Automatically Learning Compact Quality-aware Surrogates for Optimization Problems

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Abstract

Solving optimization problems with unknown parameters often requires learning a predictive model to predict the values of the unknown parameters and then solving the problem using these values. Recent work has shown that including the optimization problem as a layer in the model training pipeline results in predictions of the unobserved parameters that lead to higher decision quality. Unfortunately, this process comes at a large computational cost because the optimization problem must be solved and differentiated through in each training iteration; furthermore, it may also sometimes fail to improve solution quality due to non-smoothness issues that arise when training through a complex optimization layer. To address these shortcomings, we learn a low-dimensional surrogate model of a large optimization problem by representing the feasible space in terms of meta-variables, each of which is a linear combination of the original variables. By training a low-dimensional surrogate model end-to-end, and jointly with the predictive model, we achieve: i) a large reduction in training and inference time; and ii) improved performance by focusing attention on the more important variables in the optimization and learning in a smoother space. Empirically, we demonstrate these improvements on a non-convex adversary modeling task, a submodular recommendation task and a convex portfolio optimization task.

1 Introduction

Uncertainty is a common feature of many real-world decision-making problems because critical data may not be available when a decision must be made. Here is a set of representative examples: recommender systems with missing user-item ratings [22], portfolio optimization where future performance is uncertain [30], and strategic decision-making in the face of an adversary with uncertain objectives [25]. Often, the decision-maker has access to features that provide information about the values of interest. In these settings, a predict-then-optimize [13] approach naturally arises, where we learn a model that maps from the features to a value for each parameter and optimize using this point estimate [45]. In principle, any predictive modeling approach and any optimization approach can be applied, but using a generic loss function to train the model may result in poor decision performance. For example, a typical ratings prediction approach in recommendation system may equally weight errors across different items, but in the recommendation task, misclassifying a
trendy item can result in more revenue loss than misclassifying an ordinary item. We may instead want to train our model using a “task-based” or “decision-focused” loss, approximating the decision quality induced by the predictive model, which can be done by embedding the optimization problem as a layer in the training pipeline. This end-to-end approach improves performance on a variety of tasks \[6, 44, 9\].

Unfortunately, this end-to-end approach suffers from poor scalability because the optimization problem must be solved and differentiated through on every training iteration. Furthermore, the output of the optimization layer may not be smooth, sometimes leading to instabilities in training and consequently poor solution quality. We address these shortcomings that arise in the end-to-end approach due to the presence of a complex optimization layer by replacing it with a simpler surrogate problem. The surrogate problem is learned from the data by automatically finding a reparameterization of the feasible space in terms of meta-variables, each of which is a linear combination of the original decision variables. The new surrogate problem is generally cheaper to solve due to the smaller number of meta-variables, but it can be lossy—the optimal solution to the surrogate problem may not match the optimal solution to the original. Since we can differentiate through the surrogate layer, we can optimize the choice of surrogate together with predictive model training to minimize this loss. The dimensionality reduction offered by a compact surrogate simultaneously reduces training times, helps avoid overfitting, and sometimes smooths away bad local minima in the training landscape.

In short, we make several contributions. First, we propose a linear reparameterization scheme for general optimization layers. Second, we provide theoretical analysis of this framework along several dimensions: (i) we show that desirable properties of the optimization problem (convexity, submodularity) are retained under reparameterization; (ii) we precisely characterize the tractability of the end-to-end loss function induced by the reparameterized layer, showing that it satisfies a form of coordinate-wise quasiconvexity; and (iii) we provide sample complexity bounds for learning a model which minimizes this loss. Finally, we demonstrate empirically on a set of three diverse domains that our approach offers significant advantages in both training time and decision quality compared previous approaches to embedding optimization in learning.

Related work Surrogate models \[16, 37, 28\] are a classic technique in optimization, particularly for black-box problems. Previous work has explored linear reparameterizations to map between low and high fidelity models of a physical system \[5, 38, 4\] (e.g., for aerospace design problems). However, both the motivation and underlying techniques differ crucially from our work: previous work has focused on designing surrogates by hand in a domain-specific sense, while we leverage differentiation through the optimization problem to automatically produce a surrogate that maximizes overall decision quality.

