Learning to Optimize: A Primer and A Benchmark

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Abstract

Learning to optimize (L2O) is an emerging approach that leverages machine learning to develop optimization methods, aiming at reducing the laborious iterations of hand engineering. It automates the design of an optimization method based on its performance on a set of training problems. This data-driven procedure generates methods that can efficiently solve problems similar to those in the training. In sharp contrast, the typical and traditional designs of optimization methods are theory-driven, so they obtain performance guarantees over the classes of problems specified by the theory. The difference makes L2O suitable for repeatedly solving a certain type of optimization problems over a specific distribution of data, while it typically fails on out-of-distribution problems. The practicality of L2O depends on the type of target optimization, the chosen architecture of the method to learn, and the training procedure. This new paradigm has motivated a community of researchers to explore L2O and report their findings.

This article is poised to be the first comprehensive survey and benchmark of L2O for continuous optimization. We set up taxonomies, categorize existing works and research directions, present insights, and identify open challenges. We also benchmarked many existing L2O approaches on a few but representative optimization problems. For reproducible research and fair benchmarking purposes, we released our software implementation and data in the package Open-L2O at https://github.com/VITA-Group/Open-L2O.

Keywords: Learning to Optimize, Meta Learning, Optimization, Algorithm Unrolling

1. Introduction

1.1 Background and Motivation

Classic optimization methods are typically hand-built by optimization experts based on theories and their experience. As a paradigm shift from this conventional design, learning to optimize (L2O) uses machine learning to improve an optimization method or even generate a completely new method. This paper provides a timely and up-to-date review of the rapid growing body of L2O results, with a focus on continuous optimization.

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Figure 1: (a) Classic optimizers are manually designed; they usually have few or no tuning parameters; (b) Learned optimizers are trained in an L2O framework over a set of similar optimizees (called a task distribution) and designed to solve unseen optimizees from the same distribution.

Classic optimization methods are built upon components that are basic methods — such as gradient descent, conjugate gradient, Newton steps, Simplex basis update, and stochastic sampling — in a theoretically justified manner. Most conventional optimization methods can be written in a few lines, and their performance is guaranteed by their theories. To solve an optimization problem in practice, we can select a method that supports the problem type at hand and expect the method to return a solution no worse than its guarantee.

L2O is an alternative paradigm that develops an optimization method by training, i.e., learning from its performance on sample problems. The method may lack a solid theoretical basis but improves its performance during the training process. The training process often occurs offline and is time consuming. However, the online application of the method is (aimed to be) time saving. When it comes to problems where the target solutions are difficult to obtain, such as nonconvex optimization and inverse-problem applications, the solution of a well-trained L2O method can have better qualities than those of classic methods. Let us call the optimization method (either hand-engineered or trained by L2O) the optimizer and call the optimization problem solvable by the method the optimizee. Figure 1 compares classic optimizers and L2O optimizers and illustrates how they are applied to optimizees (yellow boxes).

In many applications of optimization, the task is to perform a certain type of optimization over a specific distribution of data repeatedly. Each time, the input data that define the optimization are new but similar to the past. We say such an application has a narrow task distribution. Conventional optimizers may be tuned for a task distribution, but the underlying methods are design for a theory-specified class of optimization problems. We often describe a conventional optimizer by the formulation (and its math properties), not the task distribution. For example, we say an optimizer can minimize a smooth-and-convex objective function subject to linear constraints. In L2O, the training process shapes the optimizer according to both the formulation and the task distribution. When the distribution is concentrated, the learned optimizer can “overfit” to the tasks and may discover “short cuts” that classic optimizers do not take.

L2O aims to generate optimizers with the following strengths:

1. An optimizer learned by L2O is expected to complete a set of optimizees from the same task distribution at a much faster speed than classic methods. In particular, such an L2O method can run even faster than so-called optimal methods such as Nesterov’s faster gradient descent (FGD) [3] on a set of problems well suited for these optimal methods.
Figure 2: L2O for compressive sensing. In our prior work [1; 2], we successfully demonstrated that L2O optimizers (LISTA, LISTA-CPSS, TiLISTA and ALISTA) converge much faster than the two popular iterative solvers (ISTA, FISTA). Left: noiseless case. Right: noisy case (SNR = 20). X-axis is the number of iterations; Y-axis is normalized mean squared error (lower is better).

2. The learned optimizer may also return a higher-quality solution to a difficult task than classic methods given a similar amount of computing budget. For instance, an L2O method can recover a sparse signal that is much more faithful (meanwhile much faster) than LASSO. Therefore, when it comes to a small set of optimization tasks, L2O presents a potential approach to break the limits of analytic methods.

L2O has started to demonstrate great potential in some areas of optimization and applications. Examples include convex $\ell_1$-minimization [4], neural network training [5], black-box optimization [6] and combinatorial optimization [7]. For example, a class of L2O methods, in the form of algorithm unrolling (which we review below), has state-of-the-art results in compressed sensing and inverse problems. The L2O methods in [1; 2] converge much faster than the classic ISTA/FISTA optimizers (see Figure 2), using an order-of-magnitude fewer iterations on unseen optimizees from the same distribution to reach the same accuracy. Application areas benefiting from L2O methods include computer vision [8–9; 10], medical imaging [11], signal processing and communication [12–13], as well as computational biology [14–15]. For example, the applications of algorithm unrolling, a popular form of model-based L2O, has been specifically reviewed in Section 3.3. As another example, in the practical application of domain adaptation, model-free L2O has demonstrated the ability to adapt a pre-trained classifier to a new domain at less computational resource costs, achieving the same accuracy or generalization performance [16; 17].

1.2 Preliminaries

L2O starts with an architecture of the optimizer with free parameters to learn, so one must prepare a set of training samples that represent the task distribution, as well as choosing a training approach. The training process (Figure 1 (b) left) executes the approach, which iteratively applies the current optimizer to the sample optimizees and uses the observed performance to update the parameters. Over the time, the optimizer adapts to the training samples. This process trains an optimizer to take a series of actions on the optimizees, analogous of classic learning where we train ML models to make predictions or decisions.

We now formalize our notation. Consider an optimization problem $\min_x f(x)$ where $x \in \mathbb{R}^d$. A classic optimizer often iteratively updates $x$ based on a handcrafted rule. For

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1. In this example, the task is to recover a sparse signal, not necessarily solving the LASSO minimization.
example, the first-order gradient descent algorithm is to take an update at each iteration $t$ based on the local landscape information at the instantaneous point $x_t : x_{t+1} = x_t - \alpha \nabla f(x_t)$, where $\alpha$ is the step size.

L2O has lots of freedom to use the available information. The information $z_t$ available at time $t$ can include the existing iterates $x_0, \ldots, x_t$ as well as their gradients $\nabla f(x_0), \ldots, \nabla f(x_t)$, etc. Then the L2O approach models an update rule by a function $g$ of $z_t$: $x_{t+1} = x_t - g(z_t, \phi)$, where the mapping function $g$ is parameterized by $\phi$.

Finding an optimal update rule can be formulated mathematically as searching for a good $\phi$ over the parameter space of $g$. In order to find a desired $\phi$ associated with a fast optimizer, [5] proposed to minimize the weighted sum of the objective function $f(x_t)$ over a time span (which is called the unrolling length) $T$ given by:

$$
\min_{\phi} \mathbb{E}_{f \in T} \left[ \sum_{t=1}^{T} w_t f(x_t) \right], \quad \text{with} \quad x_{t+1} = x_t - g(z_t, \phi), \quad t = 1, \ldots, T - 1,
$$  \hspace{1cm} (1)

where $w_1, \ldots, w_T$ are the weights and $f$ represents an optimizee from an ensemble $T$ of optimizees that represent the target task distribution. Note that the parameter $\phi$ determines the objective value through determining the iterates $x_t$. L2O solves the training problem eq. (1) for a desirable $\phi$ and the update rule $g(z_t, \phi)$.

In practice, the choices of $w_t$ vary by case and depend on empirical settings. For example, many L2O models for sparse coding unroll to a fixed length $T$ for all optimizees, and then only minimize the step-$T$ functional value [18; 19], i.e., $w_T = 1$ and $w_1 = \cdots = w_{T-1} = 0$. Paper [5], on the other hand, assigns nonzero $w_t$ values and report more efficient Backpropagation Through Time (BPTT).

In L2O, one specifies the architecture of optimizer, which consists of both learnable components $\phi$ and fixed components. We address both of them in this paper. The update rule $g$ is often parameterized by multi-layer neural networks or recurrent neural networks. In theory, neural networks are universal approximators, so L2O may discover completely new, optimal update rules without referring to any existing updates. Since this kind of L2O architecture does not need a model, we refer to it as model-free L2O.

The shortcomings of model-free L2O include: lacking convergence guarantees and requiring a high number of training samples. On tasks where classic operations — such as projection, normalization, and decaying step sizes — are critical to good performance, model-free L2O either cannot achieve good performance or require largely many training problems to discover classic operations from the scratch. To avoid these shortcomings, we consider incorporating the existing methods as base or starting points for learning, which reduce the search to fewer parameters and a smaller space of algorithms. We call this alternative approach model-based L2O.

1.3 Broader Contexts

When the tasks are general machine learning tasks, e.g., determining the parameters of a prediction model by minimizing its training loss, L2O overlaps with meta-learning [20; 21; 22]. It is worth noting that “meta-learning” in different communities has different meanings. Here, meta-learning refers to using a method to improve learning algorithm(s), also called “learning to learn” [5; 6]. Recent results in this line of research contributes a significant portion of L2O development [23]. The goal of L2O captures two main aspects of meta learning: rapid learning within tasks, and transferable learning across many tasks from
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L2O shares many similarities with AutoML [24]. The term “AutoML” broadly refers to the automation of any step(s) in the ML lifecycle. Traditionally, AutoML research focuses on model selection, algorithm selection, and hyperparameter optimization. These methods accelerate the design iterations of many types of ML algorithms, such as random forests, gradient boosting, and neural networks. AutoML recently draws (back) the mainstream attention because of its significant success in enhancing the performance of deep learning [25]. Among the topics under AutoML, those most relevant to L2O are algorithm configuration [26] and hyperparameter optimization (HPO) [27]. Algorithm configuration determines a high-performing configuration of some algorithm across a given set of problem instances. HPO tunes a set of hyperparameters specifically for an ML algorithm, via Bayesian optimization [28] or classification-based optimization [29]. [30] has combined algorithm selection and hyperparameter optimization, also known as CASH. The main difference between these works and L2O lies in that L2O can discover new optimization methods from a parameterized algorithmic space of optimizers, rather than only selecting from or tuning a few existing optimizers. Yet, the boundary between HPO and L2O is often blurry since certain L2O methods [2; 31; 32] configure or predict hyperparameters for existing optimizers only.

