How Should We Meta-Learn Reinforcement Learning Algorithms?

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Keywords: Meta-Reinforcement Learning, Algorithm Discovery.

Summary

The process of meta-learning algorithms from data, instead of relying on manual design, is growing in popularity as a paradigm for improving the performance of machine learning systems. Meta-learning shows particular promise for reinforcement learning (RL), where algorithms are often adapted from supervised or unsupervised learning despite their suboptimality for RL. However, until now there has been a severe lack of comparison between different meta-learning algorithms, such as using evolution to optimise over black-box functions or LLMs to propose code. In this paper, we carry out this empirical comparison of the different parts of the RL pipeline. In addition to meta-train and meta-test performance, we also investigate factors including the interpretability, sample cost and train time for each meta-learning algorithm. Based on these findings, we propose several guidelines for meta-learning new RL algorithms which will help ensure that future learned algorithms are as performant as possible.

Contribution(s)

1. We provide an empirical study of a number of different meta-learning algorithms when applied to a range of meta-learned algorithms for online reinforcement learning. This study considers the trade-off of meta-train and meta-test performance of learned algorithms in addition to how sample-efficient, time-consuming and interpretable different meta-learning algorithms are.

Context: Prior work has introduced a number of different meta-learning algorithms, such as using evolution to optimise black-box algorithms (Goldie et al., 2024; Lu et al., 2022), prompting LLMs to propose new functions (Lu et al., 2024), or using symbolic distillation of black-box functions to discover interpretable symbolic algorithms (Zheng et al., 2022). Due to the cost of meta-learning new algorithms, our study focuses on an intentionally diverse subset of meta-learning algorithms motivated by literature. Our study goes beyond any prior comparison of different meta-learning algorithms, improving our understanding of the field.

2. Based on our experimental results, we produce a set of recommendations for designing meta-learning pipelines. These proposals provide a 'snapshot' of the current state of the field, at the time of writing. These can help to make meta-learned algorithms as performant as possible within resource budgets.

Context: Meta-learning experiments are very time-consuming and costly. For instance, Goldie et al. (2024) uses over 2 GPU-years of compute for meta-learning optimisers in small-scale RL environments, and Metz et al. (2022b) use over 4000 TPU-months to meta-learn a large versatile optimisation algorithm. By providing rough guidelines for where researchers should start their search, this work can help to reduce redundant initial experimentation while hopefully improving downstream performance. Our recommendations are based on experimental results, which are rooted in algorithms available from current literature. Different meta-learning algorithms are likely to improve and mature into the future, and thus our recommendations may need adaptation down the line.

How Should We Meta-Learn Reinforcement Learning Algorithms?

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Abstract

The process of meta-learning algorithms from data, instead of relying on manual design, is growing in popularity as a paradigm for improving the performance of machine learning systems. Meta-learning shows particular promise for reinforcement learning (RL), where algorithms are often adapted from supervised or unsupervised learning despite their suboptimality for RL. However, until now there has been a severe lack of comparison between different meta-learning algorithms, such as using evolution to optimise over black-box functions or LLMs to propose code. In this paper, we carry out this empirical comparison of the different approaches when applied to a range of meta-learned algorithms, which each target different parts of the RL pipeline.¹ In addition to meta-train and meta-test performance, we also investigate factors including the interpretability, sample cost and train time for each meta-learning algorithm. Based on these findings, we propose several guidelines for meta-learning new RL algorithms which will help ensure that future learned algorithms are as performant as possible.

1 Introduction

The improvement of machine learning algorithms typically relies on manual design, a cumbersome process that is limited by human intuition and only rarely yields breakthroughs. An alternative, recent paradigm instead involves *meta-learning* learning algorithms from data. In this setting, algorithms are discovered computationally, with only limited need for human intervention in the design of the meta-learning process. This has particular potential for reinforcement learning (Sutton & Barto, 2020, RL), which is prone to instability (Van Hasselt et al., 2018; Achiam et al., 2019; Tang & Berseth, 2024) and often borrows algorithms from supervised and unsupervised learning that require adaptation to RL (e.g., (Parisotto et al., 2020; Obando Ceron et al., 2023; Ellis et al., 2024)).

There are numerous meta-learning algorithms, such as using evolution to optimise over neural networks for black-box algorithms, prompting a language model to propose algorithms in code or distilling from a pretrained black-box algorithm into a symbolic function. However, while many papers compare their meta-learned algorithms with handcrafted baselines, there have been few direct comparisons between methods for learning the algorithm itself. Consequently, there is little clarity on the pros and cons of different meta-learning algorithms, and to which settings they are most suited.

In this paper, we aim to address this deficit with an empirical analysis of different meta-learning algorithms. We consider a number of *meta-learned algorithms* – learned algorithms which replace certain components in RL training – and find the best *meta-learning algorithms* for each – ways for training the learned algorithm. This distinction is visualised in Figure 1, which is adapted from Goldie et al. (2024). We select meta-learned algorithms that exhibit different qualities, such as using recurrence or a large number of inputs, to provide coverage for a range of different possible algorithm features. These include learned optimisers and a learned drift function (Kuba et al., 2024).

¹We provide our code in an open-source library for meta-learning algorithms.

Our analysis focuses on the trade-offs between the different meta-learning algorithms. Principally, we consider the performance of each approach, both within its meta-training domain and in generalisation to new environments. In reinforcement learning, this is particularly important since algorithms often show limited ability to transfer (e.g., (Jackson et al., 2023)). In addition, due to the significant cost incurred by meta-learning experiments, which can require thousands of TPU-months of compute (Metz et al., 2022b), and the need for environment simulation in RL that is not present in supervised and unsupervised learning, we also consider the time and compute cost for training. Finally, we discuss the interpretability of the learned algorithms, which is useful for analysing the behaviour of an algorithm and its corresponding safety implications.

In our results, we find that: language models can find effective RL algorithms in a sample-efficient way, so long as there is a good algorithm from which to kickstart meta-training; distillation of learned algorithms into other networks sometimes improves performance without increasing samples; and symbolic representations do not scale well to recurrent algorithms or those with many inputs. Based on these findings, we propose several recommendations for better ways to meta-learn new RL algorithms, such as suggesting that many systems could benefit from using LLMs in the loop or that distillation from a black-box algorithm into another network is usually worth trying for a potential cheap performance boost. We hope that these guidelines can help reduce the cost of research in meta-RL while ensuring that meta-learned algorithms are as capable as possible.

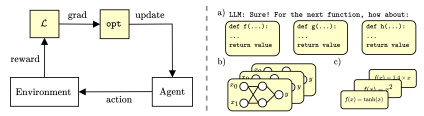


Figure 1: In the RL training loop on the left, we show different components of reinforcement learning which could be replaced by meta-learned algorithms. For example, OPEN (Goldie et al., 2024) is a learned optimiser replacing opt, while LPO (Lu et al., 2022) is a learned loss function which replaces \mathcal{L} . On the right, we demonstrate a few meta-learning algorithms, such as: a), prompting an LLM to propose new functions; b), evolving a black-box algorithm; or c), using symbolic evolution.

2 Related Work

2.1 Learned Algorithms

The practice of *meta-learning* algorithms is growing in popularity for both RL (Beck et al., 2024) and machine learning in general. There are many learned optimisation algorithms in supervised and unsupervised learning (e.g., (Andrychowicz et al., 2016; Metz et al., 2019b; 2020; Almeida et al., 2021)). Unlike these works, which present new meta-*learned* algorithms, we focus on understanding how the meta-*learning* algorithm affects a number of factors in RL, like generalisation. This is particularly important due to the instability of RL (Van Hasselt et al., 2018; Igl et al., 2021a) and the importance of transfer between environments (Finn et al., 2017; Duan et al., 2016; Jia et al., 2022).

Instead of meta-learning black-box algorithms represented by neural networks, some approaches discover *symbolic* algorithms defined as interpretable mathematical functions. Symbolic algorithms fit naturally into an LLM-based pipeline, since they are easily represented in code. Symbolic programs can be found through symbolic evolution (e.g., Lion (Chen et al., 2023)) or by prompting LLMs to improve algorithms over meta-training (e.g., (Lehman et al., 2022; Lu et al., 2024; Romera-Paredes et al., 2024)). In part of this work, we explore when symbolic algorithms are better than black-box ones, as suggested by Chen et al. (2023).

In RL, a pioneering meta-learned algorithm is Learned Policy Gradient (Oh et al., 2020, LPG), which replaces the actor-critic update, although there are many learned RL algorithms (e.g., (Kirsch et al., 2020; Jackson et al., 2023; Kirsch & Schmidhuber, 2022; Lan et al., 2024)). In addition to LPG, we focus on Learned Policy Optimisation (Lu et al., 2022, LPO), a learned alternative to proximal

policy optimisation (Schulman et al., 2017, PPO); and Optimisation for Plasticity, Exploration and Nonstationarity (Goldie et al., 2024, OPEN), a learned optimiser that uses feature engineering for meta-learning. Different to these papers, which propose new meta-learned algorithms for RL, we instead seek to understand how the meta-learning algorithm itself affects performance.

Generalisation after meta-training is important for learned algorithms to be applied in new settings. Jackson et al. (2023) explore using curricula based on unsupervised environment design (Dennis et al., 2021; Parker-Holder et al., 2022) for meta-training as a way to improve LPG generalisation. In this work, we consider how different meta-learning algorithms affect generalisation. As a separate component of the meta-training process, our study is complementary to that of Jackson et al. (2023).

2.2 Distillation

Distillation, which trains a *student* to imitate a *teacher* (Hinton et al., 2015), relates to many metalearning algorithms. Distillation is often applied to policies (Rusu et al., 2016; Jia et al., 2022), datasets (Wang et al., 2020; Lupu et al., 2024), handcrafted algorithms (Laskin et al., 2023; Son et al., 2025), and reasoning language models (DeepSeek-AI et al., 2025). Distillation from one network to another, called black-box distillation, usually trains a student that is *smaller* than its teacher (Hinton et al., 2015), to reduce inference costs and overfitting, or the *same size* as the teacher (Furlanello et al., 2018), since distillation *itself* acts as a regulariser (Zhang & Sabuncu, 2020; Mobahi et al., 2020). Contrary to these papers, our analysis explores whether applying black-box distillation to learned algorithms provides similar benefits as in other settings.

Rather than distilling from one network to another, symbolic distillation learns a *symbolic program* (Cranmer et al., 2020) that has a similar mapping to the neural network teacher. Symbolic distillation is often applied to physical systems (e.g., (Cranmer et al., 2020; Mengel et al., 2023; Lemos et al., 2023)) for interpretability reasons, but has been extended to learned optimisers (Zheng et al., 2022; Song et al., 2024a). Similarly, Lu et al. (2022) manually distil LPO, a black-box algorithm, into *discovered* policy optimisation. In this paper, we seek to understand *when* symbolic distillation is appropriate for meta-learned RL algorithms. While interpretability is part of our analysis, we also consider whether symbolic distillation improves generalisation of learned algorithms.