Our work is closest to the recent literature on differentiable optimization. Amos et al. \[2\] and Agrawal et al. \[1\] introduced differentiable quadratic programming and convex programming layers, respectively, by differentiating through the KKT conditions of the optimization problem. Donti et al. \[9\] and Wilder et al. \[44\] apply this technique to achieve end-to-end learning in convex and discrete combinatorial programming, respectively. Perrault et al. \[33\] applied the technique to game theory with a non-convex problem, where a sampling approach was proposed by Wang et al. \[42\] to improve the scalability of the backward pass. All the above methods share scalability and non-smoothness issues: each training iteration requires solving the entire optimization problem and differentiating through the resulting KKT conditions, which requires \(O(n^3)\) time in the number of decision variables and may create a non-smooth objective. Our surrogate approach aims to rectify both of these issues.

2 Problem Statement

We consider an optimization problem of the form: \(\min_{x \text{ feasible}} f(x, \theta_{\text{true}})\). The objective function depends on a parameter \(\theta_{\text{true}} \in \Theta\). If \(\theta_{\text{true}}\) were known, we assume that we could solve the optimization problem using standard methods. In this paper, we consider the case that parameter \(\theta_{\text{true}}\) is unknown and must be inferred from the given available features \(\xi\). We assume that \(\xi\) and \(\theta_{\text{true}}\) are correlated and drawn from a joint distribution \(D\), and our data consists of samples from \(D\). Our task is to select the optimal decision \(x^*(\xi)\), function of the available feature, to optimize the expected objective value:

\[
\min_{x^* \text{ feasible}} \mathbb{E}_{(\xi, \theta_{\text{true}}) \sim D}[f(x^*(\xi), \theta_{\text{true}})]
\]
In this paper, we focus on a predict-then-optimize framework, which proceeds by learning a model $\Phi(\cdot, w)$, mapping from the features $\xi$ to the missing parameter $\theta_{\text{true}}$. When $\xi$ is given, we first infer $\theta = \Phi(\xi, w)$ and then solve the resulting optimization problem to get the optimal solution $x^*$:

$$\min_{x} \ f(x, \theta), \quad \text{s.t.} \ h(x) \leq 0, \ Ax = b \quad (2)$$

This reduces the decision-making problem with unknown parameters to a predictive modeling problem: how to learn a model $\Phi(\cdot, w)$ that leads to the best performance.

A standard approach to solve the predict-then-optimize problem is two-stage learning, which trains the predictive model without knowledge of the decision-making task (Figure 1). The predictive model minimizes the mismatch between the predicted parameters and the ground truth: $E_{(\xi, \theta_{\text{true}}) \in D} f(\Phi(\xi, w), \theta_{\text{true}})$, with any loss metric $\ell$. Such two-stage approach is very efficient, but it may lead to poor performance when a standard loss function is used. Performance can be improved if the loss function is carefully chosen to suit the task [12], but doing so is challenging for an arbitrary optimization problem.

Gradient-based end-to-end learning approaches in domains with optimization layers involved, e.g., decision-focused learning [13, 11], directly minimize Equation 1 as the training objective, which requires back-propagating through the optimization layer in Equation 2. This end-to-end approach is able to achieve better solution quality compared to two-stage learning, in principle. However, because the decision-focused approach has to repeatedly solve the optimization program and back-propagate through it, scalability becomes a serious issue. Additionally, the complex optimization layer can also jeopardize the smoothness of objective value, which is detrimental for training parameters of a neural network-based predictive model with gradient-based methods.

### 3 Surrogate Learning

The main idea of the surrogate approach is to replace Equation 2 with a carefully selected surrogate problem. To simplify Equation 2 we can linearly reparameterize $x = Py$, where $y \in \mathbb{R}^m$ with $m \ll n$ and $P \in \mathbb{R}^{n \times m}$,

$$\min_{y} \ g_P(y, \theta) := f(Py, \theta) \quad \text{s.t.} \ h(Py) \leq 0, \ APy = b \quad (3)$$

Since this reparameterization preserves all the equality and inequality constraints in Equation 2, we can easily transform a feasible low-dimensional solution $y^*$ back to a feasible high-dimensional solution with $x^* = Py^*$. The low-dimensional surrogate is generally easier to solve, but lossy, because we restrict the feasible region to a hyperplane spanned by $P$. If we were to use a random reparameterization, the solution we recover from the surrogate problem could be far from the actual optimum in the original optimization problem, which could significantly degrade the solution quality.