L2O is closely related to the new frontier of learning-augmented algorithms [33]. Classic algorithms are designed with worst-case performance guarantees in mind and do not adapt to actual inputs. On the other hand, ML algorithms often achieve competitive performance by adapting to inputs, but their worst-case performance on (unseen) inputs degrades significantly. Learning-augmented algorithms combine the best of both worlds, using ML to improve the performance of classic algorithms, by adapting their behaviors to the input distribution. Examples include learning-augmented data structures [34; 35], streaming and sketching algorithms [36; 37], online algorithms [38], error-correcting codes [39; 40], scheduling algorithms [41], approximation algorithms [42], and safeguarded learned algorithms [19]. L2O can be counted as a subset of those learning-augmented algorithms. A comprehensive list of relevant materials can be found in [33].

Previous review articles and differences There have been a number of review articles on meta learning [20; 21] and AutoML [24; 43; 44].

Paper [45] surveys algorithm unrolling. Work [46] reviews model-based deep learning. These articles have overlapping scopes with ours as algorithm unrolling is a main (but not the only) technique of model-based L2O. This article features a more comprehensive coverage including both model-based and model-free L2O approaches.

Lastly, let us mention recent works that leverage ML for combinatorial and discrete optimization [47; 48]. A comprehensive review is [49].

1.4 Paper Scope and Organization

We draw distinctions between model-free and model based L2O approaches and review many L2O techniques. Emphases are given to recurrent network-based L2O methods, algorithm unrolling, and plug-and-play. We discuss how to train them effectively and how their designs can benefit from the structures of the optimizees and classic optimization methods.

We benchmark existing model-based and model-based L2O approaches on a few representative optimization problems. We have released our software implementation and test
cases as the Open-L2O package at [https://github.com/VITA-Group/Open-L2O], in the spirit of reproducible research and fair benchmarking.

The rest of the article is organized as follows. Section 2 and Section 3 review and categorize the existing works in model-free and model-based L2O, respectively. Section 4 describes the Open-L2O testbeds, our comparison experiments, and our result analyses. The article is concluded by Section 5 with a discussion on research gaps and future work.

2. Model-Free L2O Approaches

A model-free L2O approach in general aims to learn a parameterized update rule of optimization without taking the form of any analytic update (except the updates being iterative). The recent mainstream works in this vein [5; 6; 50; 51; 14] leverage recurrent neural networks (RNNs), most of which use the long short-term memory (LSTM) architecture. An LSTM is unrolled to perform iterative updates and trained to find short optimization trajectories. One set of parameters are shared across all the unrolled steps. At each step, the LSTM takes as input the optimizee’s local states (such as zero-order and first-order information) and returns the next iterate.

Model-free L2O shares the same high-level workflow depicted in Figure 1 (b). It is divided into two stages: an offline L2O training stage that learns the optimizer with a set of optimizees sampled from the task distribution $T$; then, an online L2O testing stage that applies the learned optimizer to new optimizees, assuming they are samples from the same task distribution. Table 1 compares a few recent, representative works in this area, from multiple lens including what optimizer architecture to select, what objectives and metrics that meta training and meta testing each use, and what input feature and other techniques are adopted. There are a great variety.

| Paper | Optimizer Architecture | Input Feature | Meta Training Objective | Additional Technique | Evaluation Metric |
|-------|------------------------|---------------|-------------------------|----------------------|------------------|
| 5     | LSTM                   | Gradient      | Meta Loss               | Transform input gradient $\nabla$ into $\log(\nabla)$ and sign($\nabla$) | Training Loss    |
| 6     | LSTM                   | Objective Value | Objective Value         | N/A                  | Objective Value  |
| 50    | LSTM                   | Gradient      | Meta Loss               | Random Scaling       | Training Loss    |
| 51    | Hierarchical RNNs      | Scaled averaged gradients, relative log gradient magnitudes, relative log learning rate | Log Meta Loss | Gradient History Attention | Training Loss |
| 52    | MLP                    | Gradient      | Meta Loss               | Unbiased Gradient Estimators | Training Loss Testing Loss |
| 53    | RNN Controller         | Loss, Gradient | Meta Loss               | Coordinate Groups    | Training Loss    |
| 54    | Searched Mathematical Rule by Primitive Functions | Scaled averaged gradients | Meta Loss | N/A | Testing Accuracy |
| 14    | Multiple LSTMs         | Gradient, momentum, particle's velocity and attraction | Meta Loss and Entropy Regularizer | Sample- and Feature- Attention | Training Loss    |
| 55    | RNN                    | Input Images, Input Gradient | Meta Loss | N/A | Training Loss and Robust Test Accuracy |
| 56    | LSTM                   | Input Gradient | Meta Loss               | N/A                  | Training Loss and Robust Test Accuracy |

2.1 LSTM Optimizer for Continuous Minimization: Basic Idea and Variants

As the optimization process can be regarded as a trajectory of iterative updates, RNNs are one natural option with a good inductive bias to learn the update rule. The first pioneering
model-free L2O method [5] proposed to model the update rule implicitly by gradient descent. This first LSTM method for L2O has often been referred to as L2O-DM in literature.

At high-level, the learned optimizer (modeled by LSTM) is updated by the gradients from minimizing the loss induced by the optimizee, and optimizees are updated by the update rule predicted by the optimizer:

$$\mathcal{L}(\phi) = \mathbb{E}_{(\theta_0,f) \in T} \left[ \sum_{t=1}^{T} w_t f(\theta_t) \right]$$

where

$$\begin{bmatrix}
\theta_{t+1} \\
h_{t+1}
\end{bmatrix} = m(\nabla \theta_t, h_t, \phi)
$$

(2)

In Eq. 2, $f$ represents an optimizee (e.g. a neural network with its loss function), with $\theta_0$ denoting the initialization of $f$. A set of optimizees and their initializations are sampled from the task distribution $T$. $\phi$ is the parameters of the L2O optimizer. $\nabla \theta_t = \nabla \theta f(\theta_t)$ is the gradient of the objective function with respect to the optimizee’s parameters. $m$ is the LSTM optimizer, and $g$ and $h$ are update and hidden state produced by $m$, respectively. $T$ is the maximal unrolling length for LSTM, often set due to the memory limitation; and $w_t = 1$ was used here for every $t$.

Two ad-hoc training techniques were utilized here. Firstly, each optimization variable (or called coordinate) of the optimizee’s parameter $\theta_t$ shared the same LSTM weights, but with different hidden states, in order to reduce the memory overhead faced when scaling up the number of optimization coordinates; this trick has been inherited by many LSTM-type L2O works. Secondly, to stabilize the optimizer’s training, the authors proposed to reduce the dynamic range of the optimizee’s gradient magnitudes, by preprocessing $\nabla \theta_t$ into $(\log(\nabla \theta_t), \text{sgn}(\nabla \theta_t))$ as the input into the optimizer.

The authors of [5] conducted a few proof-of-the-concept studies on small-scale tasks such as MNIST classification, where they showed the learned optimizer $m$ can converge faster than some stochastic gradient descent based optimizers such as SGD, RMSprop, and Adam. Despite this initial success, the scalability and generalizability of [5] remain underwhelming. Two specific questionable shortcomings are:

- **Generalizability of learned optimizer to unseen and potentially more complicated optimizees.** The L2O training set contains sampled optimizees from the target distribution. Just like typical machine learning models, we might expect a learned optimizer to both interpolate and extrapolate from the seen (meta-)training set, the latter being more challenging. Taking training deep networks for one example, during meta-testing, we might expect the learned optimizer to generalize (extrapolate) to the training of deeper or wider networks beyond instances seen in the meta training set. This generalizability is demanded also due to the memory bottleneck during meta training. To update the L2O optimizer $m(\phi)$ using back-propagation, we need to keep the gradients and the computation graph of the optimizee in memory. Therefore, the memory bottleneck arises when the unroll length of the $m(\phi)$ becomes large, or the dimension of optimizee’s parameter $\theta$ is high. Other types of generalization, such as training a network with unseen different activation functions or loss functions, are also found to be desirable yet challenging [5].

Moreover, applying the L2O optimizers to train more complex neural networks may meet more sophisticated loss landscapes, which cannot be well learned from observing simple training dynamics from small networks. For example, it is well known that training over-parameterized networks will exhibit sharply distinct behaviors at small
and large learning rates, and the two regimes usually co-exist in practical training and are separated by a phase transition [57; 58; 59].

- **Generalizability of learned optimizer to longer training iterations.** Training larger models naturally takes more iterations, which calls for longer-term L2O modeling. An LSTM model can in principle characterize longer-term dependency by unfolding more time steps (i.e., longer unrolling length $T$). However, that often results in L2O training instability due to the optimization artifacts of LSTM such as gradient explosion or vanishing, in addition to the practical memory bottleneck. Therefore, most LSTM-based L2O methods [5; 6; 50; 51; 52; 14; 15; 60] are forced to limit their maximal unroll lengths and to truncate the unrolled optimization (e.g., up to 20 steps). As a result, the entire optimization trajectory is divided into consecutive shorter pieces, where each piece is optimized by applying a truncated LSTM.

However, choosing the unrolling (or division) length faces a well-known dilemma [50]: on one hand, a short-truncated LSTM can result in premature termination of the iterative solution. Naively unfolding an LSTM-based L2O optimizer (trained with a small $T$) to more time stamps at meta-testing usually yields unsatisfactory results. The resulting “truncation bias” causes learned optimizers to exhibit instability and yield poor-quality solutions when applied to training optimizees.

The above research gaps motivate several lines of follow-up works, as summarized below:

**Debiasing LSTM Truncation** As explained above, selecting the unrolling length $T$ reflects a fundamental dilemma in LSTM-type L2O models: a longer $T$ causes optimization difficulty at L2O training, while a smaller $T$ incurs larger “truncation bias” and hampers the generalization at meta-testing.

![Figure 3: The dilemma of longer unrolling in both L2O testing and L2O training: (1) generalization failure at L2O testing: the upper figure shows the meta-testing loss of the vanilla L2O method [5] to quickly diverge, as we increase the number of steps at the meta-testing. This failure case was also observed in Figure 2 of [51] and in Figure 2 of [50]. (2) optimization failure at L2O training: the bottom figure shows the L2O training loss to also not decrease when it adopts longer unrolling lengths, due to the optimization artifacts of LSTM such as gradient explosion or vanishing.](image)

Many LSTM-type works focus on addressing this truncation bias. [51] drew the unroll length of their truncated optimization from a heavy tailed distribution. [52] proposed to replace LSTM with just MLP for the optimizer. The authors of [52] proposed to smooth
the loss landscape by using two unbiased gradient estimators with dynamic re-weighting:

\[ g_{rp} = \frac{1}{S} \sum_{s=1}^{S} \nabla_{\theta} f(\theta + n_s), \quad n_1, \cdots, n_S \sim N(0, \sigma^2 I) \text{ i.i.d.} \]  

(3)

\[ g_{es} = \frac{1}{S} \sum_{s=1}^{S} f(\tilde{\theta}_s) \nabla_{\theta} \log(P(\tilde{\theta}_s)), \quad \tilde{\theta}_1, \cdots, \tilde{\theta}_S \sim N(\theta, \sigma^2 I) \text{ i.i.d.} \]  

(4)

\[ g_{merged} = \frac{g_{rp} \sigma_{rp}^{-2} + g_{es} \sigma_{es}^{-2}}{\sigma_{rp}^{-2} + \sigma_{es}^{-2}} \]  

(5)

\( g_{rp} \) and \( g_{es} \) are gradient estimated by “reparameterization trick” [61] and “evolutionary strategies” [62], and \( \sigma_{rp} \) and \( \sigma_{es} \) are empirical estimates of the variances of \( g_{rp} \) and \( g_{es} \), respectively. In this way, they demonstrated L2O can train convolutional network networks faster in wall-clock time compared to tuned first-order methods, with reduced test losses.