3 Background

Reinforcement Learning Reinforcement learning (RL) problems are often modeled as Markov decision processes (Sutton & Barto, 2020, MDPs). An MDP is typically denoted as a tuple $\langle \mathcal{A}, \mathcal{S}, \mathcal{S}_0, P, \rho, R, \gamma \rangle$. In an MDP, an agent in state $s_t \in \mathcal{S}$, starting at $s_0 \in \mathcal{S}_0$, takes an action $a_t \in \mathcal{A}$ according to its state-conditioned, probabilistic policy $\pi(\cdot|s_t)$ and the state transitions to s_{t+1} based on the environment transition dynamics $P(\cdot|s_t, a_t)$. In response, the environment generates a reward $r = R(s_t, a_t)$. An agent's policy is trained to maximise its expected discounted return, $J^{\pi} = \mathbb{E}_{a_{0:\infty},s_0 \sim \rho, s_{1:\infty} \sim P} [\sum_{t=0}^{\infty} \gamma^t R_t]$, with a discount factor of $\gamma \in [0, 1)$.

Mirror Learning Mirror learning (Kuba et al., 2024) is a theoretical framework that provides guarantees to a class of RL algorithms including PPO (Schulman et al., 2017) and underpins the architecture of LPO (Lu et al., 2022). A mirror learning algorithm updates a policy according to

$$\pi_{k+1} = \underset{\pi \sim \mathcal{N}(\pi_k)}{\arg \max} \mathbb{E}_{s \sim \beta_{\pi_k}} [A_{\pi_k}(s, a)] - \mathbb{E}_{s \sim \nu_{\pi_k}} [\mathcal{D}_{\pi_k}(\pi|s)], \tag{1}$$

where β_{π_k} and $\nu_{\pi_k}^{\pi}$ are sampling and drift distributions over s, and A(s, a) = Q(s, a) - V(s) is the advantage. \mathcal{D} , the *drift* function, measures the difference between π and the current policy π_k and is used to penalise large policy updates. A valid drift function must uphold three conditions: be nonnegative everywhere; be zero at $\pi = \pi_k$; and have zero gradient with respect to π when $\pi = \pi_k$. For PPO, the drift function is ReLU $\left(\left[\frac{\pi(a|s)}{\pi_k(a|s)} - \operatorname{clip}\left(\frac{\pi(a|s)}{\pi_k(a|s)}, 1 \pm \epsilon\right)\right] A_{\pi_k}(s, a)\right)$.

4 Meta-Learning Algorithms

In this section, we describe different meta-learning algorithms and qualitatively discuss the possible pros and cons of each.

4.1 Black-Box Meta-Learning

Black-box algorithms are typically represented as neural networks. For example, a black-box learned optimiser might replace gradient descent with a neural network that maps from gradient to a parameter update. Most black-box algorithms are meta-trained using evolution or meta-gradients.

Meta-gradients are often calculated with backpropagation through time (BPTT) with respect to an RL objective, where the algorithm itself is treated as an agent (Oh et al., 2020) and updates are applied after fixed-length rollouts of the algorithm. Rollouts are usually truncated to prevent exploding or vanishing gradients, causing bias (Metz et al., 2022a; Wu et al., 2018). Although Jackson et al. (2024) demonstrate that evolution often learns better algorithms than meta-gradients, to provide diversity in our study, we use meta-gradients for learning LPG as proposed by Oh et al. (2020) and an evolutionary algorithm, evolution strategies (Wierstra et al., 2011; Salimans et al., 2017; Rechenberg, 1973, ES), for other algorithms.

ES is a population-based optimisation approach where a network's parameters, $\hat{\theta}$, are iteratively updated using a natural gradient estimate for fitness $F(\cdot)$. This is calculated as $\nabla_{\theta} \mathbb{E}_{\epsilon \sim \mathcal{N}(0,I)} F(\theta + \sigma \epsilon) = \frac{1}{\sigma} \mathbb{E}_{\epsilon \sim \mathcal{N}(0,I)} \{\epsilon F(\theta + \sigma \epsilon)\}$ using a sample average for N sets of randomised parameters, with mean $\tilde{\theta}$. $\tilde{\theta}$ is updated with gradient ascent to maximise $F(\cdot)$, which is often defined as an agent's final return (Lu et al., 2022; Goldie et al., 2024). Unlike symbolic evolution, which is difficult to vectorise due to each program having a separate computation trace (see Section 4.3), the structure of ES can easily exploit GPU vectorisation for parallelisation (Lu et al., 2022; Lange, 2022b).

4.2 Black-Box Distillation

One way to improve the performance of black-box algorithms may be to distil the algorithm into another neural network, as introduced in Section 2.2. In our analysis, we consider two types of black-box distillation: distilling into a network with the *same* architecture (**Same-Size Distillation**); or distilling into a *smaller* network (**Smaller Distillation**), which we implement by halving all layer widths, such that the student underfits the teacher. Underfitting may help generalisation if the teacher has overfit to its original meta-training distribution, and distillation can itself provide learning regularisation (Zhang & Sabuncu, 2020; Mobahi et al., 2020).

We distil by using L_2 regression to match the student's outputs to the teacher for synthetically generated inputs, rather than sampling from the environment. This needs no additional environment transitions and introduces more diversity than sampling from RL training, which can lead to skewed distributions. We evaluate the RL performance of the student on the original meta-training environments periodically, and select the best-performing checkpoint as the distilled algorithm.

4.3 Symbolic Discovery

Evolutionary algorithms combine mutation, which randomly changes members of a population; crossover, which mixes two members of the population; and selection, which selects individuals from a population to pass to the next generation. When evolving an abstract syntax tree (AST), which represents a symbolic program, mutation adds or changes variables, constants or functions in the tree nodes and crossover swaps the nodes, and sometimes their children, between two ASTs.

Lion (Chen et al., 2023) is an interpretable symbolic learned optimiser discovered using symbolic evolution. However, symbolic search is inefficient and requires evaluating many functions (30,000 for a single seed in Lion, despite warm-starting from handcrafted optimisers) and, while computing fitness is quick for supervised learning, RL typically requires expensive environment simulation.

Though ES can be parallelised using a GPU, since the traced computation graphs of members in the population are the same, vectorising symbolic programs is more difficult as different programs have different computation graphs. Complex hand-coded branching logic could overcome this issue, but this would lead to huge performance degradation and significant inefficiency. Therefore, we exclude direct symbolic discovery from our empirical analysis and cover it here only for completeness.

4.4 Symbolic Distillation

Rather than evaluating symbolic programs in RL, a quicker alternative is to distil black-box algorithms into symbolic programs in a supervised problem. In addition to outputting interpretable functions, this approach may lead to better generalisation (Zheng et al., 2022; Chen et al., 2023). We base our approach on Zheng et al. (2022), who distil a learned optimiser into a symbolic program. We generate input data using the same statistics as in black-box distillation, albeit generated in one large batch to make the dataset stationary. We apply symbolic evolution (Section 4.3) using PySR (Cranmer, 2023) to find a program with low L_2 loss with respect to the black-box teacher outputs. While PySR has an in-built method for selecting algorithms based on a *combination* of high fitness and low complexity, we consistently find that choosing the most fit (i.e., lowest L_2 loss) function produces better RL results. As such, we select the algorithm that most accurately fits the teacher.

4.5 LLM Proposal

Since the rise of highly capable agentic language models, many researchers have used language models for algorithm discovery (e.g., (Lu et al., 2024; Faldor et al., 2024; Romera-Paredes et al., 2024; Hu et al., 2024; Song et al., 2024b)). Generally, this research is based on the premise that language models generate intelligent proposals, making them more sample efficient than symbolic evolution. As such, LLM-driven discovery pipelines generally evaluate on the order of tens of algorithms, rather than thousands, making them much more practical for evaluating directly in RL.

Since prompt tuning can play a large part in LLM performance, we build on an existing system, DiscoPOP (Lu et al., 2024), and warm-start search from a handcrafted algorithm. The LLM must reason in-context about previous algorithm performance to make suggestions for the next algorithm. In our setting, due to a number of unconventional inputs (particularly in the case of OPEN), we provide the LLM with a brief description of all inputs to the learned algorithm. After training, we select the best in-distribution algorithm for evaluation. We use GPT o3-mini (OpenAI, 2025) as our LLM, since it is a highly capable reasoning model with good performance for coding tasks.

5 Meta-Learned Algorithms

In this section, we introduce the set of meta-learned algorithms to which we apply the meta-learning algorithms introduced in Section 4. Due to the cost of meta-learning experiments, we are both selective and deliberate about which algorithms to include. We choose algorithms based on: how many inputs they condition on; whether they are recursive or not; and the component of RL training which they affect.

Learned Policy Optimisation LPO (Lu et al., 2022) is a learned algorithm that replaces the mirror drift function in PPO (Schulman et al., 2017; Kuba et al., 2024). The LPO network has no bias, to satisfy the mirror learning conditions at $r := \frac{\pi}{\pi_k} = 1$, and passes through a ReLU for nonnegativity. Inputs to LPO are transformations of r, the policy ratio, and A, the advantage typically calculated with generalised advantage estimation (GAE) (Schulman et al., 2018), and are defined as

$$\mathbf{x} = \left[(1-r), (1-r)^2, (1-r)A, (1-r)^2 A, \log(r), \log(r)A, \log(r)^2 A \right].$$
(2)

We follow Lu et al. (2022) in initialising LPO near the PPO drift function to ease meta-learning.

Learned Policy Gradient LPG (Oh et al., 2020) meta-learns a policy update rule based on actorcritic algorithms (Sutton & Barto, 2020), which update a policy (actor) using a learned value (critic). It is typically trained with meta-gradients (Oh et al., 2020; Jackson et al., 2024) and takes inputs of

$$[r_t, d_t, \gamma, \pi(a_t|s_t), y_\theta(s_t), y_\theta(s_{t+1})], \qquad (3)$$

from fixed-length policy rollouts, using a backward-LSTM (Hochreiter & Schmidhuber, 1997). Here, r_t is a reward, d_t is a done flag, γ is a discount factor, $\pi(a_t|s_t)$ is the probability of taking action a_t in s_t and $y_{\theta}(\cdot)$ is an *n*-dimensional categorical distribution acting as a bootstrap vector. **Optimisation for Plasticity Loss, Exploration, and Non-stationarity** OPEN (Goldie et al., 2024) is a meta-learned optimiser for RL that conditions on features measuring the presence of certain difficulties in RL optimisation, in addition to typical learned optimiser inputs (Metz et al., 2020). Its design takes into account: plasticity loss (Abbas et al., 2023; Lyle et al., 2023; Dohare et al., 2024), where an agent loses the ability to learn new things, which is overcome by conditioning OPEN on neuron dormancy (Sokar et al., 2023) and allowing it to behave differently on deeper layers in the agent; exploration (Cesa-Bianchi et al., 2017; Burda et al., 2018; Aubret et al., 2023; Sukhija et al., 2025), which prevents agents from getting trapped in local minima and which is boosted in OPEN by making the update slightly stochastic, as in noisy nets (Fortunato et al., 2019) or parameter space noise (Plappert et al., 2018); and non-stationarity (Igl et al., 2021b), which is measured based on how long training has gone on (like (Jackson et al., 2024)) and how many iterations have been spent optimising with a given data batch (similar to Ellis et al. (2024)). The full set of inputs are

$$\mathbf{x} = [p, g, m_{0.1}, m_{0.5}, m_{0.9}, m_{0.99}, m_{0.999}, m_{0.9999}, t_p, b_p, \text{dorm}, l_p],$$
(4)

where p is the current parameter, g is its gradient with respect to the PPO objective and m_x is an exponential moving average of gradient with scale x. g and m_x are both transformed as $x \to \{\log(|x|+\epsilon), \operatorname{sgn}(x)\}$ to ease learning (Lan et al., 2024). The extra inputs are t_p and b_p , which measure time on the training and batch scale; dorm, which is the dormancy of the neuron downstream of the parameter; and l_p measures the depth of a parameter. The optimiser is applied to each parameter in a network independently. Whereas OPEN originally uses a *recurrent* architecture, here we explore applying different meta-learning algorithms to *both* a feed-forward and recurrent OPEN.