This is why we need to learn the surrogate and its reparameterization matrix. Because we can differentiate through the surrogate optimization layer, we can estimate the impact of the reparameterization matrix on the final solution quality. This allows us to run gradient descent to learn the reparameterization matrix $P$. The process is shown in Figure 3. Notice that the surrogate problem also takes the prediction $\theta$ of the predictive model as input. This implies that we can jointly learn the predictive model and the reparameterization matrix by solely solving the cheaper surrogate problem.
Differentiable optimization In order to differentiate through the optimization layer as shown in Figure 2, we can compute the derivative of the solution quality, evaluated on the optimal solution \( x^* \) and true parameter \( \theta_{\text{true}} \), with respect to the model’s weights \( w \) by applying the chain rule:

\[
\frac{df(x^*, \theta_{\text{true}})}{dw} = \frac{df(x^*, \theta_{\text{true}})}{dx^*} \frac{dx^*}{d\theta} \frac{d\theta}{dw}
\]

where \( \frac{dx^*}{d\theta} \) can be obtained by differentiating through KKT conditions of the optimization problem.

Similarly, in Figure 3, we can apply the same technique to obtain the derivatives with respect to the weights \( w \) and reparameterization matrix \( P \):

\[
\frac{df(x^*, \theta_{\text{true}})}{dw} = \frac{df(x^*, \theta_{\text{true}})}{dx^*} \frac{dx^*}{dy^*} \frac{dy^*}{d\theta} \frac{d\theta}{dw}
\]

\[
\frac{df(x^*, \theta_{\text{true}})}{dP} = \frac{df(x^*, \theta_{\text{true}})}{dx^*} \frac{dx^*}{dy^*} \frac{dy^*}{dP}
\]

where \( y^* \) is the optimal solution of the surrogate problem, \( x^* = Py^* \), and \( \frac{dy^*}{d\theta}, \frac{dy^*}{dP} \) can be computed by differentiating through the KKT conditions of the surrogate optimization problem.

4 Analysis of Linear Reparameterization

The following sections address three major theoretical aspects: (i) complexity of solving the surrogate problem, (ii) learning the reparameterization, and (iii) learning the predictive model.

4.1 Convexity and DR-Submodularity of the Reparameterized Problem

In this section, we fix the predictive model and the linear reparameterization. We prove below that convexity and continuous diminishing-return (DR) submodularity [7] of the original function \( f \) is preserved after applying reparameterization. This implies that the new surrogate problem can be solved by gradient descent or Frank-Wolfe [8, 23, 17] algorithm with an approximation guarantee.

**Proposition 1.** If \( f \) is convex, then \( g_P(y, \theta) = f(Py, \theta) \) is convex.

**Proposition 2.** If \( f \) is DR-submodular and \( P \geq 0 \), then \( g_P(y, \theta) = f(Py, \theta) \) is DR-submodular.

4.2 Convexity of Reparameterization Learning

In this section, we assume the predictive model \( \Phi \) is fixed. We want to analyze the convergence of learning the surrogate and its linear reparameterization \( P \). Let us denote the optimal value of the optimization problem in the form of Equation 3 to be \( \text{OPT}(\theta, P) := \min_{y \text{ feasible}} g_P(y, \theta) \in \mathbb{R} \). It would be ideal if \( \text{OPT}(\theta, P) \) is convex in \( P \) so that gradient descent would be guaranteed to recover the optimal reparameterization. Unfortunately, this is not true in general, despite the fact that we use a linear reparameterization: \( \text{OPT}(\theta, P) \) is not even globally quasiconvex in \( P \).

**Proposition 3.** \( \text{OPT}(\theta, P) = \min_{y \text{ feasible}} g_P(y, \theta) \) is not globally quasiconvex in \( P \).

