Single-Agent Optimization Through Policy Iteration Using Monte-Carlo Tree Search

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Abstract

The combination of Monte-Carlo Tree Search (MCTS) and deep reinforcement learning is state-of-the-art in two-player perfect-information games. In this paper, we describe a search algorithm that uses a variant of MCTS which we enhanced by 1) a novel action value normalization mechanism for games with potentially unbounded rewards (which is the case in many optimization problems), 2) defining a virtual loss function that enables effective search parallelization, and 3) a policy network, trained by generations of self-play, to guide the search. We gauge the effectiveness of our method in “SameGame”—a popular single-player test domain. Our experimental results indicate that our method outperforms baseline algorithms on several board sizes. Additionally, it is competitive with state-of-the-art search algorithms on a public set of positions.

Introduction

Single-agent optimization problems have been an active field of research for decades. Such problems include any domain with an agent whose goal is to maximize an objective function(s), without interference from any other agents. NP-hard problems such as the Travelling Salesman Problem (TSP) can be framed as a single-agent optimization problem. Algorithms for solving TSP have many practical uses, such as computer chip design and order-picking in warehouses (Theys et al. 2010). Single-agent optimization problems can be represented as (deterministic) single-player games. This is the term used throughout this paper to better present our contribution in relation to previous work.

Most state-of-the-art heuristic search algorithms for single-player games use Monte-Carlo simulations (Schadd et al. 2012), (Cazenave 2009), (Rosin 2011). These methods estimate the values of states using random simulations. The generality of these methods makes them applicable to a wider variety of domains. Examples include two-player board games such as Go (Gelly et al. 2012) and Hex (Arneson, Hayward, and Henderson 2010), (Huang et al. 2013), real-time domains such as Ms. Pac-Man (Pepels, Winands, and Lanctot 2014), general video game playing (Perez et al. 2019).

A recently proposed enhancement is to combine MCTS with deep reinforcement learning (Silver et al. 2018), (Anthony, Tian, and Barber 2017). These algorithms have achieved state-of-the-art performance in deterministic two-player perfect-information zero-sum games. MCTS provides the agent with the ability to look ahead, while policy and value networks are used to decrease the width and depth of the search tree, respectively. Additionally, the trained policy networks can be surprisingly strong by themselves. For instance, AlphaZero’s Go policy plays at human expert level without the need for forward search.

In this paper, our goal is to bring the ideas of these algorithms to single-player optimization problems. There are multiple differences between the two settings that make this task non-trivial. In zero-sum two-player games, the reward seen by the agent is often one of \{-1, 0, 1\}, for loss, tie, or win, whereas in single-player games, the reward are unknown. Furthermore, values found during search are a lower-bound on the optimal value. Therefore, the action selection strategy, as well as the policy target, have to be adjusted.

To address these problems, after a discussion of related work, we first introduce our policy-guided MCTS algorithm for single-agent optimization problems, then describe how we train our policy networks, and finally measure the performance of our methods in the NP-hard SameGame—a popular single-agent optimization domain—before closing with concluding remarks and suggestions for future work.

Related Work

We begin by introducing several state-of-the-art search algorithms for single-player games. Then, pioneer works on combining MCTS and deep reinforcement learning for two-player games are presented. Lastly, we discuss similar work on single-player games.

MCTS in Single-Player Games

Schadd et al. introduced Single-Player MCTS (SP-MCTS) (Schadd et al. 2012), which was the first successful application of MCTS to a single-player game with a large state and action space. Two contributions which we also use in our work are creating a tree-per-move rather than tree-per-game by playing the highest valued action, and a few value normalization methods.

Cazenave developed a recursive variant of MCTS called Nested Monte-Carlo Search (NMCS) (Cazenave 2009). NMCS estimates the values of states at level $k$ using a recursive call of level $k - 1$, where a level 1 recursion is a Monte-Carlo rollout.

Rosin’s Nested Rollout Policy Adaptation (NRPA) (Rosin 2011) combines NMCS with online policy learning. NRPA’s rollouts are guided by a policy, which is slowly adapted towards the moves with the highest return. An efficient en-
coding of states, which transforms similar states to the same value, is required for the algorithm to perform well.