Stronger LSTM Architectures  Another direction explored is to introduce stronger RNN/LSTM architecture to L2O. Instead of using a single RNN layer, Wichrowska et al. [51] leveraged three RNN layers to learn the optimization in different scales. They organized three RNN layers in a hierarchical fashion, here we simply denote them as “bottom RNN”, “middle RNN”, “upper RNN”. The “bottom RNN” directly takes the scaled gradients from the optimizee as input. Given a specific training step, the “middle RNN” receives the average all hidden states from “bottom RNN” across the optimizee’s parameter coordinates, and outputs a bias term to the “bottom RNN”. Further, the “upper RNN” takes the averaged hidden states from “middle RNN” across a certain window of optimizee’s training steps. By using smaller hidden states in “bottom RNN”, this hierarchical design of L2O optimizer achieved lower memory and compute overhead, while achieving better generalization. We refer [51] as L2O-Scale in this article.

Improved L2O Training Techniques  In order to improve the generalization of the learned optimizer to both longer unrolling (i.e. longer optimization iterations) and unseen functions, [50] proposed two training tricks. The first one, called random scaling, could be viewed as a special “data augmentation” for L2O: a coordinate-wise scale factor \( c \) was randomly generated at each iteration to scale the parameters of the optimizee during L2O training: \( f_c(\theta) = f(c\theta) \). It was motivated by the observation that in many optimization problems such as the quadratic function \( f(\theta) = \lambda \| \theta \|^2 \), the ideal update rule should achieve the same minima under varying scaling \( \lambda \). The second trick was to add a convex term during L2O training, which was inspired by the proximal algorithms [63] and avoided large random updates when the L2O optimizer is under-trained. We refer [50] as L2O-RNNprop in this article.

Lately, the authors of [64] took a deeper dive into improved training techniques for L2O models. The authors first presented a progressive training scheme, which gradually increased the optimizer unroll length to mitigate the L2O dilemma of truncation bias (shorter unrolling) versus gradient explosion (longer unrolling). Furthermore, they presented an off-policy imitation learning approach to guide the L2O training, by forcing the L2O optimizer to mimic the update rules generated by analytic optimizers. The authors of [64] evaluated their improved training techniques with a variety of state-of-the-art L2O models [5; 50; 51], and achieved boosted performance (lower training losses of unseen optimizees)
without changing the original L2O RNN architecture in each method. We refer [64] as L2O-enhanced in this article.

2.2 Other Common Implementations for Mode-Free L2O

While LSTM is so far the mainstream model, other optimizer models have also been explored. We describe two alternatives: reinforcement learning (RL), and neural symbolics. [53] proposed to learn an RL policy $\pi$ to predict the update rule, as the learned optimizer. The policy $\pi$ samples the update steps from a Gaussian distribution. The mean and variance of the Gaussian distribution are learnable parameters of the L2O policy, updated by reinforcement learning. The observation of the policy is composed of the current value of objective function (i.e. loss), the recent gradients, and changes of the objective values up to $H$ steps ($H = 25$ in their experiments). The policy receives the decrease of training loss as the reward. Logistic regression functions and two-layer neural nets were leveraged as the testbeds. Further on, [23] proposed to group coordinates under permutation invariance (e.g., weight matrix or a bias vector) into a coordinate group. This formulation reduced the computation cost of expensive optimization problems, making the proposed method extensible to wider neural networks. Generalizability of the learned policy was also studied by training the policy on shallow and narrow networks, and test on wider layers.

It is worthy to mention a special L2O work [54], that explored L2O from the neural symbolic prospective. The authors also leveraged an RL controller, but avoided directly modeling the update rules. Instead, they designed a search space of operands (gradient $g$, gradient’s running exponential moving average $\hat{m}$, etc.), unary functions of input $x$ ($e^x$, $\log(x)$, $\sqrt{|x|}$, etc.), and binary functions (mapping $(x,y)$ to $x+y$, $x-y$, $x\cdot y$, etc.). The RL controller was learned to select a sequence of elements from the search space, formulate a function to process the input, and output the update rule. Their searched best L2O optimizer (the RL controller) shows strong transferability, and improved training performance (lower training losses) on different tasks and architectures, including ImageNet classification and Google’s neural machine translation system.

2.3 More Optimization Tasks for Model-Free L2O

Black-box Optimization [6] pioneered to extend the LSTM L2O framework [5] to derivative-free or black-box function optimization. Due to the absence of the optimizee’s gradients as input, the authors of [6] instead treated the optimizee’s input-output pair as the observation of the optimizer, and formulated the optimization as an exploration-exploitation trade-off problem. They updated the optimizer’s hidden state $h_t$ by the observation from the last step $(x_{t-1},y_{t-1})$ and then chose a new query point $x_t$ to explore

$$h_t, x_t = \text{RNN}(h_{t-1}, x_{t-1}, y_{t-1}) \quad (6)$$

$$y_t \sim p(y | x_t, \ldots, x_1) \quad (7)$$

where the function value $y_t$ was incrementally sampled from a Gaussian Process ($p$) at each query point $x_t$ in the experiments. During L2O training, it was assumed that the derivatives of function value $y_t$ can be computed with respect to the input $x_t$, which means the errors will be backpropagated for L2O training, but not needed at L2O testing time. [6] demonstrated that their learned RNN optimizers are competitive with state-of-the-art Bayesian optimization packages (Spearmint [65], SMAC [26], and TPE [66]).
Particle Swarm Optimization  Current L2O methods mostly learn in the space of continuous optimization algorithms that are point-based and uncertainty unaware. Inspired by population-based algorithms (e.g. swarm optimization), [14] estimated the posterior directly over the global optimum and used an uncertainty measure to help guide the learning process. The authors designed a novel architecture where a population of LSTMs jointly learned iterative update formula for a population of samples (or a swarm of particles). The model can take as input both point-based input features, such as gradient momentum; and population-based features, such as particle’s velocity and attraction from swarm algorithms. To balance exploration and exploitation in search, the authors directly estimated the posterior over the optimum and included in the meta-loss function the differential entropy of the posterior. Furthermore, they learn feature- and sample-level importance reweighting (often called “attention” in deep learning) in the L2O model, for more interpretable learned optimization rules. Their empirical results over non-convex test functions and the protein-docking application demonstrated that this new L2O largely outperforms the off-the-shelf Particle Swarm Optimization (PSO) algorithm [67] and the vanilla LSTM-based L2O methods that are not uncertainty-aware [5]. We refer [14] as L2O-Swarm in this article.

Minimax Optimization  A latest challenging testbed for model-free L2O is to solve continuous minimax optimization, that is of extensive practical interest [68; 69; 70], yet notoriously unstable and difficult. Three prior works [55; 71; 56] tried to plug in L2O into a specific application of minimax optimization called adversarial training [69]:

$$\min_{\theta} \mathbb{E}_{(x, y) \sim D} \left\{ \max_{x' \in \mathbb{B}(x, \epsilon)} \mathcal{L} \left( f \left( x' \right), y \right) \right\},$$

(8)

where $D$ is the empirical distribution of input data, and the inner maximization is defined as the worst-case loss within a small neighborhood $\mathbb{B}(x, \epsilon)$. In both works, the L2O only predicted update directions for the inner maximization problem, while the outer minimization was still solved by classic optimizer. They empirically showed that L2O can improve the solution quality to the inner maximization optimization, hence also leading to a better minimax solution and a more robustly trained model.

A latest work [72] extended L2O to solve minimax optimization from end to end, for the first time. The authors proposed Twin L2O, consisting of two LSTMs for updating min and max variables, respectively. This decoupled design was shown by ablation experiments to facilitate learning, particularly when the min and max variables are highly non-symmetric. Several enhanced variants and training techniques were also discussed. The authors benchmarked their L2O algorithm on several relatively basic and low-dimensional test problems, and on which L2O compared favorably against state-of-the-art minimax solvers. How to scale up L2O for fully solving minimax problems of practical interest, such as adversarial training or GANs, remained to be an open challenge.

3. Model-Based L2O Approaches

We now overview model-based L2O approaches, which mark the recent movement to fuse traditional model-based optimization algorithms with powerful deep learning architectures [45]. Rather than use general-purpose LSTMs, these methods model their iterative update rules through a learnable architecture that is inspired by analytic optimization algorithms. Often, these learned methods can approximate problem solutions with tens of iterations whereas their classic counterparts make require hundreds or thousands of iterations [2].
At a high level, model-based L2O can be viewed as a “semi-parametric” option that takes advantage of both model-based structures/priors and data-driven learning capacity. The growing popularity of this approach lies in its demonstrated effectiveness in developing compact, data-efficient, interpretable and high-performance architectures, when the underlying optimization model is assumed available or can be partially inferred. There is a large design space to flexibly balance between the two. Most model-based L2O methods take one of the following mainstream approaches.

The first approach is known as plug and play (PnP), and the key idea here is to plug a previously trained neural network (NN) into part of the update for an optimization algorithm (i.e., in place of an analytic expression), and then play by immediately applying the modified algorithm to problems of interest (without additional training). We illustrate this with the common alternating direction method of multipliers (ADMM) formulation of PnP (e.g., see [73]). Here the underlying task of the original optimization algorithm is to minimize the sum of two functions $f$ and $g$, using successive proximal operations. The PnP formulation replaces the proximal for $g$ with an operator $H_\theta$ to obtain the iteration:

\begin{align}
    x^{k+1} &= H_\theta(y^k - u^k) \\
    y^{k+1} &= \text{prox}_{\alpha f}(x^{k+1} + u^k) \\
    u^{k+1} &= u^k + x^{k+1} - y^{k+1}.
\end{align}

Before inserting $H_\theta$ into the iteration above, the parameters $\theta$ are learned independently as the solution to a training problem, i.e.,

$$\theta \in \arg\min \mathcal{L}(\tilde{\theta}).$$

The loss function $\mathcal{L}$ is designed by an independent goal (e.g., to learn a natural image denoising operator). For example, one might model an image recovery problem by using total variation (TV) as a regularizer [74]. In the optimization scheme chosen (e.g., ADMM) to recover the image, one step of the process could be to perform a proximal operation with TV, which effectively acts as a denoiser. The PnP framework proposes replacement of the TV proximal with an existing denoiser (e.g., a neural network [75] or BM3D [76]).