Fortunately, we can guarantee the partial quasiconvexity of \( \text{OPT}(\theta, P) \) in the following theorem:

**Theorem 1.** If \( f(\cdot, \theta) \) is quasiconvex, then \( \text{OPT}(\theta, P) = \min_{y \text{ feasible}} g_P(y, \theta) \) is quasiconvex in \( P \) for any \( 1 \leq i \leq m \), where \( P = [P_1, P_2, \ldots, P_m] \geq 0 \).
We conduct experiments on three different domains where decision-focused learning has been applied: when the objective function is convex (14). Throughout all the experiments, we compare the performance of the surrogate model with a standard first-order method. Theorem 2. Let \( \mathcal{H}_\text{lin} \) be the hypothesis class of all linear function mappings from \( \xi \in \Xi \subset \mathbb{R}^n \) to \( \theta \in \Theta \subset \mathbb{R}^m \), and let \( P \in \mathbb{R}^{n \times m} \) be a linear reparameterization used to construct the surrogate. The expected Rademacher complexity over \( t \) i.i.d. random samples drawn from \( \mathcal{D} \) can be bounded by:

\[
\text{Rad}^t(\mathcal{H}_\text{lin}) \leq 2mC\sqrt{\frac{2p\log(2mt\|P^+\|\rho_2(S))}{t}} + O\left(\frac{1}{t}\right) \tag{4}
\]

where \( C \) is the gap between the optimal solution quality and the worst solution quality, \( \rho_2(S) \) is the pseudo-inverse. Equation 4 gives a bound on the Rademacher complexity, an upper bound on the generalization error with \( t \) samples given. Although a low-dimensional surrogate can lead to less representational power (lead to lower decision quality), we can also see that in Equation 4 when the reparameterization size \( m \) is smaller, a compact surrogate can get a better generalizability. This implies that we have to choose an appropriate reparameterization size to balance representational power and the generalizability.

5 Experiments

We conduct experiments on three different domains where decision-focused learning has been applied: (i) adversarial behavior learning in network security games with a non-convex objective (14). (ii) movie recommendation with a submodular objective (44), and (iii) portfolio optimization problem with a convex quadratic objective (14). Throughout all the experiments, we compare the performance and the scalability of the surrogate learning (surrogate), two-stage (TS), and decision-focused (DF) learning approaches. Performance is measured in terms of regret, which is defined as the difference between the achieved solution quality and the solution quality if the unobserved parameters \( \theta^* \) were observed directly—smaller is better. To compare scalability, we show the training time per epoch and inference time. The inference time corresponds to the time required to compute a decision for all instances in the testing set after training is finished. A short inference time may have intrinsic value, e.g., allowing the application to be run in edge computing settings. All methods are trained using gradient descent with optimizer Adam (26) with learning rate 0.01 and repeated over 30 independent runs to get the average. Each model is trained for at most 100 epochs with early stopping criteria when 3 consecutive non-improving epochs occur on the validation set. The reparameterization size is set to be 10% of the problem size throughout all three examples.

5.1 Adversarial Behavior Learning and Interdiction Games

Given a network structure \( G = (V,E) \), a NSG (network security game) (43) models the interaction between the defender, who places checkpoints on a limited number of edges in the graph, and an attacker who attempts to travel from a source to any of a set of target nodes in order to maximize the expected reward. The NSG is an extension of Stackelberg security games (40) (24), meaning that the defender commits to a mixed strategy first, after which the attacker chooses the path (having observed the defender’s mixed strategy but not the sampled pure strategy). In practice, the
 attacker is not perfectly rational. Instead, the defender can attempt to predict the attacker’s boundedly rational choice of path by using the known features of the nodes en route (e.g., accessibility or safety of hops) together with previous examples of paths chosen by the attacker.

Once the parameters \( \theta \) of the attacker behavioral model are given, finding the optimal defender’s strategy reduces to an optimization problem \( \max f(x, \theta) \) where \( x_e \) is the probability of covering edge \( e \in E \) and \( f \) gives the defender’s expected utility for playing mixed strategy \( x \) when the attacker’s response is determined by \( \theta \). The defender must also satisfy the budget constraint \( \sum_{e \in E} x_e \leq k \) where \( k = 3 \) is the total defender resources. We use a GCN (graph convolutional network) \([22, 27, 19]\) to represent the predictive model of the attacker. We assume the attacker follows reactive Markovian behavior \([42, 18]\), meaning that the attacker follows a random walk through the graph, where the probability of transitioning across a given edge \((u, v)\) is a function of the defender’s strategy \( x \) and an unknown parameter \( \theta_e \), representing the “attractiveness” of node \( v \). The walk stops when the attacker either is intercepted by crossing an edge covered by the defender or reaches a target. The defender’s utility is \(-u(t)\) if the attacker reaches target \( t \) and 0 otherwise, and \( f \) takes an expectation over both the random placement of the defender’s checkpoints (determined by \( x \)) and the attacker’s random walk (determined by \( x \) and \( \theta \)). Our goal is to learn a GCN which takes node features as input and outputs the attractiveness over nodes \( \theta \).