**MCTS and Deep Reinforcement Learning**

The combination of MCTS and deep reinforcement learning has been successfully applied to two-player games. One of the major contributions is the work on AlphaZero (Silver et al. 2018), which reached superhuman performance in the games of Go, Chess, and Shogi. The algorithm uses a single two-headed network, which outputs both a policy and value of a state. The policy is used during the selection step of MCTS, and the simulation phase is replaced by the value prediction. The network is trained from scratch using self-play, without the use of any human generated data.

A similar work to AlphaZero is Expert Iteration (Anthony, Tian, and Barber 2017). Expert Iteration trains a new neural network at each generation, starting with only a policy network, which is used in a similar way as AlphaZero. Once the generated data is of sufficient quality, the policy network is replaced by a two-headed network. The value prediction of the network is combined with the rollout result using a mixing parameter, which further improves the agent.

We are not the first to combine MCTS with deep reinforcement learning for single-player games. The work of (McAleer et al. 2018) trains a two-headed network to solve the Rubik’s cube, using a training procedure called Autodidactic Iteration (ADI). ADI is only suitable for problems in which there is a single known goal state. This is in contrast to our algorithm, which—as we will see—does not have this requirement.

Laterre et al. introduce Ranked-Rewards (R2 (Laterre et al. 2018)), a general algorithm that enables self-play for single-player games. This is accomplished by setting the rewards seen by the agent to a value of either 1 or −1, depending on whether the actual value of the state is higher or lower than a given percentile of previously seen rewards. This pits the agent against itself, forcing it to continuously outperform previous generations. Our algorithm does not make use of this self-play scheme.

**Policy-Guided MCTS for Single-Agent Optimization**

Our work is based on the belief that combining MCTS and deep reinforcement learning can also lead to state-of-the-art performance in single-player games. Therefore, our objectives are threefold: 1) develop an effective variant of MCTS for single-agent domains, 2) establish a suitable learning target for the policy network, and 3) create a training procedure similar to the self-play regime for two-player games.

**Integrating Policies into MCTS**

Our initial task is to develop a policy-guided MCTS algorithm for single-player domains. To this end, we first introduce a normalization strategy that is applied during the selection stage of MCTS. Having normalized values allows us to use the standard Predictor + Upper Confidence Bound For Trees (PUCT) (Silver et al. 2018) selection strategy, which we introduce in Eq. [3]. We additionally use the policy for the initial action selection in expanded nodes, as well as to guide the rollouts. Using a neural network during search is expensive however, so we define a virtual loss function that enables tree parallelization (Chaslot, Winands, and van Den Herik 2008). Parallelization allows us to do batch prediction, which increases GPU efficiency, thereby increasing the speed of the search.

**Value Normalization**

To use the PUCT selection strategy, the rewards need to be in the range of [−1, 1]. In single-player games however, the range of rewards is often different, and unknown. Therefore, we require a robust method for normalizing the values.

Several normalization techniques have been introduced in previous work (Schadd et al. 2012), (Klein 2015). In SP-MCTS, the values are not normalized, instead larger parameters are used in the selection strategy, which increases the upper confidence bound (UCB) term. The work of (Klein 2015) uses the highest score achievable in any SameGame board to normalize the values. The main problem with both strategies is that they are domain specific, and require either extensive experiments or domain knowledge to set properly.

We propose a more general normalization method—max-min scaling—that is applied locally at the node level as follows:

\[
Q_{\text{norm}}(s, a) = \frac{2(Q(s, a) - \min_a Q(s, a'))}{\max_a Q(s, a') - \min_a Q(s, a')} - 1 \quad (1)
\]

When no action has been taken in state \(s\) (i.e., max and min values are not yet defined), or they are equal, we set \(Q_{\text{norm}}(s, a)\) to 1—being optimistic. The highest and lowest values can be stored in a node directly, or calculated by looping over all edges. This approach does not require any assumptions about the lowest or highest achievable values in the domain. Furthermore, it does not require the tracking of standard deviations to relate average values and UCB exploration terms (Schadd et al. 2012). Lastly, since \(Q\) is defined locally, action values of one node do not impact other nodes, which is a desirable property.

