k}) \) by following the policy of the tree. It is important to obtain a good estimate of these gradients, to avoid splitting on noise. Therefore, during this phase, the policy is kept fixed while a suitably large number of trajectories are observed, and \( \nabla_{\psi_{L,k}} \eta(\psi_{L,k}) \) is estimated for all \( L \) and \( k \).

We choose the leaf node \( L \) and the decision index \( k \) to split on using the following criterion:

\[
L, k = \arg \max_{L,k} ||\nabla_{\psi_{L,k}} \eta(\psi_{L,k})||_q. \tag{4}
\]

It is worth reflecting on the interpretation of the \( q \)-norm of the gradient vector, in order to interpret our criterion. By using a first order Taylor expansion of the expected return, we can measure the change corresponding to a tiny step \( \Delta \psi \):

\[
\eta(\psi_{L,k} + \Delta \psi) = \eta(\psi_{L,k}) + \nabla_{\psi_{L,k}} \eta(\psi_{L,k})^T \Delta \psi. \tag{5}
\]

If we constrain \( \Delta \psi \) to lie within a small \( p \)-norm sphere with radius \( \epsilon \), we have:

\[
\max_{\Delta \psi} \nabla_{\psi_{L,k}} \eta(\psi_{L,k})^T \Delta \psi = ||\nabla_{\psi_{L,k}} \eta(\psi_{L,k})||_q \epsilon \\
\text{s.t.} \ ||\Delta \psi||_p \leq \epsilon \tag{6}
\]

where, \( \frac{1}{p} + \frac{1}{q} = 1 \).

This shows that the \( q \)-norm of the gradient represents the maximum change in the objective function within a local region of the parameter space bounded by the \( p \)-norm sphere, where \( p \) and \( q \) are the dual norms of each other (Kolmogorov...
and Fomin 1957). Figure 3 shows a graphical representation of various $p$-norm spheres.

By the same reasoning, $||\nabla_{\theta} q(\theta)||_q$ represents the maximum local improvement in the expected return that can be obtained without altering the structure of the tree. A simple stopping condition for tree growth is $||\nabla_{\psi_{L,k}} q(\psi_{L,k})||_q < \lambda||\nabla_{\theta} q(\theta)||_q$, for some $\lambda$.

The fringe gradient $\nabla_{\psi_{L,k}} q(\psi_{L,k})$ has $(N_L + 1)NP$ components, as it is defined over all the parameters corresponding to an incremented tree. However, $(N_L - 1)NP$ of these components are partial derivatives with respect to the existing parameters in the tree, and are shared by all the fringe gradients. Thus we need to measure at most $N_L D$ gradients involving $2NP$ unique parameters, and the per-step computational complexity during this phase for most policy gradient algorithms is $O(N_P N_L D)$. For the linear Gibbs softmax base policy, this is $O(N_L D^2 |A|)$. Note that the number of steps or episodes in this phase will almost always be considerably lower than the previous one, as we simply want a high accuracy gradient estimate and do not need to optimize the parameters.

### 3.3 Fringe Bias Approximation

For many base policy functions, the number of parameters will increase linearly with the number of features, making the complexity of the tree growth phase quadratic in the number of features. Here, we describe an approximation which can reduce this complexity, and applies when the base policy depends on linear functions of the features (as is the case in Equations 1 and 2). If there are $N_F$ such functions, then the number of parameters $N_P$ is equal to $N_F D$. For example, with the linear Gibbs softmax policy we have $N_F = |A|$.

The standard practice when defining a linear function is to augment the input vector $\phi$ with a bias term, usually chosen as the first term of the vector. This term, denoted as $\phi_0$, is always 1. If we had to choose a few components to represent the gradient of the fringe parameters, choosing the parameters associated with $\phi_0$ is a reasonable approximation, which we call the fringe bias approximation. Let $\psi_{L,k}^0$ denote the subset of these parameters in $\psi_{L,k}$.

The bias parameters are always present regardless of the observed features, and are thus indicative of the overall tendency of action selection. The other parameters are present only when the corresponding features are active, and so they can be thought of as context specific tendencies of action selection. The fringe bias approximation looks for features to split on for which the policy would improve if different actions were preferred independent of the rest of the context. However, it ignores improvements due to context specific changes in action selection.

To apply the fringe bias approximation, we compute the gradient of the expected return with respect to these bias parameters, or $\nabla_{\psi_{L,k}^0} q(\psi_{L,k})$. The tree growth criterion remains the same, which is to measure the norm of the gradient in this reduced space. By utilizing only the bias parameters, the computational complexity reduces by $D$ and becomes $O(N_L N_F D)$ (or $O(N_L D |A|)$ for the linear Gibbs softmax policy). As $N_P = N_F D$ for such policies, the per-step complexity for tree growth becomes the same as that for parameter optimization.