**Experimental setup:** We generate random geometric graphs of varying sizes with radius 0.2 in a unit square. We select 5 nodes uniformly at random as targets with payoffs \( u(t) \sim \text{Unif}(5, 10) \) and 5 nodes as sources where the attacker chooses uniformly at random from. The ground truth attractiveness value \( \theta_v \) of node \( v \in V \) is proportional to the proximity to the closest target plus a random perturbation sampled as \( \text{Unif}(-1, 1) \) which models idiosyncrasies in the attacker’s preferences. The node features \( \xi \) are generated as \( \xi = \text{GCN}(\theta) + 0.2N(0, 1) \), where GCN is a randomly initialized GCN with four convolutional layers and three fully connected layers. This generates random features with correlations between \( \xi_v \) (the features of node \( v \)) and both \( \theta_v \) and the features of nearby nodes. Such correlation is expected for real networks where neighboring locations are likely to be similar. The defender’s predictive model (distinct from the generative model) uses two convolutional and two fully connected layers, modeling a scenario where the true generative process is more complex than the learned model. We generate 35 random \((\xi, \theta)\) pairs for the training set, 5 for validation, and 10 for testing. Since decision-focused (DF) learning fails to scale up to larger instances, we additionally compare to a block-variable sampling approach specialized to NSG \([42]\) (block), which can speed up the backward pass by back-propagating through randomly sampled variables.

### 5.2 Movie Recommendation and Broadcasting Problem

In this domain, a broadcasting company chooses \( k \) movies out of a set of \( n \) available to acquire and show to their customers \( C \). \( k \) reflects a budget constraint. Each user watches their favorite \( T \) movies, with a linear valuation for the movies they watch. This is a variant of the classic facility location problem; similar domains have been used to benchmark submodular optimization algorithms \([29, 10]\).

In our case, the additional complication is that the user’s preferences are unknown. Instead, the company uses user’s past behavior to predict \( \theta_{ij} \in [0, 1] \), the preference score of user \( j \) for movie \( i \).

The company would like to maximize the overall satisfaction of users without exceeding the budget constraint \( k = 10 \). \( \{x_i\}_{i \in \{1, 2, \ldots, n\}} \) represents the decision of whether to acquire movie \( i \) or not. Once the preferences \( \theta_{ij} \) are given, the company wants to maximize the objective function:

\[
f(x) \coloneqq \sum_{j \in C} \max_{z_j \in \{0, 1\}, \sum_i z_i = T} \sum_{i \in \{1, 2, \ldots, n\}} x_i z_i \theta_{ij}
\]

**Experimental setup:** We use neural collaborative filtering \([21]\) to learn the user preferences. Commonly used in recommendation systems, the idea is to learn an embedding for each movie and user. The ratings are computed by feeding the concatenated user’s and movie’s embeddings to a neural network with fully connected layers. We use MovieLens \([20]\) as our dataset. The MovieLens dataset includes 25M ratings over 62,000 movies by 162,000 users. We first randomly select \( n \) movies as our broadcasting candidates. We additionally select 200 movies and use the users’ ratings on the movies as the users’ features. Then we split the users into disjoint groups on size 100 and each group serves as an instance of broadcasting task, where we want to choose \( k = 10 \) from the \( n \) candidate.
movies to recommend to the group members. Each user chooses $T = 3$ movies. 70% of the user groups are used for training, 10% for validation, and 20% for testing.

5.3 Stock Market Portfolio Optimization

Portfolio optimization can be treated as an optimization problem with missing parameters [34], where the return and the covariance between stocks in the next time step are not known in advance. We learn a model that takes features for each security and outputs the predicted future return. We adopt the classic Markowitz [30, 31] problem setup, where investors are risk-averse and wish to maximize a weighted sum of the immediate net profit and the risk penalty. The investor chooses a vector $x \geq 0$ with $\sum x_i = 1$, where $x_i$ represents the fraction of money invested in security $i$. The investor aims to maximize the penalized immediate return $f(x) := p^\top x - \lambda x^\top Q x$, where $p$ is the immediate net return of all securities and $Q \in \mathbb{R}^{n \times n}$ is a positive semidefinite matrix representing the covariance between the returns of different securities. A high covariance implies two securities are highly correlated and thus it is more risky to invest in both. We set the risk aversion constant to be $\lambda = 2$.