No Features As in Goldie et al. (2024), we consider a 'No Features' learned optimiser that is similar to OPEN but includes only parameter, gradient, and momentum information. We include it as an example of a simple learned optimiser and hence only consider a feed-forward version of it.

6 Evaluation

There is no single measure of success for meta-learning algorithms. For instance, some users may choose to sacrifice some return for the sake of interpretability. Therefore, when comparing the different meta-learning algorithms, we consider a number of performance measures. In Section 8, we propose a number of design recommendations for future meta-learned algorithms with the following qualities in mind:

- In-distribution (i.d.) return, where we evaluate the algorithm on its meta-training task or tasks;
- Out-of-distribution (o.o.d.) return, where the algorithm is evaluated for meta-*test* generalisation to environments outside its training distribution;
- The sample cost of meta-learning, where training is stopped at peak in-distribution performance;
- The meta-train runtime (wall clock) for learning the algorithm;
- The meta-test runtime (wall clock); and
- How interpretable the algorithm is, judged subjectively as low, medium, or high.

For feed-forward algorithms, we meta-learn from both a *single* environment, Ant from Brax (Freeman et al., 2021; Todorov et al., 2012), and the *multiple* environments in MinAtar (Lange, 2022a; Young & Tian, 2019), following Goldie et al. (2024). These settings are selected to enable *fast* metalearning without having overlap between the different meta-training distributions. For the recurrent implementation of OPEN, we use a pretrained optimiser from Goldie et al. (2024) instead of metatraining one ourselves, to allow for comparison against a publicly available baseline. Here, we focus only on the multiple environment setting to limit the cost of distillation, which is more expensive for recurrent algorithms. We meta-test these algorithms on a diverse set of environments: Freeway, Space Invaders, Asterix and Breakout from MinAtar (Lange, 2022a; Young & Tian, 2019)²; Humanoid, Hopper, Walker and Ant from Brax (Freeman et al., 2021; Todorov et al., 2012); Cartpole

²Seaquest is not available in the Gymnax implementation of MinAtar.

from OpenAI gym (Lange, 2022a; Brockman et al., 2016); and Craftax-Classic (Matthews et al., 2024; Hafner, 2021). For LPG, to align to prior research, we follow Oh et al. (2020) by metatraining on randomly distributed gridworlds and, as in Jackson et al. (2023), explore transfer to MinAtar. We specify all hyperparameters in Supplementary Material A, including any hyperparameters needed for the LLM proposed functions, which are tuned for the warm-start algorithm in each environment separately, before meta-training. Instead of a standardised evaluation set, we believe that our approach is more informative for the *actual* use cases of each meta-learned algorithm.

Due to the high cost and chance of failure, we do not apply symbolic distillation to recurrent algorithms. While Zheng et al. (2022) distil a recurrent learned optimiser with a single input using a fixed window of inputs, LPG has 19 inputs and OPEN has 20. For a window size of 20, as in Zheng et al. (2022), we would therefore require over 380 symbolic variables. Such a high dimensional problem is extremely difficult for symbolic regression to solve and would require so many AST nodes as to be computationally infeasible, given the search space grows exponentially with the size of the tree.

When plotting results, we normalise returns for each environment independently by dividing by the mean black-box learning score. Results are aggregated into 'In' and 'Out Of' Distribution based on the meta-training distribution and, unless otherwise stated, show the interquartile-mean (IQM) of return with 95% stratified bootstrap confident intervals for 16 environment seeds, following Agarwal et al. (2021). In addition to understanding how well each method performs in- and out-of-distribution, our in-distribution results for distillation verify whether it was successful.

We include unnormalised and unaggregated results in Supplementary B, and the symbolic and LLMproposed algorithms in Supplementary C. We show all initial LLM prompts in Supplementary D, and an example LLM discussion is in Supplementary E. We also provide extra results, from metatraining in gridworlds, in Supplementary F.

Due to the high cost of meta-learning, we follow standard procedure from the literature by metalearning each algorithm for a single seed (Goldie et al., 2024; Metz et al., 2022b; Lan et al., 2024; Metz et al., 2019a) without meta-hyperparameter tuning.

7 Results

In this section, we present results from all experiments introduced in Section 6.

7.1 Learned Policy Optimisation

We firstly consider LPO, with results shown in Figure 2. We find that all distillation examples perform similarly, and often give minor generalisation gains without harming i.d. performance. Even though the LLM-proposed algorithms perform significantly worse than the others i.d., they achieve the best o.o.d. performance. This is unsurprising: the LLM proposed algorithms in Supplementary C for each task are both very similar and related to the warm-start function, PPO, and so are expected to generalise across a wide task distribution. Based on these results, LLM proposal is the best approach if generalisation is the priority. For an algorithm which performs well both i.d. and o.o.d., same-size distillation of a black-box algorithm is possibly the best option.

We visualise the gradients of all LPO functions in Supplementary G, following Lu et al. (2022).

7.2 Feed-Forward No Features

We show performance for the No Features optimiser in Figure 3. In Ant, the black-box optimisers fail to learn; based on Goldie et al. (2024), No Features is a very weak learned algorithm for RL, making this failure unsurprising. This does highlight a clear limitation of distillation, though: if the original algorithm is poor, distillation is unlikely to *fix* it. Symbolic distillation also struggles, likely as the 8 inputs make this a relatively high dimensional problem for symbolic evolution. Overall, LLM proposal is by far the strongest baseline, both in-distribution and for generalisation.

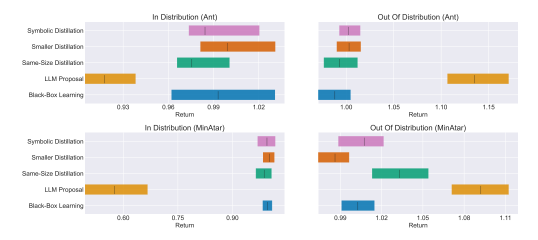


Figure 2: IQM of final returns for LPO trained on Ant (top) and MinAtar (bottom). The distillation experiments all have similar returns, with same-size distillation offering improved generalisation over the algroithm learned in MinAtar. LLM Proposal is poor in-distribution, but has strong out-of-distribution generalisation performance.

The LLM likely performs well for a few reasons: gradient-based optimisation is well covered in the LLM's training corpus; all inputs to the optimiser are easy to understand; and the LLM has access to a per-environment learning rate tuned for its initialisation of SGD, which effectively relies on few-shot meta-test evaluation. The use of hyperparameters can be seen as an advantage, for flexibility, or disadvantage, if meta-test time samples are expensive.

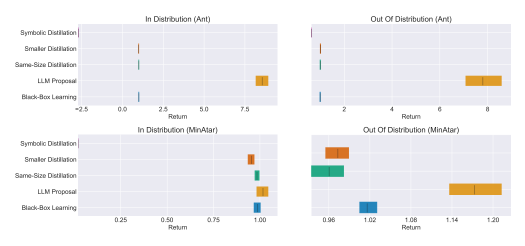


Figure 3: IQM of final returns for the No Features optimiser after meta-training in Ant (top) and MinAtar (bottom). Black-Box Learning struggles to learn in Ant, hurting the distilled optimiser performances. In MinAtar, symbolic distillation produced an optimiser which produces NaN returns in generalisation, so is omitted from the plot.

7.3 Feed-Forward OPEN

In Figure 4, we show the performance of a *feed-forward* implementation of OPEN after metatraining in Ant and MinAtar. OPEN has more inputs than the other algorithms analysed thus far, which is likely why the LLM and symbolic distillation catastrophically fail. Anecdotally, we find that symbolic distillation is unable to search the high dimensional space and instead converges to relatively simple, almost constant, algorithms, and the language model is unable to correctly use the additional input features despite explanations of their meaning. In fact, despite giving the LLM the shapes and ranges of all inputs, many of the algorithms it proposes in training produce errors. Similar to No Features, distilling into a smaller model can worsen performance. However, samesize distillation produces a small generalisation benefit for the model trained on MinAtar. It is likely that the smaller model's representational capacity is too low, but that the regularisation effect of same-size distillation aids generalisation.

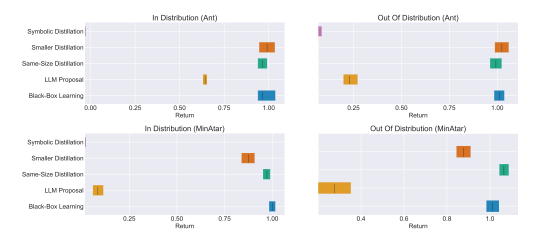


Figure 4: IQM of final returns for meta-training feed-forward OPEN in Ant (top) and MinAtar (bottom). Symbolic distillation from meta-learning in MinAtar caused NaNs out of distribution, so is omitted from the plot. The LLM and symbolic distillation both clearly struggle.

7.4 Recurrent LPG

In Figure 5, we explore the generalisation performance of meta-training LPG in gridworlds for black-box learning and distillation only. Due to the formulation in LPG of y_{θ} as a categorical distribution, finding an algorithm grounded in literature to warm-start LLM proposals, as needed in DiscoPOP, is impractical. Therefore, as well as excluding symbolic distillation of LPG as a recurrent algorithm, we omit LLM Proposal and underscore this as a

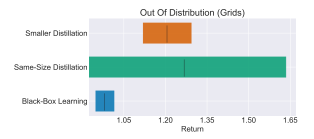


Figure 5: O.o.d. performance of LPG after training in gridworlds and transferring to MinAtar. Both types of distillation lead to higher mean performance, but wider confidence intervals.

key limitation of LLM proposal: it needs *something* to start from, which may not always be practical.