The second approach is known as algorithm unrolling, which unrolls a truncated optimization algorithm into the structure of a neural network [45]. Updates take the form

$$x^{k+1} = T(x^k; \theta^k), \quad k = 0, 1, 2, \ldots, K.$$

Upon establishing this form, the parameters are learned by an end-to-end approach:

$$\min_{\{\theta^k\}_{k=0}^{K-1}} \mathcal{L}(x^K(\{\theta^k\}_{k=0}^{K-1})), \quad \mathcal{L}$$

where $\mathcal{L}$ is the loss function we use in training. We emphasize the distinction that the parameters in unrolled schemes are trained end-to-end using the iterate $x^K$ as a function of each $\theta^k$ whereas training occurs separately for PnP. We will introduce how to design $\mathcal{L}$ later in this section. Below we discuss each of these approaches, their typical features, and which approach is suitable for various applications.
Learning to Optimize: A Primer and A Benchmark

Figure 4: A common approach in various L2O schemes is to form feed forward networks by unrolling an iterative algorithm, truncated to \( K \) iterations, and tuning parameters at each layer/iteration \( k \). This generalizes the update formula \( u^{k+1} = T(u^k; d) \) to include a dependence on weights \( \theta^k \), denoted by a subscript.

| Tunable Optimization Model | Opt | IP | PnP |
|----------------------------|-----|----|-----|
| Tunable Update Formulas    | ✓   | ✓  | ✓   |
| Training Loss tied to Model | ✓   |    |     |
| Training Loss measure \( u^* \) error | ✓   |    |     |
| Interpretable Updates     | S   | S  | ✓   |

Table 2: Comparison of properties for three types of model-based L2O methods: unrolled objective-based (Opt), inverse problem (IP), and Plug and Play (PnP) methods. Here ✓ and S mean that the corresponding property always and sometimes holds, respectively.

3.1 Plug and Play

PnP methods date back to 2013 when they were first proposed for ADMM [73]. The authors replaced one of the proximal operators in ADMM with a denoiser (e.g., K-SVD [77], BM3D [76], and non-local means [78]) and showed empirical performance surpassing the original ADMM algorithm. Several other PnP schemes were also introduced and their empirical success was shown in the literature [79; 80; 81; 82; 83; 84; 85; 86; 87; 88; 89; 90; 91; 92; 93; 94; 95]. The theoretical guarantees of convergence of PnP-ADMM was first provided in [96] under the assumption that the derivative of the denoiser was doubly stochastic. Then [97] proved the convergence of PnP-ADMM with the use of a more realistic “bounded denoisers” assumption. Some other PnP schemes were analyzed under different assumptions [98; 99; 100; 101; 102; 103; 104; 105; 106; 107; 108].

PnP was initially not connected to L2O, until some concurrent works [109; 110; 111] introduced the concept of “learning operators” into a PnP framework. Instead of using a manual-designed denoiser, they modeled the proximal operator as a deep neural network and learned it from data. The empirical performance of such an approach largely exceeded the prior PnP works. From an L2O perspective, the learned methods were able to improve performance by either accelerating execution of a subroutine in the optimization algorithm or providing a “better” solution than previously obtained.

Besides learning the operators in a PnP framework, one can also learn a functional as the regularizer. [112; 113] modeled the statistical distribution of nature images as a Denoising Autoencoder (DAE) and constructed the optimization objective by maximizing
the likelihood. The DAE is differentiable; thus, the inverse problem can be solved with gradient-based optimization algorithms. A parameterized discriminator function defined in the Wasserstein distance between two distributions was used by [114] as a learned a functional that discriminates between a ground-truth and fake images. The authors treated this learned functional as a regularizer and used it in a gradient descent algorithm for image inverse problems.

Learning is not only able to help find the denoiser/regularizer in PnP, but is also able to be used in a more meta manner to help find good parameters in PnP. [115] learned a policy network to tune the parameters in Plug-and-Play with reinforcement learning. By using the learned policy, the guided optimization can reach comparable results to the ones using oracle parameters tuned via the inaccessible ground truth.

The marriage of PnP with L2O also provides theoretical blessings. While a manually designed regularizer may not guarantee the convergence of PnP to the desired solution, a learning-based method provides the flexibility to meet the condition of convergence. [75] studied the convergence of some PnP frameworks using fixed-point iterations, showing guaranteed convergence under a certain Lipschitz condition on the denoisers. They then proposed a normalization method for training deep learning-based denoisers to satisfy the proposed Lipschitz condition. Similarly, [116] also provided an approach for building convergent PnP algorithms using monotone operator theory and constraining the Lipschitz constant of the denoiser during training. This fixed point framework was extended by [117] to the RED-PRO framework, which also showed convergence to global solutions. Independently of RED-PRO, [118] proposed a method within the RED-PRO framework for learning projection operators onto compact manifolds of true data. Utilizing assumptions about the manifold and sufficient representation by the sampled data, the authors proved their approach to constructing a PnP operator can provably approximate the projection (in probability) onto a low dimensional manifold of true data.

3.2 Algorithm Unrolling

Herein we overview L2O methods comprised of unrolling iterative optimization algorithms. We start by emphasizing there are two distinct goals of unrolled L2O methods: either to minimize an objective, or to recover a signal. The distinction is that the aim for inverse problems is to tune the parameters so that they minimize the reconstruction accuracy of some “true” signal rather than find a minimizer of the model’s objective function. The practicality for objective minimization is to speed up convergence. Keeping these categorizations in mind will help provide the reader intuitive lenses for looking at different approaches. See Table 2 for a comparison of common qualities in these methods and PnP.

Below we provide a comprehensive and organized review of existing works, along multiple dimensions: what problems they solve, what algorithms they unroll, what goals they pursue (objective minimization or signal recovery), and to what extent they freely parameterize their learnable update rule.

3.2.1 Different target problems

We broadly categorize the main problems tackled by model-based L2O in four categories: probabilistic graph models, sparse and low rank regression, differential equations, and quadratic optimization.
• **Sparse and low rank regression.** The most investigated problems in the literature of unrolling is probably the sparse regression, inspired by the first seminal work [4], which unrolls the Iterative Shrinkage Thresholding Algorithm (ISTA) or its block coordinate variant as a recurrent neural network to solve LASSO for fast sparse coding. The unrolling philosophy is also used in low rank regression as it shares some common nature with sparse regression.

  - **LASSO:** Most of unrolling works following [4] also solve LASSO-type optimization problems for sparse coding [119, 120, 121, 122, 18, 123, 124, 125, 126]. Beyond naive LASSO, a natural extension is to apply the same unrolling and truncating methodology to solve group LASSO [127], i.e. finding solutions with structured sparsity constraints.

  - **Other sparse/anti-sparse regression problems:** [128] and [129] unroll the iterative hard thresholding (IHT) algorithm that solves $\ell_0$ minimization problem instead of LASSO. Unrolling for $\ell_\infty$ minimization was also considered by [130], leading to so-called anti-sparse representation learning [131].

  - **Low Rank Regression:** [132; 133; 134; 135] all extend the unrolling idea to low-rank matrix factorization. Specifically, [134] proposes tailored pursuit architectures for both robust PCA and non-negative matrix factorization with specialized algorithms inspired by the non-convex optimization techniques to alleviate the expensive SVD involved in each iteration.

• **Probabilistic Graphical Model.** The unrolling method can also be adopted to solve probabilistic graphical models. For example, [132] interpret conventional networks as mean-field inference in Markov random fields and obtains new architectures by modeling the belief propagation algorithm to solve the Markov random field problems. [136] formulate Conditional Random Fields with Gaussian pairwise potentials and mean-field approximate inference as recurrent neural networks.

• **Differential equations.** Another line of works [137; 138; 139; 140] unroll the evolution in Partial Differential Equation (PDE) systems. While PDEs are commonly derived based on empirical observations, those recent advances offer new opportunities for data-driven discovery of (time-dependent) PDEs from observed dynamic data. One can train feed-forward or recurrent neural networks to approximate PDEs, with applications such as fluid simulation [141]. Loosely related is also significant work on connecting neural networks with Ordinary Differential Equation (ODE) systems.

• **Quadratic optimization.** Some recent works investigate the unrolling method in quadratic optimization problems [32, 142], which are easier to solve compared to the problems studied above. The focus here is more on the theoretical analysis of the convergence, and/or the interplay between the unrolled algorithm and the resultant deep model’s property (e.g., generalization and stability).

In some scenarios, we are not directly interested in the output of the unrolled model but use it for downstream tasks, e.g. clustering [143] and classification [128; 18, 125]. That will often lead to task-driven joint optimization and the end task output becomes the focus of evaluation, in place of the original unrolling algorithm’s output fidelity.
Inverse Problems  In many cases, however, optimization problems with manually designed objective functions only provide approximations to the original signals that we are really interested. This is often due to inexact prior knowledge about the original signals. For example, sparsity and total variation regularizations only partially reflect the complexity of natural images (approximated sparsity and smoothness) which are hardly true in real-world applications and do not depict the exact characteristics of natural images. Therefore, many works solve the inverse problem directly, striving to recover the original signals that we are really interested in. We return to this task in a later subsection.

3.2.2 Different Algorithms Unrolled

For the bulk of iterative optimization algorithms that have been unrolled, we classify them into three categories: forward backward splitting, primal-dual methods, and others. The first two categories consist entirely of first-order algorithms, which is due to their low computational complexity and the fact their resultant L2O methods are often more reliable to train. We also emphasize that although various works discussed below may use the same underlying algorithm as the base, they can vary greatly in their performance due to choices regarding what parameters are learned and what safeguard precautions are used to ensure convergence (discussed further in subsequent sections). We also emphasize that the majority of these algorithms revolve around obtaining some form of sparsity/low rank solution.