### 4 Experiments

We wish to evaluate the following questions empirically:

1. Can Policy Tree improve over the base policy?
2. How well does the fringe bias approximation perform?
3. Is the improvement merely due to an increase in the number of parameters, or does Policy Tree choose intelligent splits?

To answer these questions, we consider a set of domains inspired by arcade games.

#### 4.1 Domains

Our test suite is a set of 4 simple games inspired by arcade games, which have a 16x16 pixel game screen with 4 colours. A pictorial representation of them is presented in Figure 4. All of these games are episodic with a maximum episode length of 256 time steps, and every object moves with a speed of one pixel per step. Objects in these games, including the player agent, enemy agents, friendly agents or bullets are a single pixel in size, and each object type is of a distinct colour. Unless specified otherwise, the actions available to the agent are to move up, down, left, right or stay still. The objective of the learning agent is to maximize the expected undiscounted return.

For generating the binary feature vector, the location of the player agent is assumed to be known. We divide the
The game screen is divided into zones, the size of which grows exponentially with the distance from the agent. Formally, if $(x, y)$ represents the displacement of a pixel from the agent, all pixels with the same value of $\lfloor \log_2 |x| \rfloor$, $\lfloor \log_2 |x| \rfloor$, $\text{sgn}(x)$ and $\text{sgn}(y)$ belong to the same zone. For each zone, we have a feature corresponding to every colour used in the game. A feature is set to 1 if an object of the corresponding colour is found in that zone. An example of feature generation is shown in Figure 5. The left image shows a game screen for Mothership, with the black lines representing the zones centred around the agent. The right screen indicates the active features in each zone via coloured dots (7 features are active in this example). Such a choice of features allows fine distinctions to be made between objects near the agent, while ensuring that the feature size grows only logarithmically with the size of the game screen.

Monsters: The player agent is green. There are three red monsters and three blue power suits randomly located in the game area. The agent can collect and wear a power suit, allowing it to kill the next monster it collides with for one point. A power suit lasts a maximum of 16 time steps once it is worn. The monsters chase the agent when it is not powered, and stay still otherwise.

Switcheroo: There are two types of objects, red and blue. At regular intervals, groups of 8 objects of a randomly chosen type move across the screen horizontally. The player agent starts in the center of the screen, as an object of a random type. Hitting objects of the other type terminates the episode, while hitting an object of the same type earns a point. After a point is earned, the agent switches to the other type.

Mothership: The blue mothership sits at the top of the screen, while 4 rows of 8 red guard ships patrol horizontally below. The green player agent is constrained to the bottom of the screen, and cannot move up or down. It needs to shoot the mothership to collect 5 points, and then shoot the guard ships for 1 point each. Shooting the guards before destroying the mothership simply reflects bullets back at the agent, and alerts the enemy, causing the mothership to move randomly, and the guard ships to shoot bullets. The agent is only allowed to have one active bullet at a time, and there is a random wait of 0 or 1 time steps after its bullet is consumed. All the bullets are coloured yellow.

Rescue: The green player agent has to collect yellow hostages and bring them to the blue rescue location to collect a point. The rescue location is randomly located after every successful rescue, and the player starts from a random location at the top of the screen. The hostages and red enemies move across the screen horizontally at random intervals, and collision with the enemies results in instant episode termination.

These games contain elements of partial observability and non-linearity in the optimum policy function. As examples, the direction of objects in the games cannot be determined from a single game screen, and the best direction to move in Monsters is conditional on multiple variables.

4.2 Experimental Setup

We use the linear Gibbs softmax policy as the base policy for our algorithm, augmented with $\epsilon$-greedy exploration. This policy is defined as:

$$\pi_{\theta_L}(a|\phi) = (1 - \epsilon) \frac{e^{\theta_L^T \phi}}{\sum_{a' \in A} e^{\theta_L^T \phi}} + \epsilon \frac{1}{|A|},$$

The $\epsilon$ term ensures that the policy is sufficiently stochastic, even if $\theta$ takes on heavily deterministic values. This allows accurate estimation of the gradient at all times, and ensures that the policy does not quickly converge to a poor deterministic policy due to noise. Policy Tree benefits a great deal from exploration, as often highly optimized parameters are inherited from parent nodes, and need to be re-optimized after a split. We noticed that adding this exploration term helped the base policy as well.

We compare four different algorithms on these domains. The first is the standard Policy Tree algorithm with the $\epsilon$-greedy Gibbs policy as the base. The second is a version with the fringe bias approximation enabled. The third is a version of the algorithm which chooses a random split during tree growth, for the purpose of testing whether the Policy Tree is just benefiting from having a larger number parameters, or whether it makes good representation choices. And finally, we test the base policy, which represents the standard parametric approach to policy gradient.