In these results, distillation leads to improved IQM generalisation performance both when the student is smaller *and* the same size, albeit with overlapping confidence intervals when the student is the same size. Given that LPG uses a large network (\sim 200k parameters, compared to \sim 1k for OPEN), the regularisation from distillation likely helps reduce variance, improving generalisation.

7.5 Recurrent OPEN

Unlike LPG, which rolls out for only 20 steps at a time, OPEN unrolls over the *entire* course of RL training, which can potentially be tens of thousands of steps. As such, for stability and computational reasons, we cannot distil from data sequences as long as RL training. Instead, we distil a pretrained OPEN optimiser over 'Long Rollouts', where we train on sequences of 100 steps, and 'Short Rollouts', where the generated sequences are 20 steps long.

Figure 6 shows that distillation of recurrent OPEN is poor, suggesting that distilling an algorithm with long unrolls is too hard. This contrasts with feed-forward OPEN, where distillation occasionally helps and rarely hurts performance. LLM proposal, which was initialised with Adam (a better optimiser than SGD, which initialises feed-forward LLM-proposed optimisers), produces a stronger optimiser in o.o.d. environments than black-box learning. This is likely due to the fact that the best LLM algorithm is *very* similar to Adam, having been discovered early in training. It also uses a per-environment learning rate tuned for Adam and only uses extra features to have a per-layer learning rate; later attempts to incorporate more features lead to significantly worsened performance. Over-all, the black-box learning algorithm in this setting learns a performant but overfit algorithm and the LLM a simple but more general optimiser, although it does not change much from its initialisation.

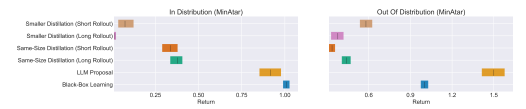


Figure 6: IQM of final returns for recurrent OPEN meta-trained in MinAtar. Distillation struggles due to the long unroll length of OPEN. The LLM performs well, but requires a tuned learning rate.

7.6 Additional Metrics

In this section we provide a more holistic evaluation of the different meta-learning algorithms, which is summarised in Table 1.

Black-box learning incurs a high sample cost since it requires many iterations of learning in an online environment, but distilling from this black-box algorithm requires no additional samples as distillation uses synthetic data. Since LLM proposal only evaluates individual algorithms produced by the language model, it requires comparatively few online interactions and is thus sample efficient.

In terms of speed, symbolic distillation can be the slowest of all techniques since its meta-training time scales exponentially with the maximum number of nodes in the AST, although it can be quicker for simpler algorithms. This contrasts with black-box distillation methods, whose speed remains broadly similar no matter the function being distilled. Using an LLM is fast, both because the search is warm-started from a known algorithm and as it only requires evaluating a small number of high quality algorithms, unlike the more random search in symbolic evolution.

Whereas black-box algorithms are almost completely uninterpretable, symbolic distillation and especially the LLM produce highly interpretable algorithms. This disparity arises because, whereas the LLM explains its proposals in plain-text at generation time, symbolic distillation generally introduces many constants into the equations that can obfuscate behaviour.

We find that symbolic distillation is unable to scale to functions with more than a small number of inputs. While LLM proposal is better, as it makes intelligent suggestions rather than randomly searching, we find that it is unable to incorporate all features from OPEN into a performant algorithm and requires warm-starting. Therefore, as the only meta-learning algorithm which can meta-train on long rollouts with many features, we believe black-box learning is the most scalable algorithm.

Approach	Samples	Train Time	Test Time	Interpretability	Scalability
Black-Box Learning	High	Slow	Slow	Bad	Best
Same-Size Distillation	No Extra	Slow	Slow	Bad	Good
Smaller Distillation	No Extra	Medium	Medium	Bad	Good
Symbolic Distillation	No Extra	Medium-Slow	Fast	Medium	Bad
LLM Proposal	Low	Fast	Fast	Good	Medium

Table 1: A summary of how each meta-learning algorithm performs across different metrics.

8 Design Recommendations

Based on the results in Section 7, we produce a set of design recommendations for future metalearning pipelines. These recommendations reflect the current state of the field, meaning they may require adaptation as meta-learning algorithms and capabilities improve. We describe them below.

- For a meta-learned algorithm with *few* inputs, or inputs which are easy to understand (i.e., an LLM can interpret them), prompting an LLM for new algorithms is a sample-efficient way to find new generalisable algorithms. This has three caveats: there must be an easy-to-define, performant function from which to start the search; it must be possible to run hyperparameter tuning for the algorithm in the meta-test environment; and in-distribution performance of the algorithm will likely be worse than learning a black-box function (especially for many meta-samples).
- As long as it is possible to define a warm-start initialisation function, it is almost always better to prompt a language model for algorithm proposals over applying symbolic distillation. In fact, besides yielding interpretable functions, symbolic distillation is unlikely to improve performance, contrary to the suggestion of Chen et al. (2023) that symbolic functions should generalise better.
- Black-box distillation can often, but not always, improve generalisation. We recommend applying black-box distillation into the same-sized network for all black-box learned algorithms that are feed-forward or have short recurrent rollouts; given there is no increased sample cost and training is quick, this can occasionally yield cheap performance gains. On balance, smaller distillation can cause bigger drops in performance for smaller potential gains.
- Black-box algorithms are practically the only way to meta-learn algorithms which use a large number of features. If a meta-learned algorithm has many inputs, like OPEN, then an LLM is unlikely to propose a performant algorithm which also incorporates all of the input features.

9 Limitations and Future Work

There are a number of possible directions for future work. Firstly, while we discuss the reliance of LLM-proposed algorithms on hyperparameters, it would be interesting to explore *how* reliant the algorithms are on hyperparameter selection, and whether they are more sensitive than handcrafted algorithms, based on approaches like Probst et al. (2019) and Adkins et al. (2025).

Secondly, an unexplored axis in our study is how the *representation* in black-box distillation affects performance. For instance, while we consider changing the black-box layer widths, we do not explore the effect of changing architectures entirely on performance. Inspired by work in algorithm distillation (Laskin et al., 2023; Son et al., 2025), it could be insightful to test distillation from recurrent or feed-forward algorithms to transformers (Vaswani et al., 2023).

Finally, we believe the findings presented here could be built upon by blending different meta learning algorithms. For instance, one avenue could test whether symbolic distillation scales better to high-dimensional problems if inputs were first encoded by a black-box network, or whether LLMs could be warm-started from a symbolically distilled algorithm. Similarly, understanding the effect of different prompting styles or agentic frameworks would be a valuable addendum to this work.

10 Conclusion

This work presents a large-scale empirical analysis comparing many different meta-learning algorithms for RL: learning a black-box algorithm; distilling the algorithm into a same-size or smaller network; distilling the algorithm into a symbolic function; or prompting a language model to propose new algorithms. Based on our results, we propose a number of recommendations for how to meta-learn algorithms for RL. These include generally using language models for discovering new algorithms, so long as search can be initialised from something performant and it is possible to tune hyperparameters, and trying same-sized black-box distillation to potentially improve generalisation. These design suggestions can be used to ensure learned algorithms are as performant as possible for RL, while hopefully reducing the need for unnecessary experiments.

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Supplementary Materials

The following content was not necessarily subject to peer review.

A Hyperparameters

In this section, we provide all hyperparameters used in this paper. Due to the number of experiments used here, we break our list of hyperparameters into multiple subsections. To prevent unnecessary hyperparameter tuning, where our implementations are based in open-source releases from other works we replicate their hyperparameters.

A.1 Policy Optimisation Hyperparameters

The learned optimiser experiments (i.e. OPEN, No Feat) use PPO for policy optimisation, and use the same hyperparameters as the LPO experiments. Our PPO hyperparameters are largely grounded in Lu et al. (2022) and Goldie et al. (2024), and are show in Table 2.

Note that, for LPO, only the LLM proposals get access to *PPO Clip* ϵ as it needs to be initialised at PPO.

Table 2: PPO and LPO hyperparameters. The Brax and MinAtar suites used common PPO parameters.

II	Environment			
Hyperparameter	MinAtar	Brax	Cartpole	Craftax
Number of Environments N _{envs}	64	2048	4	1024
Number of Environment Steps N_{steps}	128	10	128	20
Total Timesteps T	1e7	3e7	5e5	3e7
Number of Minibatches $N_{minibatch}$	8	32	4	16
Number of Epochs L	4	4	4	2
Discount Factor γ	0.99	0.99	0.99	0.99
$G\!A\!E\lambda$	0.95	0.95	0.95	0.95
PPO Clip ϵ	0.2	0.2	0.2	0.2
Value Function Coefficient c_1	0.5	0.5	0.5	0.5
Entropy Coefficient c_2	0.01	0.0	0.01	0.01
Max Gradient Norm	0.5	0.5	0.5	0.5
Layer Width W	64	64	64	64
Number of Hidden Layers H	2	2	2	2
Activation	relu	tanh	tanh	tanh

LPG uses a different set of hyperparameters since it has a different algorithmic backbone. We use hyperparameters from Jackson et al. (2023). We use the 'all_shortlife' class of gridworlds for meta-training, and show the LPG hyperparameters in Table 3.

A.2 Optimiser Hyperparameters

Depending on whether the algorithm was feed-forward or recurrent, the learned optimisers require a per-environment learning rate tuned for either SGD or Adam. We provide all optimiser hyperparameters for LPO and PPO with learned optimisers in Tables 4-6, and for LPG in Table 7. For LPO and PPO, we tune optimiser hyperparameters *individually* per environment. We round all values for SGD. LPO uses a slightly different learning rate than the learned optimisers in some cases, since we used standard β values of [$\beta_1 = 0.9, \beta_2 = 0.999$] for the learned optimisers but tuned them for LPO, as they are not part of the learned algorithm. All learning rates use linear annealing over the course of training.

Uynomonomoton	Environment		
Hyperparameter	Gridworld	MinAtar	
Number of Environments N_{envs}	64	64	
Number of Environment Steps N_{steps}	20	20	
Total Timesteps T	3e6	1e7	
Number of Minibatches $N_{minibatch}$	64	64	
Discount Factor γ	0.99	0.99	
$G\!A\!E\lambda$	0.95	0.95	
Entropy Coefficient c_2	0.01	0.01	
Max Gradient Norm	0.5	1.0	
Layer Width W	32	32 (conv)	
Number of Hidden Layers H	1	2	
Activation	relu	relu	

Table 3: Hyperparameters for	policy	optimisation	and the agent in	LPG experiments.
2 1 1		1	U	1

Table 4:	PPO and LPO	hyperparameters	for MinAtar	environments.