We begin with forward backward splitting (FBS). The simplest learned scheme is Gradient Descent, which is where the the gradient descent operation is applied (i.e., the forward operator) and the backward operator is simply the identity. This class of methods is studied in many works (e.g., see [122; 87; 144; 145; 142; 146; 147; 148; 149; 137; 114; 150; 126; 151; 152]). However, the most popular focus in the literature is on problems with sparsity constraints, which are usually modeled by $\ell_1/\ell_0$-minimization. The former, also known as LASSO, can be solved by the iterative shrinkage-thresholding algorithm (ISTA) [153] and its variants. This has yielded great interest in L2O schemes, as evidenced by the fact the works [4; 143; 127; 8; 9; 1; 2; 154; 123; 155; 156; 124; 157; 158], among others, provide various ways to unroll the original ISTA algorithm. Additionally, [120; 121; 159; 157] unroll a Nesterov accelerated ISTA known as FISTA (Fast ISTA) [160]. Continuing in the vein of sparsity, [161; 12; 162] unroll another algorithm, called approximate message passing (AMP), which introduces Onsager correction terms that whitens the noise in intermediate signals while [163; 155; 154] unroll Orthogonal AMP [164], an extension to the original AMP. For the $\ell_0$-minimization problems, the iterative hard-thresholding (IHT) algorithm, which replaces the soft-thresholding function in ISTA with hard-thresholding, is unrolled in [128; 129]. Switching gears, for problems that involve minimize an objective that is the sum of several terms, the incremental proximal gradient method has also been unrolled [165]. Further generalization of FBS is given by an abstract collection of flexible iterative modularization algorithms (FIMAs) [166], that perform updates using the composition of two learned operators (provided the composition monotonically decreases an energy). A caveat of this particular approach is that the learned operator is also composed with a classic proximal-gradient operation.

The next category of unrolled L2O algorithms consists of primal-dual schemes. These typically include variations of primal-dual hybrid gradient (PDHG) and the alternating direction method of multipliers (ADMM) (e.g., see [119; 167; 110; 147; 168; 125; 169]). An-
other variation used is the *Condat-Vu primal-dual hybrid gradient* [170]. As above, many of these methods also focus on leveraging the sparse structure of data.

The remaining group of miscellaneous methods take various forms. These include another first-order algorithm, *Half Quadratic Splitting* (HQS), for image restoration [111; 88]. Using a substitution and soft constraint, the HQS approach solves a problem using a model that approximates variational problems (VPs). Beyond first-order algorithms, unrolling is also applied to *second-order (Newton or Quasi-Newton) methods* [171; 172], *Differential Equations* [138; 173; 174] and *Interior Point method* [10; 175], and *Frank-Wolfe algorithm* [176]. *Conjugate Gradient* was proposed [177] to help yield a more accurate enforcement of the data-consistency constraint at each iteration than comparable proximal-gradient methods. Lastly, [178] propose an unfolded version of a greedy pursuit algorithm, i.e., *orthogonal matching pursuit* (OMP), that directly targets at the original combinatorial sparse selection problem. Their methods called Learned Greedy Method (LGM) can accommodate a dynamic number of unfolded layers, and a stopping mechanism based on representation error, both adapted to the input. Besides, there are also L2O schemes that appear to be based entirely on heuristic combinations of classic optimization methods (e.g., see [179]).

### 3.2.3 Objective-Based v.s. Inverse Problems

**Objective Based.** The simplest L2O unrolling scheme is objective based. Training is applied here to yield rapid convergence for a particular distribution of data. The training loss can take various forms, including minimizing the expectation of an objective function [123], the objective’s gradient norm, the distance to the optimal solution, or the fixed point residual of the algorithm [19] (e.g., if \( T \) is the update operator, then \( \| x - T(x) \| \) is the fixed point residual). Examples of objectives that have been extensively studied in the literature are presented in Section 3.2.1.

In addition, *safeguarding* can be used in this situation, for guiding learned updates to ensure convergence [180; 19]. This can be accomplished in multiple ways. A typical approach is to generate a tentative update using an L2O scheme and then check whether the tentative update satisfies some form of descent inequality (e.g., yields a lesser energy value or fixed point residual). If descent is obtained, then the tentative update is used; otherwise, a classic optimization update is used. These safeguarded schemes provide the benefit of reducing computational costs via L2O machinery while maintaining theoretical convergence guarantees.

**Inverse Problems** Several L2O methods attempt to solve inverse problems (IPs) (e.g., see [181; 179; 170; 182; 152; 183; 166; 150]). Here the task is to reconstruct a signal \( x^* \) from indirect noisy measurements. The measurements are typically expressed by \( d \in \mathbb{R}^m \) and are related to a forward operator \( A : \mathbb{R}^n \rightarrow \mathbb{R}^m \) by

\[
d = A(x^*) + \varepsilon,
\]

where \( \varepsilon \) is noise. A few repeated themes arise the L2O IP literature. The typical process is to i) set up a variational problem (VP) as a surrogate model, ii) choose a parameterized optimization algorithm that can solve \( \varepsilon \) (VP), and iii) perform supervised learning to identify the optimal parameter settings. We expound upon these themes and their nuances below.

---

2. We mean to say that, for some choice of parameters, the algorithms solves (VP).
First, by creating a variational problem, one assumes \( u^* \) approximately solves
\[
\min_{x \in \mathbb{R}^n} \ell(d, A(x)) + J(x), \tag{VP}
\]
where \( \ell: \mathbb{R}^m \times \mathbb{R}^m \to \mathbb{R} \) is a fidelity term that encourages the estimate \( u \) to be consistent with the measurements \( d \) and \( J: \mathbb{R}^n \to \mathbb{R} \) is a regularizer. Learning comes into play since \( u^* \) is usually not the solution to (VP), but instead some “close” variant. Thus, upon choosing the form of \( \ell \) and \( J \), one includes tunable parameters in the model and unrolls a parameterized optimization scheme for (typically) a fixed number \( K \) of iterations. Algorithm updates can be parameterized beyond their classic form (e.g., replace a fixed matrix with a matrix that has tunable entries as done in \([4]\)). In most cases, the training loss takes the form of minimizing the expected value of the square of the Euclidean distance between the output estimate \( u^K \) and the true signal \( u^* \), i.e., the parameters \( \Theta \) are trained to solve
\[
\min_{\Theta} \mathbb{E}_{d \sim D} \left[ \| x^K(\Theta, d) - x^*_d \|_2^2 \right]. \tag{14}
\]

The primary alternative training loss for L2O is to use estimates of the Wasserstein-1 distance between the distribution of reconstructed signals and the distribution of true signals (e.g., see \([114]\)), which yields unsupervised training.

### 3.2.4 Learned Parameter Roles

A key aspect of unrolled L2O methods is to determine how to parameterize each update. This subsection discusses some of the common considerations and roles for these parameters.

**Learning parameters in the iterations** First, the direct method of parameterization is to convert scalars/vectors/matrix/filters used in iterative algorithms into learnable parameters and learn them through data-driven training process. In this type of parameterization, learning can overcome the need to hand choose hyperparameters. For example, LISTA \([4]\) unrolls ISTA, which usually has update formulation
\[
x^{k+1} = \eta_{\lambda/L} \left( x^k - \frac{1}{L} A^T (Ax^k - d) \right), \tag{15}
\]
where \( \lambda \) is the coefficient before the \( \ell_1 \) regularization in LASSO and \( L \) is the largest eigenvalue of \( A^T A \). Then LISTA parameterizes ISTA as a recurrent formulation
\[
x^{k+1} = \eta_{\theta} \left( W_e d + S x^k \right), \tag{16}
\]
where taking \( W_e \equiv A^T / L, S \equiv I - A^T A / L \) and \( \theta \equiv \lambda / L \) reduces (16) to ISTA. A large amount of unrolling works follows the same methodology, but differ from each other in specific perspectives during the parameterization process, e.g. drop the recurrent structure in \([4]\) and use feed-forward modeling by untying the learnable parameters in different layers and update them independently \([132; 161]\). This untying formulation can enlarge the capacity of the unrolled model \([132]\) but can also cause the training instability due to the overwhelming parameter space and the difficulty of theoretical analysis on the convergence and other properties of the unrolled model.

To this end, there has been a line of efforts that strive to reduce the number of trainable parameters by relating and coupling different parts of parameterization \([129; 1]\) and
analytically generate parameters that satisfy certain constraints \[2\]. ALISTA \[2\] learns thresholding and step size parameters, reducing the number of parameters significantly – to two scalars per layer, which stabilizes the data-driven process while achieving state-of-the-art recovery performance at the time with theoretical convergence guarantee.

Besides, recent trends also started to apply model-free L2O methods, to predict hyperparameters in classic iterative optimization algorithms. This is seen as the fusion among model-free L2O (since the algorithmic techniques are LSTM- or RL-based), model-specific L2O (as the update rule is eventually based on the classic iterates), and more traditional hyperparameter optimization. Examples include learning an adaptive learning rate schedule for training deep networks with stochastic gradient descent \[31; 184; 185\]; coordinating specifically for layer-wise training \[186\] or domain adaptation speeds \[17\]; predicting the update combination weights and decay rates in Adam \[187\]; or estimating training sample importance \[185; 188\], among others.

**Deep priors learned from data** Another popular parameterization in unrolling is to use a deep model to learn a data-dependent prior on the interested signals to replace hand-designed priors such as sparsity and total variation, which are not accurate in real-world applications and thus introduce bias. However, previous works have various ways to use the “learned prior”. For example in \[110; 162; 87; 111\], people used data-driven training to learn a proximal or a projection operator that is iteratively used in the unrolled algorithm. The learned operator takes recovered signals contaminated by noises and artifacts as inputs and outputs a refined estimation. Sometimes the learned operator is found to fit a denoiser. The main difference of these learned prior methods from Plug-and-Play methods is that the operator is usually learned in a end-to-end way, making it overfitting to the current task or data distribution and not be able to be plugged into other algorithms. \[189; 190\] supposed that the relevant vectors lie near the range of a generative model, and hence used generative adversarial networks (GANs) as a learning-based prior for image compressive sensing. \[114; 150\] perceive the prior as a loss function that can distinguish between coarsely recovered signals without considering any priors, and real on-domain signals with good quality. The prior as a loss function is adversarially trained to output high losses for coarse recoveries and low losses for real signals. Then we use the (sub-)gradient generative by the learned network via back-propagation in the unrolled algorithms e.g. gradient descent.

**Others** Besides the above two major ways of parameterization, there are also works that learn black-box agents that directly generate next-step estimations given the current iterate and historical information \[146; 171; 149\]. For instance, \[158\] learns an LSTM network that generates the step sizes and threshold parameters within each layer in ALISTA, instead of training them using back-propagation.

### 3.3 Applications

Unrolled L2O schemes have found many applications. Below we identify several of these and note that there is some overlap among the three approaches discussed in this section.

- **Image Restoration and Reconstruction.** The model-based L2O methods, including both Plug-and-Play and unrolling methods, are widely used for various tasks in image restoration, enhancement, and reconstruction. Popular application examples include denoising \[144; 149; 111; 137; 191; 2; 168; 114; 192\], deblurring \[144; 111; 109; 193; 10; 192; 150; 91; 194\], super-resolution \[110; 122; 149; 111; 137; 193; 91\], inpainting
and compressive sensing [110; 162; 1; 144; 121; 195; 9; 155; 192], JPEG artifacts reduction [8; 137; 196], demosaicking [109], dehazing [88] and deraining [197]. Note that not all those works identically stick to the algorithm’s original architecture; instead many only follow loosely the idea, and replace various components with convolutions or other modern deep learning building blocks.