Note that max-min normalization maximizes the spread of mapped action values: we can expect at least one action with a value of 1, and after several simulations, at least one action with a value of −1. Assuming a scenario where the maximum and minimum values are very similar, and the reward bounds are known, max-min normalization can lead to a higher level of exploitation than standard UCT (Kocsis and Szepesvári 2006). One method for reducing the spread of mapped action values is to use max-min normalization with the actual rewards, rather than average state-action values. Since both methods normalize values to range [−1, 1], we can expect UCB-based selection strategies to converge to the globally optimal solution.

**MCTS Stages**

In what follows, we describe the four stages of our MCTS algorithm that uses max-min scaling. PUCT, and virtual loss enabling effective parallelization:

1. **Selection.** The selection strategy is applied recursively until an edge of a leaf node or an edge leading to a terminal node is selected. At each node, the action with the highest sum of value and upper confidence bound, corrected for virtual loss, is selected:

\[
a' = \arg\max_a \left( (Q(s, a) - L(s, a))_{\text{norm}} + U(s, a) \right), \quad (2)
\]
where the upper bound \( U(s,a) \) is given by PUCT, which is calculated as follows:

\[
U(s,a) = c_{\text{puct}} \pi_\theta(s,a) \sqrt{\frac{N(s)}{1+N(s,a)}}. \tag{3}
\]

This selection strategy is initially focused on actions with high prior probability and low visit count, but asymptotically prefers actions with high values. When an edge is selected, the virtual loss count (which is described below) and visit count are increased: \( W(s,a) \leftarrow W(s,a) + 1 \), \( N(s,a) \leftarrow N(s,a) + 1 \).

2. Expansion. A leaf node \( n_L \) is expanded on the first visit and added to the tree. All child edges, one per legal action \( a \), are created and initialized to \( \{N(s_L,a) = W(s_L,a) = Q(s_L,a) = Q_{\text{total}}(s_L,a) = 0 \} \), where \( s_L \) is the state of node \( n_L \). The prior probability is also stored in the edge, which is the re-normalized output of the network after filtering out illegal actions. We lock the node and put the thread to sleep until the network is finished evaluating the state.

3. Simulation. The edge with the highest prior probability is selected from the newly expanded node \( n_L \) as the first action in the simulation. The rest of the simulation is either uniformly random among all valid actions, or is guided by the current policy. The policy-based rollout selects an action in state \( s \) by sampling from the policy. The simulation is finished once a terminal state is reached.

Right after the first rollout is finished, the \( \bar{Q} \) value of all the edges is set to the value of the simulation. This gives the other edges the opportunity to be selected independently of the success of the first simulation. This initialization strategy is “optimistic”, as the greedy action is selected in a newly expanded node. Another option for optimistic initialization is to give edges that have not yet been tried a normalized value of 1.

The policy-based rollout is far more informed, but is much slower than the random counterpart, since a network prediction is required at each step of the simulation. We use policy-guided simulations for training, and compare the two simulation strategies when testing the strength of the final policy.

4. Backpropagation. The result \( R \) of the rollout is propagated to the root, updating edge statistics along the way as follows: \( Q_{\text{total}}(s,a) \leftarrow Q_{\text{total}}(s,a) + R, \bar{Q}(s,a) \leftarrow Q_{\text{total}}(s,a)/N(s,a), W(s,a) \leftarrow W(s,a) - 1 \).

Tree Parallelization and Virtual Loss The speed of Policy-MCTS is considerably lower than plain MCTS, since it requires a prediction from the policy network at each expanded node, and potentially at every step of the rollout. The slowdown caused by the network can be reduced by using batch predictions, which requires a parallel version of the algorithm. Common parallelization strategies for MCTS are root, leaf, and tree parallelization (Chaslot, Winands, and van Den Herik 2008). In our implementation, we use tree parallelization with node and edge mutexes, and virtual loss.