Hunounovomotov	Environment				
Hyperparameter	Asterix	Breakout	Freeway	SpaceInvaders	
LPO Learning Rate	3e-3	1e-2	1e-3	7e-3	
eta_1	0.9	0.9	0.9	0.9	
β_2	0.999	0.99	0.99	0.99	
SGD Learning Rate	0.52	1.02	0.56	1.17	
L2O Adam Learning Rate	3e-3	7e-3	1e-3	3e-3	

Table 5: PPO and LPO hyperparameters for Brax environments
--

Umanananatan	Environment				
Hyperparameter	Ant	Humanoid	Walker	Hopper	
LPO Learning Rate	3e-4	3e-4	1e-3	$8e{-4}$	
β_1	0.99	0.9	0.9	0.9	
β_2	0.99	0.999	0.999	0.999	
SGD Learning Rate	0.17	0.053	0.52	0.27	
L2O Adam Learning Rate	$3e{-4}$	3e-4	1e-3	8e-4	

Table 6: PPO and LPO hyperparameters for Car	tpole and Craftax.
--	--------------------

Uunonnonomoton	Environment		
Hyperparameter	Cartpole	Craftax	
LPO Learning Rate	1e-3	5e-4	
β_1	0.9	0.9	
eta_2	0.999	0.999	
SGD Learning Rate	$2.5e{-4}$	0.46	
L2O Adam Learning Rate	3e-3	5e-4	

Table 7: LPG optimiser hyperparameters.

II	Enviro	nment
Hyperparameter	Gridworld	MinAtar
Learning Rate	1e-3	5e-4

A.3 Meta-Learning Hyperparameters

In tables 8 and 9 we provide all necessary hyperparameters for *meta-learning*. In table 10, we include hyperparameters for symbolic distillation. We do not tune hyperparameters for black-box learning due to the computational cost of meta-learning. We run a small sweep over learning rates for black-box distillation. For symbolic distillation, we mostly follow the implementations in Cranmer (2023), albeit using a custom set of possible symbolic functions and generally allowing more complex programs.

For distillation, we sweep over learning rates in [0.1, 0.02, 0.001]. For smaller distillation, we halve all layer widths.

		Meta-Learned Algorithm				
Hyperparameter	LPO	No Features	Feed-Forward OPEN	Recurrent OPEN		
ES Learning Rate	3e-2	3e-2	3e-2	-		
ES LR Decay	0.999	0.999	0.999	-		
$\mathit{ES}\sigma_{init}$	3e-2	3e-2	3e-2	-		
$\mathit{ES}\sigma_{decay}$	0.999	0.999	0.999	_		
Population Size	64	64	64	-		
Number Dense Layers	1	2	2	2		
GRU Size (MinAtar)	_	_	-	16		
Dense Layer Size (MinAtar)	128	32	32	32		
GRU Size (Ant)	_	-	-	8		
Dense Layer Size (Ant)	128	16	16	16		
Dense Layer Size (Ant)	128	10	10	10		

Table 8: Meta-learning hyperparameters for LPO and learned optimisers.

Table 9: Meta-learning Hyperparameters for LPG, following Jackson et al. (2023)

Hyperparameter	Environment Gridworld
Num Steps	5000
Embedding Width	16
GRU width	256
Target Width	8
Agent Target KL Divergence	0.5
Learning Rate	1e-4
LPG Max Grad Norm	0.55
Num Agent Updates	5
LPG Policy Entropy Coeff	5e-2
LPG Target Entropy Coeff	1e-3
LPG Policy L_2 Coeff	5e-3
LPG Target L ₂ Coeff	1e-3

Hyperparameter	Environment	
	LPO	Feed-Forward OPEN/No Features
Max Size	40	60
Populations	31	160
Number Iterations	10	10
Batch Size	5000	5000
Weight Optimise	0.001	0.001

Table 10: Symbolic Distillation Hyperparameters, following Cranmer (2023). We use warm-starting after every RL evaluation of the best fit algorithm; as such, while the PySR 'Number Iterations' is 10, we loop over this process 40 times (effectively leading to 400 iterations).

B Returns by Environment

In this section, we include plots of the returns achieved by all learned algorithms in all of the environments we test. Unlike in the main body of the paper, we do not aggregate any results here.

All algorithms were run for 16 environment seeds. We plot IQM with 95% stratified bootstrap confidence intervals, following (Agarwal et al., 2021). For clarity, we separate the returns into two rows; each pair of rows corresponds to a single trained algorithm.

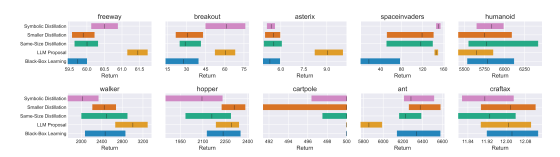


Figure 7: IQM of final meta-test returns after meta-training LPO in Ant.

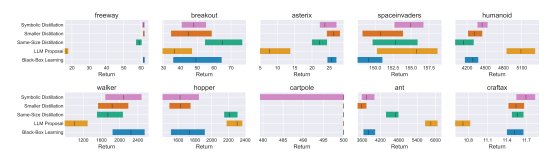


Figure 8: IQM of final meta-test returns for LPO meta-trained in MinAtar.



Figure 9: IQM of final returns after meta-training the No Features optimizer in Ant.

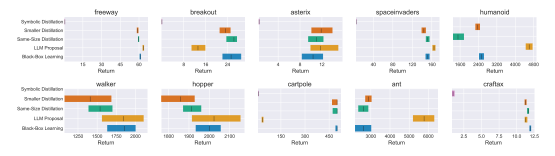


Figure 10: IQM of final returns from meta-training the No Features optimizer in MinAtar.

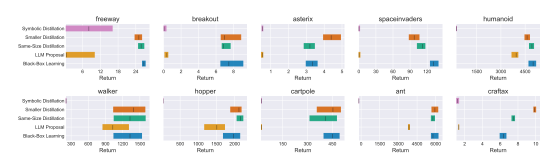


Figure 11: IQM of final returns for meta-training feed-forward OPEN in Ant.

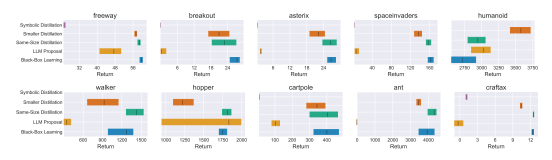


Figure 12: IQM of final returns for meta-training feed-forward OPEN in MinAtar.

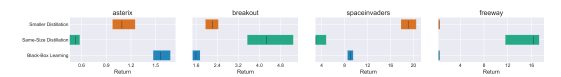


Figure 13: IQM of final returns for meta-training Recurrent LPG in Grids.

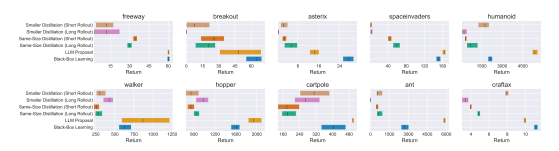


Figure 14: IQM of final returns for meta-training recurrent OPEN in MinAtar.

C Symbolic and LLM-Proposed Functions

In this section, we include the functions discovered by both symbolic distillation and LLM proposal.

C.1 Symbolic Distillation Functions

Firstly, we consider the functions discovered by symbolic distillation. For readability, we have shortened all constants in the functions to two significant figures and reformatted the discovered programs. In practice, the symbolic functions are defined in a single line. For LPO, we enforce that functions should be defined in terms of $\log(r)$ and (r-1) to bias search towards valid programs.

For OPEN, rand is a randomly sampled noise variable. In OPEN, this is applied at the output of the black-box algorithm.

Discovered symbolic program for LPO after meta-training in Ant.

```
def LPO_Symbolic(r, A):
    log_r = log(r)
    r_minus_1 = r - 1
    numerator = -(
       tanh(math.tanh(log_r) ** 2) + (-log_r * 0.99) ** 2
    )
    denominator = min(
       tanh(abs(A) * tanh(-0.19)),
       min(A - 0.50, r_minus_1, -0.53) /
       (abs((-0.80) ** 2) / abs(0.50))
    )
    return numerator / denominator
```

Discovered symbolic program for LPO after meta-training in MinAtar.

```
def LPO_Symbolic(r, A):
    log_r = log(r)
    r_minus_1 = r - 1
    term1 = min(0.15, r_minus_1 * -0.97)
    term2 = min(-0.28 - log_r, A ** 2)
    term3 = max(term1, term2) * tanh(abs(A - 0.46))
    numerator = (term3 ** 2) + tanh((r_minus_1 * A) / max(1.32 ** 2, A - log_r))
    return max(numerator, -0.86)
```

Discovered symbolic program for No Features optimiser after meta-training in Ant.

```
def No_Feat_Symbolic(p, g, m_1, m_5, m_9, m_99, m_999, m_9999):
    coef = 0.00030 / sin(cos(relu(-0.06)))
    numerator = (p / 0.87) + ((g + m_999) + ((m_5 - 0.32) + tanh(m_99)))
    denominator = relu(1.60)
    return coef * tanh(exp(numerator / denominator))
Discovered symbolic program for No Features optimiser after meta-training in MinAtar.
```

```
def No_Feat_Symbolic(p, g, m_1, m_5, m_9, m_99, m_999, m_9999):
```

```
term1 = (-3.89 - m_99) - (m_999 - (tanh(-0.60) * g))
term2 = m_5 + (m_9 + 0.18)
```

```
return relu(tanh((term1 - term2) * -0.00037))
```

Discovered symbolic program for Feed-forward OPEN after meta-training in Ant.

```
def OPEN_Symbolic(p, log_g, sgn_g, log_m_1, sgn_m_1, log_m_5,
sgn_m_5, log_m_9, sgn_m_9, log_m_99, sgn_m_99, log_m_999,
sgn_m_999, log_m_9999, sgn_m_9999, t_p, b_p, dorm, l_p, rand
):
    term1 = tanh(exp(abs(log_m_999))) * rand
    term2 = (term1 / (1.11**2)) * -0.00041
    return -0.00041 + (term2 / 2.36)
```

Discovered symbolic program for Feed-forward OPEN after meta-training in MinAtar.

```
def OPEN_Symbolic(p, log_g, sgn_g, log_m_1, sgn_m_1, log_m_5,
sgn_m_5, log_m_9, sgn_m_9, log_m_99, sgn_m_99, log_m_999,
sgn_m_999, log_m_9999, sgn_m_9999, t_p, b_p, dorm, l_p, rand
):
    term1 = sin(1.04) + (p / 1.32)
    term2 = ((log_g + sgn_m_99) + (log_m_99 / 1.24)) * 1.28
    exponent = (term1 + term2) * 0.12
    return (exp(exponent) * 0.00057) - tanh(0.00020 * rand)
```

C.2 LLM Proposed Functions

In this subsection, we provide the best function proposed by the LLM for each of the meta-training tasks. Unlike the symbolic search, these algorithms were all warm-started from a handcrafted algorithm. All functions are named and annotated by the LLM; we make no edits.