**Medical and Biological Imaging.** We specifically separate Medical and Biology Imaging applications from the previous Image Restoration part because the former has its own special scenario and challenges. Imaging techniques in medical such as MRI and CT require accurate reconstructions of images with as few measurements as possible that result in minimal discomfort or side-effect. It is also challenging to extend the model-based methods in natural images to properly deal with the complex-valued inputs [167]. Other work that applies model-based L2O to medical and biology imaging includes [147; 198; 146; 144; 114; 199; 150; 124; 200].

**Wireless Communication.** Tasks in wireless communication systems can also be solved by unrolling methods, e.g. resource management [201; 154; 202], channel estimation [163], signal detection [163] and LDPC coding [151]. For example, MIMO detection, which can be formulated as a sparse recovery problem, was shown to benefit from L2O methods based on “deep unfolding” of an iterative algorithm added with trainable parameters [13], such as LISTA. The model-based L2O approaches have already exhibited superior robustness and stability to low signal-to-noise (SNR), channel correlation, modulation symbol and MIMO configuration mismatches [163]. We refer the readers to a seminal survey about unfolding methods in communication systems [13].

**Seismic Imaging.** Another important application of model-based L2O is seismic imaging [203; 204; 205; 206]. Most of them adopt a more plug-and-play manner to learn CNN-based projectors that are first trained using data and then integrated into classic iterative updates, due to the desirable emphasis on the physical modeling.

**Miscellaneous Applications:** such as clustering [143; 207; 208] and classification [128; 18; 125], phase retrieval [209], RNA second structure prediction [15], speech recognition and source separation [132; 210], remote sensing [211], smart grid [212], graph recovery [213], and photometric stereo estimation [129].

### 3.4 Theoretical Efforts

Although successful empirical results show significant potential of L2O, limited theory exists due to black-box training pertaining to such learned optimizers. That important gap often makes the broader usability of L2O questionable. We note that, the specific optimization problem and algorithm structure in model-based L2O often offer more opportunities for their theoretical analysis, by bridging us to the wealth of classic optimization tools, compared to model-free L2O. Therefore, model-based L2O in a few specific problems has been the main focus of existing theoretical efforts so far.

To explain why and how model-based L2O methods outperform traditional optimization algorithms, there are several essential questions at the heart of model-based L2O:

- **(Capacity).** Given the model of L2O, do there exist parameters in the model that makes L2O provably outperform traditional optimization algorithms, over the task
distribution (i.e., “in distribution”)? Is there any “safeguard” mechanism available on L2O to ensure them at least as good as traditional algorithms even on examples out of the task distribution (i.e., “out of distribution” or OoD)?

- (Trainability). Given the existence of such parameters, what training method should we use to obtain those parameters? Do guarantees exist that the training method converges to the ideal parameters?

- (Generalization). Does the trained model generalize, say, to testing instances from the same source of training instances (i.e., “interpolation”)? Can the trained models “extrapolate”, e.g., on testing instances more complicated than any training one?

- (Interpretability). How can we explain what the L2O models have learned?

We list brief answers to the four questions here. Capacity and interpretability have been partially solved on signal/image-processing related problems by some recent works, as to be detailed in the subsections below. Generalization gains increasing attention recently and some works provide bounds of generalization gap of some specific L2O models. Lastly, to the best of our knowledge, there has been very limited theoretical work on the trainability of L2O, due to the high nonconvexity of the training objectives (see Section 3.2.3). The main exception here is the recent work [32] discussing the local minima and gradient explosion/vanishing in L2O training for quadratic minimization. This was also partially addressed by [118], where the authors used sorting activation functions [214] and Lipschitz networks to encourage gradient norm preservation.

### 3.4.1 Capacity

To the best of our knowledge, [201] is the first effort on theories of L2O. It approximates a traditional method WMMSE with a full-connected neural network and proves that the output of the neural network can be arbitrarily close to the result of WMMSE as long as the number of layers and units in the network is large enough. In other words, this work estimates the approximation capacity of the neural network.

Approximation capacity is a generic notion in deep learning study. Specifically for L2O, convergence can be used to describe the model capacity: Do there exist parameters \( \{\theta^k\}_k \) in (11) that make \( \{x^k\} \) converge better than classic optimization algorithms as \( k \to \infty \)? The work [1] adopts this way to describe the convergence of L2O on the sparse recovery problem and give a convergence rate of LISTA which is better than that of classic algorithms ISTA/FISTA. [2; 215; 157; 123; 216; 217] improve the theoretical result of LISTA by designing delicate models, in another word, designing operator \( T \) in (11). [168] analyzes the convergence rate of differentiable linearized ADMM.

Instead of studying parameters \( \theta^k \), another approach to establish convergence is to propose mathematical conditions on the operators (for example, operator \( T \) in (11) and operator \( H \) in (9a)) in the models. These mathematical conditions should not only guarantee the convergence but also can be satisfied practically. [75] proposes a continuity condition that can be satisfied by specifically designed training method. [218] assumes smoothness of the operator that is satisfied by choosing smooth activation functions in a neural network (e.g., the sigmoid function). [150] assumes convexity of the regularizer in their math proof and proposes to parameterize the regularizer by a convex neural network. [118] proves the convergence under assumptions about the manifold and sufficient representation by the sampled data, which are usually satisfied in practice.
While the above efforts focus on studying the convergence and acceleration effect of L2O over the target task distribution, a parallel and same important topic is to bound or characterize the convergence of L2O under OoD inputs: how much the L2O convergence can degrade when applied to optimizees deviating from the task distribution? Seemingly daunting at the first glance, that goal may be fulfilled by L2O with a safeguard mechanism, that can provide a way to establish convergence independent of the parameters and data. The most common approach is to i) compute a tentative L2O update using any method under the sun, ii) check if the tentative update yields a reduction in the value of an energy, and iii) accept the L2O update if the energy is less than some relative bound (e.g., the energy at the current iterate). If the tentative L2O update is rejected, then a fallback scheme is applies the update of a classic optimization algorithm. Because the energy is monotonically decreasing, it can be shown that the overall algorithm converges to a minimizer of the energy. The energy is typically defined to be the objective function in the variational problem (VP) or its differential (e.g., see [193; 219; 166; 180]). An alternative approach is define the energy to measure the residual between updates (e.g., [19]). That is, if $T$ is the update operator for a classic algorithm, then at a tentative update $u^k$ we check if the residual $\|u^k - T(u^k)\|$ is less than some relevant bound.

3.4.2 Interpretability

Interpretability is significant to a learning model, now even more than ever. Some efforts have been made on the interpretability of L2O, mainly about linking or reducing their behaviors to those of some analytic, better understood optimization algorithms. [129] studies unrolled iterative hard-thresholding (IHT) and points out that unrolled IHT adopts better dictionary and a wider range of RIP condition than IHT. [120] demonstrates that the mechanism of LISTA is related to a specific matrix factorization of the Gram matrix of the dictionary. [122] explains the success of LISTA with a tradeoff between convergence speed and reconstruction accuracy. [2] shows that the weights in LISTA has low coherence with the dictionary and proposes an analytic approach to calculate the weights. [213] analyzes alternating minimization algorithm quantitatively on a graph recovery problem and reveals the analogy between the learned L2O model and solving a sequence of adaptive convex programs iteratively. [145] points out the analogy between deep-unfolded gradient descent and gradient descent with Chebyshev step-size and shows that the learned step size of deep-unfolded gradient descent can be qualitatively reproduced by Chebyshev step-size.

3.4.3 Generalization

Generalization is an important topic for L2O, just like for any other ML domain. The relationship between generalization gap and the number of training instances provides us an estimate on how many samples we should use. Both [142] and [220] studied the Rademacher complexity, an upper bound of generalization gap. [142] estimates the Rademacher complexity of gradient descent and Nesterov’s accelerated gradients on a parametrized quadratic optimization problem; [220] estimates the Rademacher complexity of a deep thresholding network on sparse linear representation problem.

Stability is highly related with generalization [142], some L2O works analyze the stability of their methods. For example, [182] estimates Lipschitz constants of their method to provide stability analysis results both with respect to input signals and network parameters. This yields a growth condition inequality. Another measure of stability is to ensure stability
of the model as the weighting of the regularizer term in (VP) tends to zero \cite{152, 183}. A third approach is that of Φ-regularization used with null space networks \cite{221}, which learns a regularizer based on the null space of A. Each of these methods yields different insights about the stability of L2O schemes for inverse problems.

4. The Open-L2O Benchmark

The community has made diverse attempts to explore L2O and generated a rich literature for solving different optimization tasks on different data or problems, as well as different software implementations on various platforms. Each L2O method has its own training recipe and hyperparameters. Because a common benchmark for the field has not yet been established, comparisons among different L2O methods are inconsistent and sometimes unfair. In this section, we present our efforts toward creating a comprehensive benchmark that enables fair comparisons. To the best of our knowledge, this is the first time of such an attempt.

**Testbed problems:** We choose some popular and representative test problems that have been used in the existing L2O literature: (i) convex sparse optimization, including both sparse inverse problems and LASSO minimization; (ii) minimizing the nonconvex Rastrigin function, and (iii) training neural networks (NNs), which is a more challenging nonconvex minimization problem.

**Task distributions:** Inspired by both research practice and real-world demands, we define task distributions in the following problem-specific way: (i) for sparse optimization, the optimizees during training and testing are optimization problems with the same objective function and decision variables but different data; (ii) in the Rastrigin-function test, during both training and testing, the optimizees have different decision variables and use random initializations; (iii) in the NN training test, the training optimizees use the same network architecture and dataset but random initializations; however, testing samples optimizees from a different distribution, that is, the L2O optimizer is applied to train a network of an unseen architecture and on a different dataset.

**Compared Methods and Evaluation Settings:** For each problem, we choose applicable approaches that include both model-free and/or model-based ones, implement them in the TensorFlow framework, ensure identical training/testing data, and evaluate them in the same but problem-specific metrics. After presenting the results, we draw observations from these benchmark experiments.

Our datasets and software are available as the Open-L2O package at: https://github.com/VITA-Group/Open-L2O. We hope that Open-L2O will foster reproducible research, fair benchmarking, and coordinated development efforts in the L2O field.

4.1 Test 1: Convex Sparse Optimization

4.1.1 Learning to Perform Sparse Optimization and Sparse-Signal Recovery

**Problem definition** The sparse recovery problem has been widely studied in the model-based L2O literature \cite{4, 1, 2}. The task is to recover a sparse vector from its noisy linear measurements:

\[
b_q = A x_q^* + e_q, \tag{17}\]
where $x^*_q \in \mathbb{R}^n$ is a sparse vector, $\varepsilon_q \in \mathbb{R}^m$ is an additive noise, and $q$ indexes an optimizee instance. While $x^*_q, b_q, \varepsilon_q$ change for each $q$, the measurement matrix $A \in \mathbb{R}^{m \times n}$ is fixed across all training and testing. In practice, there is a matrix $A$ associated with each sensor or sensing procedure.