We used the GPOMDP/Policy Gradient algorithm with optimal baseline (Peters and Schaal 2006) to measure the policy gradient in all of the algorithms. For Policy Tree, we used a parameter optimization stage of 49000 episodes, and a tree growth stage of 1000 episodes. Splits are therefore made after every 50000 episodes. The value of $\epsilon$ used was 0.001. For the tree growth criterion, we choose the $q = 1$ norm of the gradient. A stopping criterion parameter of $\lambda = 1$ was used.

For each algorithm, we performed 10 different runs of learning over 500000 episodes. The average return was measured as the moving average of the total undiscounted return per episode with a window length of 50000.

4.3 Results

The learning curves of the Policy Tree algorithm as compared to the base policy are shown in Figures 6 to 9. The
standard error in the results, across the 10 runs, is represented by the vertical error bars in the graphs. These results allow us to answer the questions posed earlier. On our four domains, we see that:

1. The Policy Tree algorithm improves upon the underlying base policy with statistical significance.

2. The fringe bias approximation trails the exact measure in most domains, but improves over the linear parametrization significantly, without enduring additional computational complexity during tree growth.

3. An arbitrary increase in the number of parameters via the random splitting does not improve performance at all. This shows that the our tree growth criterion contributes significantly to the effectiveness of the algorithm.

The Monsters domain shows the biggest improvement, as well as the greatest gap between the exact and approximate versions. In this domain, the feature that represents whether or not the agent is powered is very informative. Policy Tree chooses this as the first split in 80% of the runs, while this drops to 20% with the approximation enabled. However, we observed that the approximate version outperforms the base policy even when this split is never made, indicating that the chosen splits are not meaningless.

5 Future Work

There are a number of areas in which this work can be expanded. The current version of our algorithm has fixed lengths for the parameter optimization and tree growth phases. One could test for convergence of the objective function during optimization, allowing the tree to get the best possible performance before we try to expand the structure. However, highly optimized policies tend to be highly deterministic, making re-optimization of the parameters after a split trickier. The Gibbs policy, for instance, has a low gradient in regions of high determinism. It is possible that the use of natural gradients would alleviate this problem, by measuring the change of the objective function with respect to the actual change in the policy distribution, rather than the change in parameter values.

The algorithm as described in this paper is restricted to policy functions defined on binary features. However, in general one could use any arbitrary base policy (defined, say, on continuous features), as long as there exists a method to perform a binary coding of these features, which the decision tree can split on. Future work could deal with better ways to handle continuous features, similar to the Continuous U-Tree algorithm (Uther and Veloso 1998).

Due to the presence of noise in the gradients, or due to the locality of our improvement measuring criterion, it is possible that some splits do not enhance the policy significantly. This causes needless increase in the number of parameters, and slows down learning by splitting the data available to each branch. A possible fix for this would be to prune the tree if optimization after a split does not significantly change the parameters.

The splits in the decision tree are currently based on the value of a single observation variable. In general, we could
define a split over a conjunction or a disjunction of observation variables. This would increase the size of the fringe used during tree growth, but would allow the algorithm to find splits which may be significantly more meaningful. A different kind of split in the decision nodes would be on a linear combination of features. This can be viewed as a split on a non-axis aligned hyperplane in the observation space. As there are infinite such splits, it is not possible to measure them using our fringe structure. However, there may be ways to develop alternative criterion in order to grow a tree with such a representation. Prior research suggests that such an architecture is useful for classification but not for regression (Breiman et al. 1984). It is unclear how useful it would be in the search for an optimal policy.

The two phase structure of our algorithm is slightly sample inefficient, as the experience during tree growth is not used to optimize the parameters. In our experiments, 2% of the episodes are unused for optimization. Due to the requirement to stay on-policy to estimate the gradient, this is difficult to avoid. One possible solution would be to use importance sample weighting and utilize off-policy trajectories to compute the gradient. This would in fact avoid the necessity of keeping the policy fixed during the tree growth stage. The use of importance sampling in policy gradient has been studied previously (Jie and Abbeel 2010). However, the weights for the off-policy samples would reduce exponentially with the horizon of the problem, and we cannot be certain whether it is possible to have a reliable off-policy estimate of the gradient in most domains.

6 Conclusion

We presented the Policy Tree algorithm, and demonstrated its utility on a variety of domains inspired by games. This algorithm has the same convergence guarantees as parametric policy gradient methods, but can adapt its representation whenever doing so improves the policy. We also present an approximate form of this algorithm, which has a linear computational complexity per time-step in the number of parameters during the entire learning phase. We believe that these results indicate that the gradient of the expected return is a useful signal in the search for representation in reinforcement learning.

References