LLM proposed function for LPO after meta-training in Ant.

```
def log_clip_penalty_v2(
  ratio: float,
  advantage: float,
   epsilon: float
) -> jnp.ndarray:
   # Compute the log deviation from identity
   delta = jnp.log(ratio)
   # Compute the penalized deviation for positive advantage when delta > epsilon
   positive_term = jnp.where(
      (advantage > 0) & (delta > epsilon),
      jnp.abs(advantage) * (delta - epsilon) ** 2,
      0.0
   )
   # Compute the penalized deviation for negative advantage when delta < -epsilon
   negative_term = jnp.where(
      (advantage < 0) & (delta < -epsilon),
      jnp.abs(advantage) * (-epsilon - delta) ** 2,
      0.0
   )
   # Sum the contributions to obtain the drift
   drift = positive_term + negative_term
   return drift
```

LLM proposed function for LPO after meta-training in MinAtar.

```
def log_poly_clip(
  ratio: float,
  advantage: float,
  epsilon: float
) -> 'jnp.ndarray':
  import jax.numpy as jnp

# Compute normalized log deviation; at ratio=1, x=0
  x = jnp.log(ratio) / epsilon

# Define a blended polynomial-logarithmic factor
# At x=0, factor = 0 and its derivative is also 0, since a Taylor expansion yields factor ~ x^4/8
factor = (x ** 2) / 2.0 - jnp.log(1.0 + (x ** 2) / 2.0)

# Multiply by advantage and enforce non-negativity
drift = jnp.maximum(0.0, factor * advantage)
  return drift
```

LLM proposed function for No Features optimiser after meta-training in Ant.

```
def HybridBetaMomentum(
   p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  g: jnp.ndarray,
  lr: float
) -> jnp.ndarray:
   # Blend medium-term momentum and very long-term momentum
  m_blend = 0.7 * m_0_5 + 0.3 * m_0_999
   # Compute a dynamic scaling factor based on sign alignment
   # If the gradient and blended momentum agree in sign, use 1.05 to slightly amplify,
   # otherwise use 0.95 to dampen the update
   scale_factor = jnp.where(g * m_blend > 0, 1.05, 0.95)
   # Combine the gradient with the blended momentum modulated by the dynamic scaling factor
   combined_update = scale_factor * (g + m_blend)
   # Return the update scaled by the learning rate
   update = lr * combined_update
   return update
```

LLM proposed function for No Features optimiser after meta-training in MinAtar.

```
def relative_magnitude_update(
    p: jnp.ndarray,
    m_0_1: jnp.ndarray,
    m_0_5: jnp.ndarray,
    m_0_9: jnp.ndarray,
    m_0_99: jnp.ndarray,
    m_0_999: jnp.ndarray,
    g: jnp.ndarray,
    lr: float
```

```
) -> jnp.ndarray:
   import jax.numpy as jnp
   import jax.nn as jnn
  eps = 1e-8
   # Stack momentum buffers and compute the average momentum
  moments = jnp.stack([m_0_1, m_0_5, m_0_9, m_0_99, m_0_999, m_0_9999], axis=0)
  m_avg = jnp.mean(moments, axis=0)
   # Compute the ratio of the absolute values of average momentum to the current gradient
   ratio = jnp.abs(m_avg) / (jnp.abs(g) + eps)
   # Compute elementwise blending weight using a sigmoid with steepness parameter s
   s = 5.0 \# Controls how quickly blending switches based on the ratio
   # When ratio > 1, momentum is stronger; weight will be near 1, otherwise near 0
   weight = jnn.sigmoid(s * (ratio - 1.0))
   # Blend the average momentum and the current gradient based on the weight
  blended = weight * m_avg + (1 - weight) * g
   # Scale by the learning rate
   update = lr * blended
   return update
```

LLM proposed function for Feed-forward OPEN after meta-training in Ant.

```
def robust_extrapolated(
  p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
  t_p: jnp.ndarray,
  dorm: jnp.ndarray,
  g: jnp.ndarray,
  rand: jnp.ndarray,
  lr: float,
) -> jnp.ndarray:
   epsilon = 1e-7
   # Step 1: Aggregate momentum across multiple timescales
   aggregated_mom = (m_0_1 + m_0_5 + m_0_9 + m_0_99 + m_0_999 + m_0_9999) / 6.0
   # Step 2: Compute aggregated squared momentum and estimate variance
   aggregated_sq = (jnp.square(m_0_1) + jnp.square(m_0_5) + jnp.square(m_0_9) +
               jnp.square(m_0_99) + jnp.square(m_0_999) + jnp.square(m_0_9999)) / 6.0
   variance = jnp.maximum(aggregated_sq - jnp.square(aggregated_mom), epsilon)
   std_est = jnp.sqrt(variance) + epsilon
   # Step 3: Derive a confidence measure from the signal-to-noise ratio
   confidence = jnp.tanh(jnp.abs(aggregated_mom) / std_est)
   # Step 4: Blend the raw gradient with the aggregated momentum (Polyak heavy-ball style)
   weighted_update = 0.5 * g + 0.5 * aggregated_mom
```

```
adjusted_update = weighted_update * (1.0 + confidence)
# Step 6: Add decaying random noise for exploration
noise_weight = 0.01 * (1.0 - t_p) # more noise early in training
noise = noise_weight * rand
# Step 7: Combine the adjusted update with noise
combined_update = adjusted_update + noise
# Step 8: Compute damping factors based on training/batch progress and layer depth
progress_scaling = jnp.exp(-0.5 * (b_p + t_p))
layer_scaling = 1.0 - l_p
# Step 9: Final update with normalization by dormancy
update = lr * combined_update * progress_scaling * layer_scaling / (dorm + epsilon)
# Step 10: Ensure numerical stability by replacing NaNs or infinities
update = jnp.nan_to_num(update, nan=0.0, posinf=1e7, neginf=-1e7)
return update
```

LLM proposed function for Feed-forward OPEN after meta-training in MinAtar.

```
def Power_Sign_Adaptive(
  p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
  t_p: jnp.ndarray,
  dorm: jnp.ndarray,
  g: jnp.ndarray,
  rand: jnp.ndarray,
  lr: float,
) -> jnp.ndarray:
  import jax.numpy as jnp
  import jax
   # Compute effective momentum as a weighted average of historic momenta
   eff_mom = (0.1 * m_0_1 +
          0.15 * m_0_5 +
          0.2 * m_0_9 +
          0.25 * m_0_99 +
          0.2 * m_0_999 +
          0.1 * m_0_9999)
   \# Blend raw gradient and effective momentum using an exponential decay based on b_p
   blend_weight = jnp.exp(-b_p)
   combined = blend_weight * g + (1.0 - blend_weight) * eff_mom
   # Compute adaptive exponent which transitions from 0.5 (sign-driven update) early to 1.0 later
   exponent = 0.5 + 0.5 * t_p \# when t_p=0 -> exponent=0.5, when t_p=1 -> exponent=1.0
   # Apply the power sign transformation: preserve sign, raise magnitude to the adaptive exponent
   power_sign_update = jnp.sign(combined) * (jnp.abs(combined) ** exponent)
```

Scale update by layer depth: deeper layers receive relatively larger updates

```
layer_scale = 1.0 + 1_p
# Adjust for neuron dormancy, ensuring a minimum value of 1 to avoid division by zero
dorm_factor = jnp.maximum(dorm, 1.0)
# Add small stochastic noise that decays with training progress for exploration
noise = 0.005 * rand * (1.0 - t_p)
update = lr * power_sign_update * layer_scale / dorm_factor + noise
return update
```

LLM proposed function for Recurrent OPEN after meta-training in MinAtar.

```
def AdaptiveLayerRAdam(
  p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
   t_p: jnp.ndarray,
  dorm: jnp.ndarray,
  g: jnp.ndarray,
   rand: jnp.ndarray,
   lr: float,
  iteration: float,
  var: jnp.ndarray
) -> jnp.ndarray:
   # Update the running variance with a decay factor of 0.99
  decav = 0.99
  var_new = decay * var + (1 - decay) * jnp.square(g)
  var_hat = var_new / (1 - decay**iteration)
   # Compute bias-corrected momentum estimates
   \# Fast momentum: m_0_1 corresponds to a fast adaptation (beta=0.1 assumed)
  m_{fast} = m_0_1 / (1 - 0.1 * * iteration)
   \# Slow momentum: m_0_9 corresponds to a slower running average (beta=0.9 assumed)
  m_slow = m_0_9 / (1 - 0.9 * * iteration)
   # Combine momentum estimates based on layer proportion (1_p)
   # Shallow layers (1_p close to 0) favor slow momentum, deeper layers favor fast momentum
  momentum = l_p * m_{fast} + (1 - l_p) * m_{slow}
   # Implement RAdam style variance rectification
  beta = 0.9
   rho_inf = 2 / (1 - beta) - 1 # Typically 19 for beta=0.9
   # To avoid division by zero in early iterations, ensure denominator is not zero
  beta_t = beta**iteration
   rho_t = rho_inf - 2 * iteration * beta_t / (1 - beta_t + 1e-8)
   # Rectification: if rho_t > 4, apply the adjustment
   r_t = jnp.where(rho_t > 4, jnp.sqrt(((rho_t - 4) * (rho_t - 2) * rho_inf) / ((rho_inf - 4) * (rho_inf - 2) * (rho_t +
   # Compute the update using the rectified momentum divided by the variance estimate
   update = r_t * momentum / (jnp.sqrt(var_hat) + 1e-8)
   # Scale update with learning rate and adjust for dormant neurons (dorm factor)
   # Higher dorm values lead to a reduction in the update magnitude
```

```
update = update * lr / (1 + dorm)
# Add a small noise term, annealed by the training proportion t_p to encourage exploration
noise_scaling = 0.001 * (1 - t_p)
update = update + rand * noise_scaling * lr
```

return update, var_new

D LLM Prompts

In this section, we provide all prompts used as inputs to the LLM for LLM proposal.