Data generation  We generate 51,200 samples as the training set and 1,024 pairs as the validation and testing sets, following the i.i.d. sampling procedure in [1]. We sample sparse vectors $x^*_q$ with components drawn i.i.d. drawn from the distribution Ber(0.1) · $N(0, 1)$, yielding an average sparsity of $\sim 10\%$. We run numerical experiments in four settings:

- **Noiseless.** We take $(m,n) = (256, 512)$ and sample $A$ with $A_{ij} \sim N(0, 1/m)$ and then normalize its columns to have the unit $\ell_2$ norm. The noise $\varepsilon$ is always zero.

- **Noisy.** The same as above except for Gaussian measurement noises $\epsilon_q$ with a signal-to-noise ratio (SNR) of 20dB.

- **Coherent.** A Gaussian random matrix is highly incoherent, making sparse recovery relatively easy. To increase the challenge, we compute a dictionary $D \in \mathbb{R}^{256 \times 512}$ from 400 natural images in the BSD500 dataset [222] using the block proximal gradient method [223] and, then, use it as the measurement matrix $A$, which has a high coherence. Other settings remain unchanged from the first setting above.

- **Larger scale.** We scale up the noiseless case to $(m,n) = (512, 1024)$; the other settings stay the same.

Model and training settings  All our model-based L2O approaches take measurements $b_q$ as input and return estimates $\hat{x}_q \approx x^*_q$. They are trained to minimize the mean squared error $E_q \sim \| \hat{x}_q - x^*_q \|_2^2$. We adopt a progressive training scheme following [12; 1; 215]. We use a batch size of 128 for training and a learning rate of $5 \times 10^{-4}$. Other hyperparameters follow the default suggestions in their original papers. After training, the learned models are evaluated on the test set in NMSE (Normalized Mean Squared Error) in the decibel (dB) unit:

$$\text{NMSE}_{\text{dB}}(\hat{x}_q, x^*_q) = 10 \log_{10} \left( \frac{\| \hat{x}_q - x^*_q \|^2}{\| x^*_q \|^2} \right).$$

We compare the following model-based L2O approaches, all of which are unrolled to 16 iterations: (1) a feed-forward version LISTA as proposed in [4], with untied weights across layers; (2) LISTA-CP, a variant of LISTA with weight coupling [129; 1; (3) LISTA-CPSS, a variant of LISTA with weight coupling and support selection techniques [1; 2; (4) ALISTA [2], the variant of LISTA-CPSS with minimal learnable parameters; (5) LAMP [12], an L2O network unrolled from the AMP algorithm; (6) LFISTA [224], an L2O network unrolled from the FISTA algorithm; (7) GLISTA [215], an improved LISTA model by adding gating mechanisms.

Results  Figure 5 summarizes the comparisons; its four subfigures present the NMSEs of sparse recovery under noiseless, noisy, coherent dictionary, and large scale settings, respectively. We make the following observations:

- In the noiseless setting, ALISTA yields the best final NMSE, and the support selection technique in LISTA-CPSS contributes to good performance. LAMP and GLISTA achieve slightly worse NMSEs than LISTA-CPSS but are much better than LISTA. Surprisingly, LFISTA fails to outperform LISTA, which we attribute to its heavier parameterization in [224] than other models and consequently harder training.
In the noisy setting of 20dB SNR, the performance of LAMP degrades severely while LISTA-CPSS and ALISTA still perform robustly. GLISTA outperforms all the others in the setting.

In setting of a coherent dictionary, ALISTA suffers significantly from the coherence while the other methods that learn the weight matrices from data cope this issue better. In particular, GLISTA has the best performance thanks to the gate mechanisms.

When it comes to a larger scale setting \((m,n) = (512,1024)\), LISTA, LISTA-CP, LFISTA and GLISTA have performance degradation due to the higher parameterization burden. In comparison, ALISTA becomes the best among all since it has the fewest parameters to learn: only the layer-wise thresholds and step-sizes. Therefore, ALISTA is least impacted by the problem scale and produces nearly the same performance as it does the smaller setting of \((m,n) = (256,512)\).

### 4.1.2 Learning to minimize the Lasso model

**Problem definition** Instead of sparse recovery, which aims to recover the original sparse vectors, our goal here is to minimize the LASSO objective even though its solution is often different from the true sparse vector:

\[
x_q^{\text{Lasso}} = \arg \min_x f_q(x), \quad \text{where } f_q(x) = \frac{1}{2} \|Ax - b_q\|_2^2 + \lambda \|x\|_1,
\]

where \(A \in \mathbb{R}^{m \times n}\) is a known, fixed, and normalized dictionary matrix, whose elements are sampled i.i.d. from a Gaussian distribution. An optimizee instance with index \(q\) is
characterized by a sparse vector \( x^*_q \) and \( b_q \in \mathbb{R}^{m \times 1} \) is the observed measurement under \( A \) from \( x^*_q \) following the same linear generation model in the previous subsection. \( \lambda \) is a hyperparameter usually selected manually or by cross-validation, and is chosen to be 0.005 by default in all our experiments.

Uniquely, we compare both model-based and model-free L2O methods in this section. For the former, we can adopt similar algorithm unrolling recipes as in the previous subsection. For the latter, we treat the problem as generic minimization and apply LSTM-based L2O methods. To our best knowledge, this is the first comparison between the two distinct L2O mainstreams. We hope the results provide quantitative understanding how much we can gain from incorporating problem-specific structures (when available) into L2O.

**Data generation** We run the experiments with \((m,n) = (5,10)\), as well as \((m,n) = (25,50)\). We did not go larger due to the high memory cost of LSTM-based model-free L2O methods. We sample 12,800 pairs of \( x^*_q \) and \( b_q \) for training and 1,280 pairs for validation and testing. The samples are noiseless. During the testing, we set 1,000 iterations for both model-free L2O methods and classic optimizers. We run model-based L2O ones with a smaller fixed number of iterations (layers), which are standard for them.

For each sample \( b_q \), we let \( f^*_q \) denote the optimal Lasso objective value, \( f_q(x^{\text{Lasso}}) \). The optimization performance is measured with a modified relative loss:

\[
R_{f,Q}(x) = \frac{\mathbb{E}_{q \sim Q}[f_q(x) - f^*_q]}{\mathbb{E}_{q \sim Q}[f^*_q]},
\]

where the optimal solution is generated by 2,000 iterations of FISTA and the expectations are taken over the entire testing set.

We run four categories of methods to solve the LASSO optimization problem:

- **Three sub-gradient descent algorithms**: GD, ADAM and RMSProp, which serve as “natural baselines” with neither learning nor problem-specific structure. All these algorithms use a step size of \( 10^{-3} \). We verified that changing step sizes did not notably alter their performance. (Even though GD and its generalizations do not handle nonsmooth objectives without smoothing or using proximal operators, they are popular optimizers that people try on almost everything, so we choose to test them anyway.)

- **Two problem-specific analytic optimizers**: ISTA [153], a forward backward splitting algorithm using a soft-thresholding backward operator and its Nesterov-accelerated version, FISTA [160]. For both methods, we use a step size of \( 1/L \), where \( L \) is the largest eigenvalue of \( A^T A \), and a threshold of \( \lambda/L \) for the soft-thresholding function.

- **Two model-based L2O models**: vanilla LISTA [4] and ALISTA [2], both unrolled to 16 layers. We follow the same training setting in the last subsection except that we replace the training loss from the mean squared error (w.r.t. the true solution) with the LASSO loss. Since other compared methods take far more iterations, we also tried to expand LISTA/ALISTA to more iterations during testing by appending FISTA iterations.

- **Three model-free L2O methods**: L2O-DM [5], L2O-RNNprop [50] and L2O-enhanced optimizers [64] (Section 2.1). All L2O optimizers are trained with the Lasso objective function as the reward for 100 epochs and 1,000 iterations per epoch. The reported evaluation is the average performance over 10 random starting points.

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Figure 6: Evaluation comparisons among analytic, model-based L2O, and model-free L2O optimizers on Lasso. y-axis represents the modified relative loss (19), and x-axis denotes the number of iterations, both in the logarithmic scale.

Results From Figure 6, we make the following observations:

- At the small problem size \((5,10)\), both ISTA and FISTA converge fast in tens of iterations with FISTA being slightly ahead. Both model-based L2O models, ISTA and ALISTA of 16 iterations (layers), converge to solutions of precision compared to what FISTA can achieve after hundreds of iterations and better than what ISTA can do at 1,000 iterations. The advantage of ISTA/ALISTA can sustain beyond 16 iterations using FISTA updates.

- In comparison, at the small problem size \((5,10)\), model-free L2O methods exhibit far worse performance. L2O-RNNprop has slower convergence than ISTA/FISTA and only produces reasonably good solutions after 1,000 iterations – though still much better than analytic optimizers Adam and RMSProp. L2O-DM, L2O-enhanced, and vanilla GD completely fail to decrease the objective value.

- At the larger problem size \((25,50)\), LISTA and ALISTA still converge to high-precision solutions with only 16 iterations with sustained advantages from the FISTA extension. Interestingly, ISTA now becomes much slower than FISTA. All the sub-gradient descent and model-free L2O methods remain to perform poorly; only L2O-RNNprop can converge faster than ISTA and comparable to FISTA, though reaching lower precision.

- Our experiments clearly demonstrate the dominant advantage of incorporating problem-specific structures to the optimizers when it comes to both analytic and learned optimizers.

4.2 Test 2: Minimization of non-convex function Rastrigin

We now turn to non-convex minimization. One popular non-convex test function is called the Rastrigin function:

\[
f(x) = \frac{1}{2} \sum_{i=1}^{n} x_i^2 - \sum_{i=1}^{n} \alpha \cos (2\pi x_i) + \alpha n,
\]  

(20)
where $\alpha = 10$. We consider a broad family of similar functions $f_q(x)$ that generalizes Rastrigin function:

$$f_q(x) = \frac{1}{2} \| A_q x - b_q \|_2^2 - \alpha c_q \cos(2\pi x) + \alpha n,$$

where $A_q \in \mathbb{R}^{n \times n}$, $b_q \in \mathbb{R}^{n \times 1}$ and $c_q \in \mathbb{R}^{n \times 1}$ are parameters whose elements are sampled i.i.d. from $\mathcal{N}(0, 1)$. Obviously, the function (20) is a special case in this family with $A = I$, $b = \{0, 0, \ldots, 0\}^T$, $c = \{1, 1, \ldots, 1\}^T$. For training, we sample 1,280 triplets of $\{A_q, b_q, c_q\}$ on two problem scales: $n = 2$ and $n = 10$. For evaluation, we sample another 128 combinations of $A_q, b_q$ and $c_q$ and report the average performance over 10 random starting points. The number of steps for evaluation is 1,000.

![Figure 7](image_url)

Figure 7: Evaluation and comparison across model-free L2O and analytic optimizers on the generalized family of Rastrigin functions. y-axis represents the average loss of function values, and x-axis denotes the number of iterations in the logarithm scale.