Experimental setup: We use historical daily price and volume data from 2004 to 2017 downloaded from the Quandl WIKI dataset [36]. We evaluate on the SP500, a collection of the 505 largest companies representing the American market. Our goal is to generate daily portfolios of stocks from a given set of candidates. Ground truth returns are computed from the time series of prices, while the ground truth covariance of two securities at a given time step is set to be the cosine similarity of their embeddings. We chronologically split the dataset into training, validation, and test sets with 70%, 10%, and 20% of the data respectively.
6 Discussion of Experimental Results

Performance: Figures 4(a), 5(a), and 6(a) compare the regret of our surrogate approach to the other approaches. In the non-convex (Figure 4(a)) and submodular (Figure 5(a)) settings, we see a larger improvement in solution quality relative to decision-focused learning. This is due to the huge number of local minima and plateaus in these two settings where two-stage and decision-focused approaches can get stuck. For example, when an incorrect prediction is given in the movie recommendation domain, some recommended movies could have no users watching them, resulting in a sparse gradient due to non-smoothness induced by the max in the objective function. The surrogate approach can instead spread the sparse gradient by binding variables with meta-variables, alleviating gradient sparsity. We see relatively less performance improvement (compared to decision-focused) when the optimization problem is strongly convex and hence smoother (Figure 6(a)), though the surrogate approach still achieves similar performance to the decision-focused approach.

Scalability: When the objective function is non-convex (Figure 4(b), 5(c)), our surrogate approach yields substantially faster training than standard decision-focused learning approaches (DF and block). The boost is due to the dimensionality reduction of the surrogate optimization problem, which can lead to speedups in solving the surrogate problem and back-propagating through the KKT conditions. While the two-stage approach avoids solving the optimization problem in the training phase (trading off solution quality), at test time, it still has to solve the expensive optimization problem, resulting in a similarly expensive inference runtime in Figure 4(c).

When the objective function is submodular (Figure 5(b), 5(c)), the blackbox optimization solver [41] we use in all experiments converges very quickly for the decision-focused method, resulting in training times comparable to our surrogate approach. However, Figure 5(a) shows that the decision-focused approach converges to a solution with very poor quality, indicating that rapid convergence may be a symptom of the uninformative local minima that the decision-focused method becomes trapped in.

Lastly, when the optimization problem is a quadratic program (Figure 6(b), 6(c)), solving the optimization problem can take cubic time, resulting in around a cubic speedup from the dimensionality reduction offered by our surrogate. Consequently, we see 7-fold faster training and inference times.

Visualization: We visualize the reparameterization for the NSG and movie recommendation domains in Figure 7. The initial reparameterization is shown in Figure 7(a) and 7(b). Initially, the weights put on the meta-variables are randomly chosen and no significant problem structure—no edge or circle colors—can be seen. After 20 epochs of training, in Figure 7(a), the surrogate starts putting emphasis on some important cuts between the sources and the targets, and in Figure 7(b), the surrogate focuses on the movies with higher average ratings and standard deviations. This allows the surrogate layer to extract the underlying structure of the optimization problem using fewer meta-variables.
7 Conclusion

In this paper, we focus on the shortcomings of scalability and solution quality that arise in end-to-end decision-focused learning due to the introduction of the differentiable optimization layer. We address these two shortcomings by learning a compact surrogate, with a learnable linear reparameterization matrix, to substitute for the expensive optimization layer. This surrogate can be jointly trained with the predictive model by back-propagating through the surrogate layer. Theoretically, we analyze the complexity of the induced surrogate problem and the complexity of learning the surrogate and the predictive model. Empirically, we show this surrogate learning approach leads to improvement in scalability and solution quality in three domains: a non-convex adversarial modeling problem, a submodular recommendation problem, and a convex portfolio optimization problem.