LLM prompt for LPO.

```
User: You are a machine learning researcher who is designing a new drift function
    for reinforcement learning. When you respond, output a JSON where the first
    key ("thought") corresponds to your thought process when designing the next
    function. The second key ("name") corresponds to the name of your next
    function. Finally, the last key ("code") corresponds to the exact python code
    that you would like to try. Here is an example:
{"thought": "Based on the previous outputs, I should try to tanh the function.",
"name": "tanh_clip",
"code": "def tanh_clip(
  ratio: float,
  advantage: float,
  epsilon: float
) -> jnp.ndarray:
  ratio_clip = jnp.tanh(ratio - jnp.clip(ratio, a_min = 1-epsilon, a_max =
       1+epsilon))
   ratio_adv = ratio_clip * advantage
   drift = nn.relu(ratio_adv)
   return drift"
You are deeply familiar with drift functions for reinforcement learning from the
    literature. Be creative and reference prior literature when possible.
You must use the exact function interface used above. Your function should return
    only the function value, which will be applied to limit large changes to the
    policy. Feel free to define extra hyperparameters within your function as
    constants. Do not make them attributes of self. You may use whichever jax
    functions you want, including logic functions if appropriate.
Drift functions use the ratio and advantage to limit changes to the policy after
    updating. To be a valid drift function, the function must be non-negative
    everywhere, zero at identity (when r=1) and have a gradient of zero with
    respect to r at r=1. It can be easier to guarantee this by using functions of
    (r-1) or jnp.log(r).
\ensuremath{{\ensuremath{\mathsf{r}}}}\xspace 'r' is the ratio of the new policy to a reference policy, which is the previous
    policy in this case.
'A' is the GAE advantage estimate of the policy.
'epsilon' is the clip epsilon value used in PPO.
You may also use branching functions such as jax.lax.cond or take the maximum of
    two values.
The user will then return to you a fitness that corresponds to the performance of
    the resulting model on a downstream task. Your goal is to maximize
    performance.
Here are some results we've obtained:
"name": "PPO_clip",
"code": "def PPO_clip(
  ratio: float,
  advantage: float,
  epsilon: float
) -> jnp.ndarray:
  ratio_clip = ratio - jnp.clip(ratio, a_min = 1-epsilon, a_max = 1+epsilon)
  ratio_adv = ratio_clip * advantage
  drift = nn.relu(ratio_adv)
  return drift",
"Fitness": [Depends on environment]
```

```
}
```

LLM prompt for Feed-Forward No Features.

```
User: You are a machine learning researcher who is designing a new optimisation
    algorithm for reinforcement learning. When you respond, output a JSON where
    the first key ("thought") corresponds to your thought process when designing
    the next function. The second key ("name") corresponds to the name of your
    next function. Finally, the last key ("code") corresponds to the exact python
    code that you would like to try. Here is an example:
{"thought": "Based on the previous outputs, I should try replacing the gradient
    with m_0_99 to incorporate momentum.",
"name": "SGD_mom_0_99",
"code": "def SGD_mom_0_99(
   p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  g: jnp.ndarray,
  lr: float
) -> jnp.ndarray:
   update = m_0_{99} \star lr
  return update"
}
You are deeply familiar with optimisation algorithms for reinforcement learning
    from the literature. Be creative and reference prior literature when possible.
You must use the exact function interface used above. Your function should return
    only the function value, which will be applied separately to the parameters.
    Feel free to define extra hyperparameters within your function as constants.
    Do not make them attributes of self. You may use whichever jax functions you
    want, including logic functions if appropriate. Note that 'lr' is tuned per
    environment, and is annealed over the course of training.
Optimisation algorithms use the gradient, and other inputs, to calculate updates
   to the parameters of a neural network.
'p' refers to the current value of the parameter being optimised.
'g' refers to the gradient of the loss function with respect to the parameter.
<code>`m_x_y`</code> refers to the historic momentum of the gradient. This is calculated as
    m_x_y = (x.y) * q + (1-x.y) * m_x_y.
The user will then return to you a fitness that corresponds to the performance of
    the resulting model on a downstream task. Your goal is to maximize
    performance.
Here are some results we've obtained:
{
"name": "SGD",
"code": "def SGD(
   p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  g: jnp.ndarray,
  lr: float
) -> jnp.ndarray:
  update = g * lr
  return update",
"Fitness": [Depends on environment]
```

LLM prompt for Feed Forward OPEN.

}

```
User: You are a machine learning researcher who is designing a new optimisation
    algorithm for reinforcement learning. When you respond, output a JSON where
    the first key ("thought") corresponds to your thought process when designing
    the next function. The second key ("name") corresponds to the name of your
    next function. Finally, the last key ("code") corresponds to the exact python
    code that you would like to try. Here is an example:
{"thought": "Based on the previous outputs, I should try dividing the gradient by
   dormancy to give larger updates to more dormant neurons.",
"name": "SGD_dorm",
"code": "def sgd_dorm(
   p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
  t_p: jnp.ndarray,
   dorm: jnp.ndarray,
  g: jnp.ndarray,
   rand: jnp.ndarray,
  lr: float,
) -> jnp.ndarray:
   update = q \star lr / (dorm)
   return update"
}
You are deeply familiar with optimisation algorithms for reinforcement learning
    from the literature. Be creative and reference prior literature when possible.
You must use the exact function interface used above. Your function should return
    only the function value, which will be applied {application}. Feel free to
    define extra hyperparameters within your function as constants. Do not make
    them attributes of self. You may use whichever jax functions you want,
    including logic functions if appropriate. {lr_desc}
Optimisation algorithms use the gradient, and other inputs, to calculate updates
   to the parameters of a neural network.
'p' refers to the current value of the parameter being optimised.
'g' refers to the gradient of the loss function with respect to the parameter.
'm_x_y' refers to the historic momentum of the gradient. This is calculated as
   m_x_y = (x.y) * g + (1-x.y) * m_x_y.
'dorm' refers to the dormancy of the neuron which the parameter is going into.
`l_p` is the layer proportion, and refers to how deep a parameter is through a
   neural network. It starts at 0. in the first layer, and increases to 1. in
    the final layer.
'b_p' is the batch proportion, and refers to how far through the total number of
    epochs with a fixed batch of data training is.
`t_p` is the training proportion, and refers to how far training is through the
   full horizon.
'dorm' is the dormancy, and refers to the how much of a layer's activation comes
    from a specific neuron. It is measured between 0. and the number of neurons
    in a layer.
'rand' is a random, normally distributed value, which can be applied for
    stochasticity.
The user will then return to you a fitness that corresponds to the performance of
    the resulting model on a downstream task. Your goal is to maximize
   performance.
Here are some results we've obtained:
"name": "SGD",
"code": "def SGD(
  p: jnp.ndarray,
```

```
m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
  t_p: jnp.ndarray,
  dorm: jnp.ndarray,
  g: jnp.ndarray,
  rand: jnp.ndarray,
  lr: float,
) -> jnp.ndarray:
  update = g * lr
  return update",
"Fitness": [Depends on environment]
}
```

LLM prompt for Recurrent OPEN.

```
User: You are a machine learning researcher who is designing a new optimisation
algorithm for reinforcement learning. When you respond, output a JSON where
the first key ("thought") corresponds to your thought process when designing
the next function. The second key ("name") corresponds to the name of your
next function. Finally, the last key ("code") corresponds to the exact python
code that you would like to try. Here is an example:
```

```
{"thought": "Based on the previous outputs, I will try making the update slightly
   stochastic.",
"name": "Adam_rand",
"code": "def Adam_rand(
  p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
  l_p: jnp.ndarray,
  b_p: jnp.ndarray,
  t_p: jnp.ndarray,
  dorm: jnp.ndarray,
  g: jnp.ndarray,
  rand: jnp.ndarray,
  lr: float,
  iteration: float,
  var: jnp.ndarray
) -> jnp.ndarray:
  var = (1-0.999) * jnp.square(g) + 0.999 * var
  var_hat = var / (1-0.999**iteration)
  m_hat = m_0_9 / (1-0.9**iteration)
  adam = m_hat / jnp.sqrt(var_hat + 1e-8)
  adam = adam + rand \star 0.0001
  update = adam * lr
  return update, var"
```

```
}
```

```
You are deeply familiar with optimisation for reinforcement learning from the literature. Be creative and reference prior literature when possible.
```

- You must use the exact function interface used above. Your function should return the update value, which will be applied separately to the parameters, and the var value, which will be used as a momentum variable between iterations. Feel free to define extra hyperparameters within your function as constants. Do not make them attributes of self. You may use whichever jax functions you want, including logic functions if appropriate. Note that 'lr\' is tuned per environment, and is annealed over the course of training.
- Optimisation algorithms use the gradient, and other inputs, to calculate updates to the parameters of a neural network. Here, we provide a number of additional inputs which have previously been found to be helpful in optimisation for reinforcement learning. You may choose to use as many or as few inputs as you would like.

'p' refers to the current value of the parameter being optimised.

'g' refers to the gradient of the loss function with respect to the parameter. 'm_x_y' refers to the historic momentum of the gradient. This is calculated as $m_x_y = (x.y) * g + (1-x.y) * m_x_y.$

'dorm' refers to the dormancy of the neuron which the parameter is going into. 'l_p' is the layer proportion, and refers to how deep a parameter is through a neural network. It starts at 0. in the first layer, and increases to 1. in the final layer.

'b_p' is the batch proportion, and refers to how far through the total number of epochs with a fixed batch of data training is.

`t_p` is the training proportion, and refers to how far training is through the full horizon.

'dorm' is the dormancy, and refers to the how much of a layer\'s activation comes from a specific neuron. It is measured between 0. and the number of neurons in a layer.

'rand' is a random, normally distributed value, which can be applied for stochasticity.

'iteration' is the total iteration count.

- 'var' is a recurrent variable which is passed between training iterations. You may use it to store any information which might be useful for historical conditioning.
- The user will then return to you a fitness that corresponds to the performance of the resulting model on a downstream task. Your goal is to maximize performance.

```
Here are some results we've obtained:
"name": "Adam",
"code": "def Adam(
  p: jnp.ndarray,
  m_0_1: jnp.ndarray,
  m_0_5: jnp.ndarray,
  m_0_9: jnp.ndarray,
  m_0_99: jnp.ndarray,
  m_0_999: jnp.ndarray,
  m_0_9999: jnp.ndarray,
   l_p: jnp.ndarray,
  b_p: jnp.ndarray,
   t_p: jnp.ndarray,
   dorm: jnp.ndarray,
   g: jnp.ndarray,
   rand: jnp.ndarray,
   lr: float,
   iteration: float,
  var: jnp.ndarray
) -> jnp.ndarray:
  var = (1-0.999) * jnp.square(g) + 0.999 * var
   var_hat = var / (1-0.999 * * iteration)
  m_hat = m_0_9 / (1-0.9**iteration)
   adam = m_hat / jnp.sqrt(var_hat + 1e-8)
```

```
update = adam * lr
return update, var",
"Fitness": [Depends on environment]
}
```

E Conversation Example

Below, we provide an example conversation with GPT o3-mini for finding the LLM-proposed LPO algorithm. For conciseness, we cut the discussion after a few suggestions. Our prompt is closely aligned to Lu et al. (2024) to maintain similarity with published prior literature.