In our tree parallelization, all search threads work on the same tree, with mutexes used to avoid data corruption. However, given that both UCT and PUCT are deterministic, we can expect the majority of threads to take similar paths down the search tree. To discourage this behaviour, we can add a temporary virtual loss to actions as they are selected (Chaslot, Winands, and van Den Herik 2008). In two-player games, virtual loss corresponds to virtual rollouts that resulted in a loss. However, in our setting, we have a score with unknown bounds, so we have to define what a “loss” means. We define virtual loss as \( L(s,a) = wW(s,a)|\bar{Q}(s,a)| \), where \( W(s,a) \) is the virtual loss count stored in the edges of the tree, and \( w \) is the global virtual loss weight, which is subject to optimization. Therefore, the loss is relative to the current state-action value. A benefit of this approach is that it requires no knowledge about the bound of rewards.

We optimized \( w \) with respect to the final average score obtained. The experiments used plain MCTS on 100 randomly generated 15 × 15 boards, with 5 runs per board. We discovered that a value of 0.01 provided the highest search speed, and was within a few points of the best average (not statistically significant); we fixed \( w = 0.01 \) in all of our experiments.

Policy Training Target In two-player adversarial games, assuming a limited search time budget for MCTS, the edge with the highest visit count—as opposed to the highest valued edge—is often selected as the action to play (e.g., (Silver et al. 2018) and (Anthony, Tian, and Barber 2017)). Actions with higher visit counts are considered more robust, as they guard against the case in which a newly analyzed move with higher value, but much fewer simulations, is overconfidently chosen.

By contrast, in single-player games, which are non-adversarial, the values of simulations starting in a state are a lower bound on the maximum achievable value. This means that the action with the highest simulation value is currently the best action, regardless of how often other actions were attempted. In the limit, the action with the highest visit count will also have the highest value (Kocsis and Szepesvári 2006). However, this might not be the case when given a limited search budget.

Given the above observation, we set the target for the policy as follows, with ties between equal valued actions being broken randomly: \( \pi(s_i,a) = 1 \) if \( a = a_i \) and 0, otherwise.

Using this policy target, training reduces to a supervised learning task that seeks to minimize the average cross-entropy loss between the target policy \( \pi \) and its approximation \( \pi_\theta \), for all training samples. I.e., for a mini-batch of size \( B \), its loss \( L \) is given by: \( L = -\frac{1}{B} \sum_{i=1}^{B} \pi(s_i)^T \log \pi_\theta(s_i) \).

Data Generation In many single-player games, the initial actions have a large impact on the final score the agent can obtain. An intuitive choice, then, is to let the agent spend the majority, if not the entire MCTS budget, on determining the best initial move. This would allow the agent to come up with the best first move it possibly can. However, once the search budget is spent, the entire sequence, and not just the first move, is reported. Since the entire search budget has been spent, there is no opportunity to optimize the rest of these moves further. It has been shown in (Schadd et al. 2012), as well as confirmed by our own experiments, that committing to an action
after a fraction of the search budget is spent, thereby allowing the search to optimize the remaining move sequence, can produce better results.

Committing to actions corresponds to playing a game; once a decision has been made, it cannot be reversed. This allows MCTS to spend more time in deeper sections of the search tree, meaning actions near the middle and end of the tree receive more of the search budget than they would otherwise. Therefore, we can expect the resulting action sequences to be better.

Inspired by this observation, our agent interleaves planning and playing. In each state, the agent receives a constant planning budget of $k$ MCTS simulations. Once the planning budget is spent, the best action, which is the one with the highest value, is taken. Although this forces the agent to commit to earlier actions, it opens up the opportunity to better optimize subsequent moves. This process is repeated until a terminal state is reached. Then, all state-action pairs taken to reach the final state are returned, to be used as training data.

While training, we add Dirichlet noise to the prior probability of all actions $a$ at the root of the tree — using $(1 - \epsilon)\pi_0(s_{root}) + \epsilon \text{Dir}(\alpha)$ instead of $\pi_0(s_{root})$. Dir$(\alpha)$ is a random vector with $L_1$-norm of 1. Using a low $\alpha$ value will add a high value of noise to a few moves, whereas a higher value will add a more uniform amount of noise to a larger number of moves. The added noise has the potential to increase the UCB value of actions the current policy believes to be bad, which encourages exploration.