Two groups of methods are compared: (1) Four traditional gradient descent algorithms, including ADAM with a $10^{-1}$ step size, RMSProp with a $3 \times 10^{-1}$ step size, GD with the line-searched step size started from $10^{-1}$, and NAG (Nesterov Accelerated Gradient) with the line-searched step size started from $10^{-1}$. All other hyperparameters are tuned by careful grid search. (2) Five model-free L2O methods, including L2O-DM [5], L2O-enhanced [64], L2O-Scale [51], L2O-RNNProp [50] (Section 2.1), and L2O-Swarm [14] (Section 2.3). All L2O optimizers are trained for 100 epochs and 1000 iterations per epoch. At the testing stage, we evaluate and report the logarithmic loss of unseen functions from the same family, which are plotted in Figure 7.

**Results** From Figure 7 we draw the following observations:

- ADAM and RMSProp converge slightly more quickly than GD and NAG with line search, especially in the early stage, for both $n = 2$ and $n = 10$. All analytic optimizers converge to local minima of similar qualities for $n = 2$, and NAG finds a slightly better solution for $n = 10$.

- L2O-DM, L2O-enhanced, L2O-RNNProp, and L2O-Scale perform similarly to analytic optimizers regarding both solution quality and convergence speed, showing no obvious advantage over analytic optimizers. For $n = 10$, L2O-enhanced finds a solution of better quality than the other two. But the three model-free L2Os have larger error bars at $n = 10$, which indicates model instability.
• Although not converging faster than others, L2O-Swarm locates higher quality solutions (lower losses) for both $n$ values, especially $n = 10$. Since L2O-Swarm is the only method that leverages a population of LSTM-based L2O “particles”, it explores a larger landscape than the other methods, so it is not surprising that its higher search cost leads to better solutions.

• The oracle objectives (dashed black lines) in Figure 7 are generated by the Run-and-Inspect method [225], which can provably find a global minimum if the objective function can be decomposed into a smooth strongly convex function (e.g., a quadratic function) plus a restricted function (e.g., sinusoidal oscillations). For $n = 2$, almost all methods reach a similar loss lied in $[5, 5, 6.0]$ in the end. For $n = 10$, L2O-Swarm performs significantly better than other methods in terms of the achieved final loss, though it converges more slowly than most of the other approaches.

4.3 Test 3: Neural Network Training

Our last test is training multi-layer neural networks (NNs), one of the most common tasks of L2O since its beginning. This has been the playground for model-free L2O methods. There are few problem-specific structures to explore. Common optimizers are stochastic gradient descent and their variants. We hope this test to address two questions:

• Can model-free L2O optimizers outperform analytic ones on neural network training? If so, how much is the margin?

• Can model-free L2O optimizers generalize to unseen network architectures and data?

To fairly compare different methods and answer these questions, we train L2O optimizers on the same neural network used in [5]: a simple Multi-layer Perceptron (MLP) with one 20-dimension hidden layer and the sigmoid activation function, trained on the MNIST dataset to minimize the cross-entropy loss. Therefore, the task distribution becomes optimizing the same MLP model with different random initializations.

We probe the (out of distribution) generalizability of the learned optimizers by testing them on two unseen optimization tasks, following the practice in [5; 64]:

1. Train another MLP with one 20-dimension hidden layer, but using the ReLU activation function, on the MNIST dataset.

2. Train another ConvNet on the MNIST dataset, consisting two convolution layers, two max pooling layers, and one last fully connected layer. The first convolution layer uses 16 $3 \times 3$ filters with stride 2. The second convolution layer uses 32 $5 \times 5$ filters with stride 2. The max pooling layers have size $2 \times 2$ with stride 2.

We compare four model-free L2O optimizers: L2O-DM [5], L2O-enhanced [64], L2O-Scale [51], and L2O-RNNprop [50] (Section 2.1), all following the hyperparameters suggested in their original papers. SGD, Adam, and RMSProp are analytical optimizers served as the baselines. All L2O optimizers are trained with the single model from 10,000 different random initializations drawn from $\mathcal{N}(0, 0.1)$. On each optimizee, now corresponding to a random initialization, the optimizers run for 100 iterations. The training uses a batch size of 128. During each run of testing, we evaluate learned optimizers on an unseen testing optimizee for 10,000 steps, which is much more than the training iteration number. We
then report the training loss of the optimizee. We perform 10 independent runs and report the error bars.

It is worth mentioning that current L2O-optimizers can hardly scale to training large neural networks with over $1 \times 10^6$ parameters. This is because to update the L2O optimizer using back-propagation, we need to keep the gradients during unrolling and the computation graph of the optimizee in memory, therefore, consuming too much memory. This remains an open challenge.

![Figure 8](image)

Figure 8: Evaluation and comparison across model-free L2O and analytic optimizers on neural network training. y-axis represents the average training loss values, and x-axis denotes the number of iterations. Each curve is the average of ten runs, and error bars are plotted.

**Results** From Figure 8, we make the following observations:

- The relative performance of analytic optimizers is consistent: both Adam and RMSProp achieve similar training losses on both ReLU-MLP and ConvNets, and better than SGD. However, the learned optimizers exhibit different levels of transferability. L2O-DM completely diverges on ConvNet, but works on ReLU-MLP when the number of iterations is small, showing poor generalization.

- L2O-Scale and L2O-RNNprop can generalize to ReLU-MLP as they achieve similar or lower training losses than analytic optimizer), but they cannot effectively optimize ConvNet. L2O-RNNprop only works for a small number of iterations less than 4,000. L2O-Scale generalizes better to larger iteration numbers, thanks to its random scaling during training.

- L2O-enhanced, which adopts increasing unroll length and off-line imitation strategy, achieves the best performance on both unseen networks after extended (optimizee) training iterations.

- The issue of lacking stability during testing exists for all L2O optimizers. On both ReLU-MLP and ConvNet, all L2O optimizers suffer from larger loss variances as the training iteration grows, causing the errors to fluctuate more severely that those of the analytic optimizers.
4.4 Task-home messages

• When appropriate model or problem structures are available to be leveraged, model-based L2O methods clearly outperform model-free ones and analytic optimizers. On optimizees sampled from the same task distribution, model-based L2O optimizers have solid, consistent, and reliable performance.

• Although in many cases we can also observe performance gains of model-free L2O approaches over analytic ones, the gains are not consistent. The benefits of current model-free L2O approaches come to questions when they are applied to larger optimizees or applied with more iterations during testing, let alone to “out-of-distribution” optimizees. We find no clear winner in existing model-free L2O approaches that can consistently outperforms others across most tests.

5. Concluding Remarks

This article provided the first “panoramic” review of the state-of-the-arts in the emerging L2O field, accompanied with the first-of-its-kind benchmark. The article reveals that, despite its promise, the L2O research is still in its infancy, facing many open challenges as well as research opportunities from practice to theory.

On the theory side, Section 3.4 has listed many open theoretical questions for us to understand why and how model-based L2O methods outperform traditional optimizers. Besides those, for model-free L2O methods, the theoretical foundation has been scarce if any at all. For example, although the training of L2O optimizers is often successful empirically, there has been almost no theoretical result established for the convergence performance of such a L2O training process, putting the general feasibility of obtaining L2O models in question. Also, for both model-based and mode-free L2O approaches, the generalization or adaption guarantees of trained L2O to optimizees out of the task distribution are understudied yet highly demanded.

On the practical side, the scalability of model-free L2O methods is perhaps the biggest hurdle for them to become more practical. That includes scaling up to both larger and more complicated models, and to more iterations during meta-testing. For model-specific L2O, the current empirical success is still limited to a few special instances in inverse problems and sparse optimization, and relying on case-by-case modeling. An exploration towards broader applications, and perhaps building a more general framework, is demanded.

Furthermore, there is no absolute border between mode-based and model-free L2O methods, and the spectrum between the two extremes can hint many new research opportunities. [45] suggested a good perspective that unrolled networks (as one example of model-based L2O) is an intermediate state between generic networks and analytic algorithms, and might be more data-efficient to learn. The view was supported by [226], which further advocated that the unrolled model might be considered as a robust starting point for subsequently data-driven model search. We also believe that end-to-end learning approaches can be improved with current continuous optimization algorithms to benefit from the theoretical guarantees and state-of-the-art algorithms already available.

So, to conclude this article, let us quote Sir Winston Churchill: “Now this is not the end. It is not even the beginning of the end. But it is, perhaps, the end of the beginning.” Although most approaches we discussed in this paper are still at an exploratory level of deployment, and are apparently not yet ready as general-purpose or commercial solvers, we
are strongly confident that machine learning has just began to feed the classic optimization
field, and the blowout of L2O research progress has yet to start.

Appendix A. List of Abbreviations

| Abbreviation | Full Name | Description |
|--------------|-----------|-------------|
| Adam         | Adaptive Moment Estimation | A first-order gradient-based optimization algorithm of stochastic objective functions, based on adaptive estimates of lower-order moments. |
| ADMM         | Alternating Direction Method of Multipliers | An optimization algorithm that solves convex problems by breaking them into smaller pieces, each of which are then easier to handle. |
| AutoML       | Automated Machine Learning | The process of automating the process of applying machine learning to real-world problems. |
| CASH         | Combined Algorithm Selection and Hyperparameter Optimization | A model selection strategy that consider the choosing of data preparation, learning algorithm, and algorithm hyperparameters as one large global optimization problem. |
| GAN          | Generative Adversarial Network | A class of machine learning frameworks’ 08s, where two neural networks contest with each other in a zero-sum game, to learn to generate new data with the same statistics as a given training set. |
| HPO          | Hyperparameter Optimization | The problem of choosing a set of optimal hyperparameters for a machine learning algorithm. |
| L2O          | Learning to Optimize | Learnable optimizers to predict update rules (for optimizers) fit from data. |
| LASSO        | Least Absolute Shrinkage and Selection Operator | A linear regression method that uses shrinkage to encourages simple, sparse models. |
| LSTM         | Long-short Term Memory | A variant of artificial recurrent neural network (RNN) architecture, typically used to process sequence data. |
| MLP          | Multi-layer Perceptron | A class of feedforward artificial neural network (ANN) composed of multiple layers of perceptrons (with threshold activation). |
| PGD          | Projected Gradient Descent | Projected gradient descent minimizes a function subject to a constraint. At each step we move in the direction of the negative gradient, and then “project” onto the feasible set. |
| ReLU         | Rectified Linear Unit | An activation function used in neural networks, defined as $f(x) = \max(x, 0)$. |
| RL           | Reinforcement Learning | RL is an area of machine learning regarding how to make intelligent agents take actions in an environment to maximize the cumulative reward. |
| RMSProp      | Root Mean Square Propagation | RMSprop gradient descent maintains a moving (discounted) average of the square of gradients, and then divide the gradient by the root of this average. |

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