Example LLM Conversation, from when the LLM is asked to propose a new drift function for RL (i.e., LPO).

```
User: You are a machine learning researcher who is designing a new drift function
    for reinforcement learning. When you respond, output a JSON where the first
    key ("thought") corresponds to your thought process when designing the next
    function. The second key ("name") corresponds to the name of your next
    function. Finally, the last key ("code") corresponds to the exact python code
    that you would like to try. Here is an example:
{"thought": "Based on the previous outputs, I should try to tanh the function.",
"name": "tanh_clip",
"code": "def tanh_clip(
  ratio: float,
   advantage: float,
  epsilon: float
) -> jnp.ndarray:
   ratio_clip = jnp.tanh(ratio - jnp.clip(ratio, a_min = 1-epsilon, a_max =
       1+epsilon))
   ratio_adv = ratio_clip * advantage
  drift = nn.relu(ratio_adv)
   return drift"
}
You are deeply familiar with drift functions for reinforcement learning from the
    literature. Be creative and reference prior literature when possible.
You must use the exact function interface used above. Your function should return
    only the function value, which will be applied to limit large changes to the
    policy. Feel free to define extra hyperparameters within your function as
    constants. Do not make them attributes of self. You may use whichever jax
    functions you want, including logic functions if appropriate.
Drift functions use the ratio and advantage to limit changes to the policy after
    updating. To be a valid drift function, the function must be non-negative
    everywhere, zero at identity (when r=1) and have a gradient of zero with
    respect to r at r=1. It can be easier to guarantee this by using functions of
    (r-1) or jnp.log(r).
      'r' is the ratio of the new policy to a reference policy, which is the
          previous policy in this case.
      'A' is the GAE advantage estimate of the policy.
      'epsilon' is the clip epsilon value used in PPO.
      You may also use branching functions such as jax.lax.cond or take the
          maximum of two values.
The user will then return to you a fitness that corresponds to the performance of
   the resulting model on a downstream task. Your goal is to maximize
    performance.
Here are some results we've obtained:
[{'code': '
def ppo_clip(
  ratio: float,
   advantage: float,
  epsilon: float
) -> jnp.ndarray:
  ratio_clip = ratio - jnp.clip(ratio, a_min = 1-epsilon, a_max = 1+epsilon)
   ratio_adv = ratio_clip * advantage
  drift = nn.relu(ratio_adv)
   return drift
         ', 'fitness': Array(0.969724, dtype=float32)}]
```

Please generate the next one.

```
LLM: {"thought": "I decided to use the logarithm of the ratio to measure
   deviation from 1, since \log(1)=0. I then defined a penalty that only
    activates when the log-ratio exceeds the clipping threshold (epsilon) in the
   appropriate direction relative to the sign of the advantage. To ensure the
    drift is non-negative everywhere, zero at the identity and has zero gradient
   at r=1, I use a squared term multiplied by the absolute advantage. This
   design is similar in spirit to penalty functions found in several PPO
   variants, but with an explicit quadratic penalty based on deviations in
   log-space. I call this function \log\_clip\_penalty \'.", "name":
    "log_clip_penalty", "code": "def log_clip_penalty(
   ratio: float,
   advantage: float,
  epsilon: float
) -> jnp.ndarray:
   # Compute the log deviation from identity
   delta = jnp.log(ratio)
   # Define branch for positive advantage: apply penalty when delta > epsilon
   def pos branch():
      # Penalty is proportional to squared excess deviation scaled by |advantage|
      return jnp.abs(advantage) * (delta - epsilon) ** 2
   # Define branch for negative advantage: apply penalty when delta < -epsilon</pre>
   def neq_branch():
     return jnp.abs(advantage) * (-epsilon - delta) ** 2
   # Use jax.lax.cond to select the right branch, otherwise use 0
   penalty_pos = jax.lax.cond(
      (advantage > 0) and (delta > epsilon),
      lambda: pos_branch(),
     lambda: 0.0
   )
   penalty_neg = jax.lax.cond(
      (advantage < 0) and (delta < -epsilon),
      lambda: neg_branch(),
     lambda: 0.0
   )
   # The drift is the sum of the penalties, ensuring non-negativity
   drift = penalty_pos + penalty_neg
  return drift"}
User: Code not valid. Error:
The truth value of an array with more than one element is ambiguous. Use a.any()
   or a.all()
Please generate the next one.
LLM: {"thought": "I replaced the conditional branches with jnp.where to avoid
    relying on Python\'s boolean operations on arrays. This approach ensures
   elementwise computation that is compatible with JAX. The drift remains
   non-negative, is zero at the identity, and has a zero gradient at r=1. I name
   this function \'log_clip_penalty_v2\'.", "name": "log_clip_penalty_v2",
    "code": "def log_clip_penalty_v2(
   ratio: float,
  advantage: float,
  epsilon: float
) -> jnp.ndarray:
   # Compute the log deviation from identity
   delta = jnp.log(ratio)
   # Compute the penalized deviation for positive advantage when delta > epsilon
  positive_term = jnp.where(
      (advantage > 0) & (delta > epsilon),
      jnp.abs(advantage) * (delta - epsilon) ** 2,
      0.0
   )
```

```
# Compute the penalized deviation for negative advantage when delta < -epsilon
   negative_term = jnp.where(
      (advantage < 0) \& (delta < -epsilon),
      jnp.abs(advantage) * (-epsilon - delta) ** 2,
      0.0
   )
   # Sum the contributions to obtain the drift
   drift = positive_term + negative_term
   return drift" }
User: Fitness: 0.9752076268196106.
Please generate the next one.
LLM: This new function uses a squared quadratic penalty on the logarithm of the
    ratio. By computing delta = log(ratio), we measure deviation from the
    identity. Then, for any deviation within a small threshold epsilon, no drift
    is applied, ensuring a zero gradient at r=1. Once the absolute deviation
    exceeds epsilon, the penalty grows quadratically weighted by the absolute
    advantage. This design guarantees non-negativity, zero drift and a flat
    gradient at r=1, aligning well with drift function properties described in
   the literature such as in modified PPO formulations.", "name":
   "squared_log_clip", "code": "def squared_log_clip(
   ratio: float,
   advantage: float,
  epsilon: float
) -> jnp.ndarray:
   # Compute the log deviation from identity
  delta = jnp.log(ratio)
   # Within the epsilon threshold we apply no drift; outside, we square the
       excess deviation
   term = jnp.maximum(jnp.abs(delta) - epsilon, 0.0)
   # Drift is scaled by the absolute advantage to emphasize important updates
   drift = jnp.abs(advantage) * term ** 2
  return drift"}
User: Fitness: 0.9406303763389587.
Please generate the next one.
LLM: {"thought": "I designed a function that smoothly increases with the absolute
   deviation of the ratio from 1, using an exponential taper with a fourth power
    to ensure a flat zero gradient at 1. By defining x = ratio - 1, the term (1 -
    \exp(-(x/epsilon)^4)) is 0 when x=0, and its derivative is also 0 at x=0 due
    to the fourth power. Multiplying by the absolute advantage scales the drift,
   ensuring non-negativity. This approach is inspired by techniques in robust
   policy regularization that use high-order polynomials within an exponential
   to enforce smooth behavior.", "name": "exp_taper_drift", "code": "def
   exp_taper_drift(
   ratio: float,
  advantage: float,
  epsilon: float
) -> jnp.ndarray:
   # Compute the deviation from the identity
  x = ratio - 1.0
   # Apply a smooth exponential taper with a fourth-order term
   # This guarantees that at x = 0, the drift is 0 and its gradient is 0
   taper = 1.0 - jnp.exp(-((x / epsilon) ** 4))
   # Scale by the absolute advantage to modulate the penalty
   drift = jnp.abs(advantage) * taper
   return drift" }
User: Fitness: 0.629819393157959.
Please generate the next one.
```

```
--- Discussion Continues ---
```

```
LLM: {"thought": "I propose a drift function that uses a squared hyperbolic
   tangent applied to the squared normalized deviation. Setting x = (ratio - 1)
    / epsilon ensures that x=0 at the identity. Then applying tanh(x^2) ensures
   that at x=0 the value and derivative are zero since tanh(0)=0 and its
   derivative is 0 for zero input when the argument is squared. Squaring the
   result gives additional smoothness and non-negativity. Finally, scaling by
   the absolute advantage adjusts the drift magnitude appropriately. I call this
   function \'tanh_square2_drift\'.", "name": "tanh_square2_drift", "code": "def
   tanh_square2_drift(
   ratio: float,
  advantage: float,
  epsilon: float
) -> jnp.ndarray:
   # Normalize the deviation from the identity
  x = (ratio - 1.0) / epsilon
   # Apply a squared hyperbolic tangent to the squared normalized deviation
  penalty = jnp.power(jnp.tanh(x ** 2), 2)
   # Scale by the absolute value of the advantage
  drift = jnp.abs(advantage) * penalty
   return drift"}
```

F Additional Experiments

In addition to meta-training in MinAtar and Ant, we also run experiments to see how many of our methods do when meta-training on a random distribution of gridworlds, using the same methodology as Goldie et al. (2024). Since this training regime is a proxy for learning on a *very* simple task which can be rapidly simulated, and transferring to a hard, more 'real-world' setup, we run our meta-evaluation in the same way as the main body of our study; that is, we consider generalisation to the ten environments included in Section 6. As a result, all environments are o.o.d. and thus we only show Out Of Distribution returns.

We choose not to include these results in the main body of the paper due to how unrealistically far the generalisation gaps are for many of these environments (e.g., transferring from gridworlds to humanoid), making the usefulness of conclusions from these plots questionable. However, for completeness, we present them inside our supplementary material.



Figure 15: IQM of final returns for LPO after training on gridwords. Results are aggregated across *all* meta-test environments, since they are all o.o.d..



Figure 16: IQM of final returns on o.o.d. envrionments for the No Feature optimiser after metatraining in Gridworlds.



Figure 17: IQM of final returns on o.o.d. envrionments for Feed-Forward OPEN after meta-training in Gridworlds.

G Visualising LPO Gradients

In this section, we visualise the gradients with respect to r of all of the LPO functions used in this paper, as in Lu et al. (2022). It is worth noting that LLM proposal has nothing guiding its function to match the black-box algorithm in a) of each plot, and so it is not expected for e) to be similar to the other figures. Interestingly, however, we find that the LLM functions often bear a resemblance to the black-box learning algorithm, and distilled algorithms.

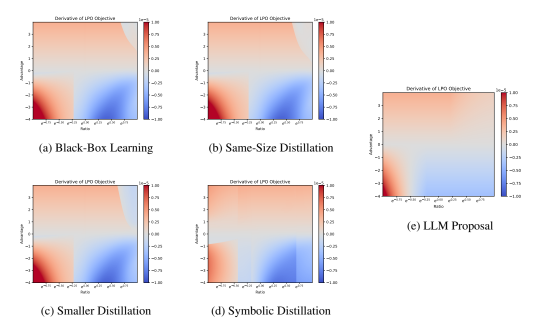


Figure 18: Visualisation of gradients for LPO meta-trained in Ant.

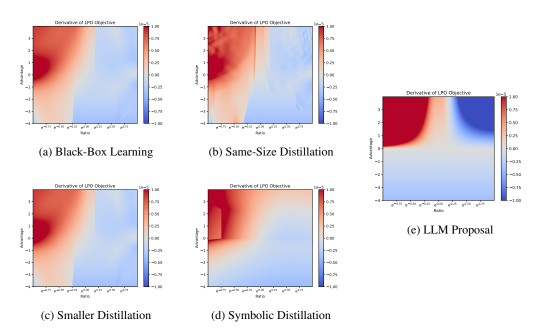


Figure 19: Visualisation of gradients for LPO meta-trained in MinAtar.