The value of $\alpha$ was chosen experimentally. We used $\epsilon = 0.25$, as proposed in (Silver et al. 2018), and tried a variety of $\alpha$ values during training. These experiments were run for a few generations, and the $\alpha$ that produced the highest average value during training was selected for the experiments presented in the experiments section. The value of $\alpha$ is dependent on the average number of legal moves per game: for the $7 \times 7$ board, we tried the following values $\{0.5, 0.75, 1.0, 1.25\}$, whereas for the $15 \times 15$ board we tried $\{0.15, 0.25, 0.4\}$. From these, we extrapolated a value for the $10 \times 10$. These parameters are presented in Table 1.

**Training Procedure**

Our training procedure alternates between policy evaluation and policy improvement, in a process known as policy iteration. Policy iteration is guaranteed to converge to an optimal policy in the tabular case. This guarantee no longer holds in the function approximation case. In spite of this, deep neural networks trained by policy iteration have surpassed human players in multiple games (Silver et al. 2018), (Anthony, Tian, and Barber 2017).

Using MCTS in conjunction with the current policy to play complete games constitutes the policy evaluation step. Because MCTS helps the agent to find better moves than what is suggested by the policy alone, it acts as an policy improvement operator. MCTS is integral to the process: the strength of the trained policy is correlated with the effectiveness of MCTS as the policy improvement operator.

Our policy is trained in generations, with the data from previous generations used to train the next generation’s policy network. The training of each generation is synchronous, and constitutes a complete policy iteration step. The first policy network is trained using data generated by MCTS using the uniform random policy. Subsequent generations combine MCTS and the current iteration of the policy to generate data. To jump-start the learning process, the first generation is run using more simulations per step; since no policy network is used, there is not a large run-time cost to this.

The training procedure (Algorithm 1) works as follows: We use two queues, $B_{\text{training}}$ and $B_{\text{validation}}$, for training which store state-action pairs $(s_t, a_t)$. As each run finishes, the resulting pairs are stored in a temporary buffer, which only contains data produced in the current generation. Once all runs in a generation are finished, the pairs stored in the temporary buffer are shuffled and split (e.g., 90-to-10), and appended to both buffers, respectively. This ensures that both buffers will contain data from multiple generations. Note that using a single buffer and randomly splitting it before training is not equivalent to this procedure.

The sizes of the buffers determine the amount of data that is kept from older generations; given more data, we can expect the policy to be better (Anthony, Tian, and Barber 2017), but the training time will be longer. The size of the buffers is also directly related to the amount of data generated per run. We use the validation buffer to avoid overfitting by early stopping.

**Experimental Setup**

SameGame

SameGame is a tile-matching game with the goal of maximizing the final score. In each move, the player can clear horizontally or vertically connected groups of size two or more of equal colour. The blocks above created holes will always fall down and then move left, if possible. When an entire column is cleared, the columns to the right are moved to the left. Each move scores $\#\text{Blocks Cleared} - 2$. The

**Algorithm 1:** Policy training procedure

1. **Input:** #generations $G$, #runs $N$, training/validation buffer lengths $l_t, l_v$, training-validation split percentage $\lambda$
2. **Output:** Trained policy network
3. $\pi_0 \sim \text{Uniform-Over-Valid-Actions}()$
4. $B_{\text{training}} = \text{queue}(\text{len}=l_t)$
5. $B_{\text{validation}} = \text{queue}(\text{len}=l_v)$
6. for $g = 1, ..., G$ do
7.   $B = []$
8.   for $r = 1, ..., N$ do
9.     $s \leftarrow \text{Generate-Random-State}()$
10.    $[(s_1, a_1), ..., (s_T, a_T)] \leftarrow \text{Policy-MCTS}(s, \pi_{g-1})$
11.    $B \leftarrow B + [(s_1, a_1), ..., (s_T, a_T)]$
12. end
13. $B_{\text{training}} \leftarrow B_{\text{training}} + T$
14. $B_{\text{validation}} \leftarrow B_{\text{validation}} + V$
15. $\pi_g \leftarrow \pi_{g-\text{random}}$
16. $\text{Train}(\pi_g, B_{\text{training}}, B_{\text{validation}})$
17. end
18. **Return:** $\pi_G$
game is over when the player cannot take an action anymore. In addition, if the board is cleared, the player is awarded an additional 1,000 points. Otherwise, the player receives a penalty based on the total number of blocks left, which is calculated as follows: \( \sum (\text{Blocks Left}, -2)^2 \).

Deciding whether a general SameGame instance with at least five colours and two columns can be fully cleared has been shown to be NP-complete (Schadd et al. 2008). We use boards of size 7 × 7, 10 × 10, and 15 × 15, with five different block colours. All boards used for training and testing are randomly generated. We slightly simplify the action space by only allowing the agent to select the lowest left block in a group; in the actual game, a player can click on any of the blocks in a group to clear it.

Policy Network Architecture

We use the same network architecture for all of the experiments to limit its impact on the results. The complexity of the network makes it suitable for the 15 × 15 boards, but it is likely too deep for the simple 7 × 7 boards.

Our architecture is similar to the one used by (Anthony, Tian, and Barber 2017). In particular, the input to the network is an “image” of size \( d \times d \times (c + 1) \), where \( d \) is the dimension of the board and \( c \) is the number of block colours, and 1 is added to encode empty tiles. That is, the board is represented as \( c + 1 \) binary layers to one-hot-encode the tile state. The input is padded by 1 on all four sides, increasing the dimension to \( (d+2) \times (d+2) \times (c+1) \). This ensures that the information at the edges of the board is not lost during convolutions. The padded input is passed to 13 convolution layers, all of which have 64 filters, stride of 1, and ELU activation (Clevert, Unterthiner, and Hochreiter 2015). Layers 11 and 13 have a kernel size of \( 1 \times 1 \), and all others have size \( 3 \times 3 \). The dimension of the input is kept for layers 1-8 and 12, and reduced in layers 9-11 and 13. The output of the final convolution layer is flattened and passed to a linear layer with softmax activation, which represents the policy.

We use Adam as the optimizer, with a learning rate of \( 5 \times 10^{-4} \). Data is fed to the network in mini-batches of size 256. For regularization, early stopping on validation loss is used. The early stopping point is the first epoch after which the validation errors do not decrease for 3 consecutive epochs. We set the size of the buffers \( B_{\text{training}} \) and \( B_{\text{validation}} \) to 1.5M and 150K, respectively, for all experiments. Given the number of runs per generation, as provided in Table 1, the buffers contain data from roughly 3 to 5 previous generations.

Experiments

In this section we will validate our contributions with several experiments in the SameGame domain. The first compares the performance of Policy-MCTS against plain MCTS, which is MCTS with a uniform random policy among all valid actions. Our experiments are run on three different board sizes using several time budgets. To demonstrate the effectiveness of our parallelization method, we also compare the performance of single and multi-threaded variants of both algorithms. To put our results into perspective, we also use Policy-MCTS to solve a standard set of 20

| Parameter          | 7 × 7 | 10 × 10 | 15 × 15 |
|--------------------|-------|--------|--------|
| Generations        | 50    | 50     | 66     |
| Runs/Generation    | 20,000| 10,000 | 5,000  |
| Simulations/Move   | 100   | 50     | 25     |
| \( c_{\text{puct}} \) | 30    | 4      | 2      |
| Threads/Run        | 1     | 1      | 1      |
| Dirichlet Noise    | 0.75  | 0.40   | 0.25   |
| Training Time      | 3 days| 3.5 days| 6.5 days|
| CPU                | Intel | Intel  | 2 x Intel |
| GPU                | Nvidia| Nvidia | 4 x Nvidia |
| - Type/RAM         | i7-7700K | i7-8700K | Xeon 6148 |
| - Cores/Speed      | 4/4.2 GHz | 6/3.7 GHz | 20/2.4 GHz |
| GPU                | Nvidia| Nvidia | 4 x Nvidia |
| - Type/RAM         | 1070/8GB | 1070/8GB | V100/16GB |

Figure 1: The average score of the 15 × 15 policy at each generation of training.

SameGame test positions. These boards are commonly used to benchmark state-of-the-art search algorithms for single-player games. We demonstrate that a policy trained using only 25 simulations per move is competitive with these methods.

Comparing with Plain MCTS

We use our algorithm to train policy networks for board sizes 7 × 7, 10 × 10, and 15 × 15. The 7 × 7 board is small and relatively simple. The other two sizes are increasingly more complex, with the 15 × 15 board often used as benchmark problem in the literature. The average solution length of 10 × 10 board games is roughly twice that of 7 × 7, and half of 15 × 15 games. To keep training time manageable, we decrease both the number of runs and simulations as the board size increases.

The parameters we used for training is shown in Table 1. Note that while we put effort into optimizing the parameters, they are most likely not optimal. This is because each experiment takes several days to run, making a grid search over these parameters not feasible, given our computation budget. Training on 7 × 7 and 10 × 10 boards converged, but the 15 × 15 policy did not (Figure 1).

Given the same node budget, we can expect Policy-MCTS to outperform plain MCTS. However, using a GPU incurs a large cost on the speed of the search. Therefore, to make a more fair comparison, we use wall-clock time for all of our

Table 1: Parameters used for training for each board size.
Table 2: Number of simulations, node expansions, and leaf node expansions for single-threaded and multi-threaded experiments on the 15 × 15 boards.

| Parameter       | 1 Second | 5 Seconds | 15 Seconds |
|-----------------|----------|-----------|------------|
|                 | MCTS     | P-MCTS Random | P-MCTS Informed |
| Simulations     | 20,106   | 1,274     | 27         |
| Expansions      | 12,074   | 1,261     | 27         |
| Leaf Expansions | 155      | 1         | 0          |
|                 | 403,243  | 80,580    | 398        |
|                 | 27,089   | 6,128     | 131        |
|                 | 809      | 150       | 0          |
|                 | 2,297,200| 376,698   | 2,575      |
|                 | 697,810  | 49,538    | 864        |
|                 | 310,313  | 46,295    | 864        |
|                 | 5,522    | 110       | 0          |
|                 | 15,524   | 2,691     | 0          |

|                 | MCTS     | P-MCTS Random | P-MCTS Informed |
|                 | 95,335   | 9,652       | 201           |
|                 | 91,140   | 9,652       | 201           |
|                 | 148      | 0           | 0             |
|                 | 2,297,200| 376,698    | 2,575         |
|                 | 697,810  | 49,538     | 864           |
|                 | 310,313  | 46,295     | 864           |
|                 | 5,522    | 110        | 0             |
|                 | 15,524   | 2,691      | 0             |

Since we are using wall-clock time, we also present the averages of simulation count, terminal node expansions, and node expansions for the experiments on the 15 × 15 boards in Table 2. The simulation count also contains simulations starting (and ending) at terminal leaf nodes, i.e., terminal states that are part of the search tree. The number of terminal node expansions is included in the node expansions count.

We use a test set of 500 randomly generated boards for each size, to compare Policy-MCTS with plain MCTS. We do 5 runs per board, for a total of 2,500 runs in each experiment. Results at three different time settings of 1, 5, and 15 seconds are shown for plain MCTS, and Policy-MCTS with random and policy guided simulations, respectively.

The same parameters are used for both the single-threaded and multi-threaded experiments. Additionally, the \( c_{puct} \) value used during training is kept for all runs. The number of threads used by MCTS is 16, which maximizes CPU usage, and Policy-MCTS uses 100 threads, which is sufficient to keep the GPU fully loaded. By using such a high number of threads, we are assuming that the extra exploration caused by virtual loss is mitigated by the policy network.

All experiments were run using the same computer that was used to train the 10 × 10 network. The experimental results are shown in Figure 2. We observe that the multi-threaded versions of all three algorithms outperform their single-threaded counterparts; for the 15 × 15 board, the average score of plain MCTS at the 5 second mark rises from 1,684 to 2,393, an increase of nearly 50%. This provides some evidence that our parallelization strategy is effective. Furthermore, we can see that using multiple threads is more beneficial for both versions of Policy-MCTS. This is expected, as using many threads allows for batch prediction, which utilizes the GPU more effectively. We can also clearly observe the downside of using a neural network when given a short time limit, with plain MCTS outperforming both versions of Policy-MCTS at the 1 second mark, in all configurations.

Another important observation is that as the decision complexity increases, the strength difference between plain MCTS and Policy-MCTS also increases. Given a simpler problem and a limited budget, it is probably more beneficial to use plain MCTS. However, as the complexity of the problem increases, so too does the benefit of using a pol-
icy. This is clearly demonstrated in the 15 × 15 graphs, with Policy-MCTS far outperforming plain MCTS.

A rather surprising result is the weak performance of multi-threaded Policy-MCTS with informed rollouts, when compared to plain MCTS, on the 15 × 15 board. This leads us to speculate that given a limited time budget on a problem with a large state and action space, creating a deeper tree by performing many weak rollouts is better than a much smaller, more informed number of rollouts.

Detailed performance data for these experiments is provided in Table 2. For the 15 × 15 board, even with multi-threading, only 2,575 policy guided rollouts were finished, compared to over 2 million for plain MCTS. Note that in the single-threaded setting, 400 informed rollouts outperform over 400k random rollouts from plain MCTS. This provides evidence that our algorithm has trained a competent policy. We can also observe that the multi-threaded versions are far more explorative. For example, for Policy-MCTS with random rollouts in the 5 second setting, node expansions per simulation increases from 0.45 to 0.93.

**Comparison to State-of-the-Art Algorithms**

In this section, we compare the performance of our parallel MCTS and Policy-MCTS algorithms against published state-of-the-art search methods on 20 public test positions. We use the network trained on 15 × 15 boards from the previous section, and run Policy-MCTS with random and guided rollouts, respectively; the data for the former is not presented as it performed worse than the latter. We additionally run our parallel version of MCTS to gain a better perspective of the results obtained in the last section. $c_{\text{exact}}$ is set to 5 and 10 and threads to 120 and 80 for Policy-MCTS and Parallel-MCTS, respectively. The same machinery that was used to train the 15 × 15 policy is used. All algorithms are run only once per position. The results of the experiment are provided in Table 3.

Note that we give each algorithm only 2 hours total per position, which is similar to SP-MCTS (Schadd et al. 2012). The results of Dist-NRPA(5) (Negrevergne and Cazenave 2017) are also achieved in 2 hours, but they use 160 CPU’s for each position. Algorithms NMCS(4), Sel-NRPA(4) (Cazenave 2016), and Div-NRPA(4) (Edelkamp and Cazenave 2016) take more than half a day per position; the number in the brackets represents the nesting level used by the algorithm. Most of these algorithms, with the exception of Div-NRPA(4) and Dist-NRPA(5), also make use of hand-crafted heuristics to guide the rollouts.

The score can potentially be increased in multiple ways. We could evenly distribute the time budget over the average number of moves, as done in SP-MCTS. Additionally, an implementation capable of using multiple GPUs could potentially achieve much higher scores with the same policy. Using a transposition table to store predictions will also increase the speed of the system. Smaller gains could be obtained by fine-tuning the search parameters; we did not tune any of the parameters to these positions. Lastly, we can increase the simulation count of MCTS during training. This greatly increases the training time, but it also strengthens the policy evaluation and improvement steps, which can lead to a much better policy.

**Conclusion and Future Work**

In this paper, we presented three main contributions. We first introduced a novel action value normalization method, which is more general and therefore applicable to a wider range of domains. Then, we defined a general virtual loss function for the single-player setting, which enabled effective tree parallelization of MCTS. Lastly, we introduced a policy training procedure for single-agent optimization tasks. The process uses a neural network to represent the policy and MCTS for policy improvement.

In our experiments on SameGame, we demonstrated the effectiveness of our policy training procedure, with the trained policy producing competitive results to state-of-the-art search algorithms on a public test set. Our results demonstrate a promising direction for future AI research in single-player optimization domains.

Another potentially fruitful research direction is to replace the policy network with a two-headed policy and value network. The predicted value can be combined with rollout results using a mixing parameter or replaced entirely. Our preliminary work on this subject has produced weaker results than plain MCTS. Whether it is beneficial to use value prediction for single-player optimization problems with unknown score bounds remains an open question. Lastly, we can look for other suitable search algorithms for policy improvement, as MCTS might not be the ideal choice.

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