8 Broader impact:

End-to-end approaches can perform better in data-poor settings, improving access to the benefits of machine learning systems for communities that are resource constrained. Standard two-stage approaches typically requires enough data to learn well across the data distribution. In many domains focused on social impact such as wildlife conservation, limited data can be collected and the resources are also very limited. End-to-end learning is usually more favorable than two-stage approach under these circumstances; it can achieve higher quality results despite data limitations compared to two-stage approaches. This paper reduces the computational costs of end-to-end learning and increases the performance benefits.

But such performance improvements may come with a cost in transferability because the end-to-end learning task is specialized towards particular decisions, whereas a prediction-only model from the two-stage predict-then-optimize framework might be used for different decision making tasks in the same domain. Thus, the predictive model trained for a particular decision-making task in the end-to-end framework is not necessarily as interpretable or transferable as a model trained for prediction only. For real-world tasks, there would need to be careful analysis of cost-benefit of applying an end-to-end approach vis-a-vis a two-stage approach particularly if issues of interpretability and transferrability are critical; in some domains these may be crucial. Further research is required to improve upon these issues in the end-to-end learning approach.

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Appendix

9 Preservation of Convexity and Submodularity

**Proposition 1.** If $f$ is convex, then $g_{P}(y, \theta) = f(Py, \theta)$ is convex.

**Proof.** The convexity can be simply verified by computing the second-order derivative:

$$
\frac{d^2 g}{dy^2} = \frac{d^2 f(Py, \theta)}{dy^2} = P^T \frac{d^2 f}{dx^2} P \succeq 0
$$

where the last inequality comes from the convexity of $f$, i.e., $\frac{d^2 f}{dx^2} \geq 0$. \qed

**Proposition 2.** If $f$ is DR-submodular and $P \succeq 0$, then $g_{P}(y, \theta) = f(Py, \theta)$ is DR-submodular.

**Proof.** Assume $f$ has the property of diminishing return submodularity (DR-submodular) \[7\]. According to definition of continuous DR-submodularity, we have:

$$
\nabla^2_{x_i, x_j} f(x, \theta) \leq 0 \quad \forall i, j \in [n], y
$$

After applying the reparameterization, we can write:

$$
g_{P}(y, \theta) = f(x, \theta)
$$

and the second-order derivative:

$$
\nabla^2_2 g_{P}(y, \theta) = P^T \nabla^2_2 f(x, \theta) P \leq 0
$$

Since all the entries of $P$ are non-negative and all the entries of $\nabla^2_2 f(x, \theta)$ are non-positive by DR-submodularity, the product $\nabla^2_2 g_{P}(y, \theta)$ also has all the entries being non-positive, which satisfies the definition of DR-submodularity. \qed

10 Quasiconvexity in Reparameterization Matrix

**Proposition 3.** $OPT(\theta, P) = \min_{y \text{ feasible}} g_{P}(y, \theta)$ is not globally quasiconvex in $P$.

**Proof.** Without loss of generality, let us ignore the effect of $\theta$ and write $g_{P}(y) = f(Py)$. In this proof, we will construct a strongly convex function $f$ where the induced optimal value function $OPT(P) := \min_{y} g_{P}(y)$ is not quasiconvex.

Consider $x = [x_1, x_2, x_3]^T \in \mathbb{R}^3$. Define $f(x) = \left\| x - \begin{pmatrix} 1 \\ 1 \end{pmatrix} \right\|^2 \geq 0$ for all $x \in \mathbb{R}^3$. Define $P = \begin{pmatrix} 1 & 0 & 0 \\ 1 & 0 & 2 \\ 1 & 0 & 2 \end{pmatrix}$ and $P' = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 1 & 2 \\ 0 & 1 & 2 \end{pmatrix}$. Apparently, $x^* = \begin{pmatrix} 1 \\ 1 \\ 1 \end{pmatrix} = P\begin{pmatrix} 1 \\ 0.5 \\ 0.5 \end{pmatrix}$ and $x^* = \begin{pmatrix} 1 \\ 0.5 \\ 1 \end{pmatrix}$ are both achievable. So the optimal values $OPT(P) = OPT(P') = 0$. But the combination $P'' = \frac{1}{2}P + \frac{1}{2}P'$ cannot, which results in an optimal value $OPT(P'') = \min_{y} g_{P''}(y) \Rightarrow 0$

since $\begin{pmatrix} 1 \\ 1 \end{pmatrix} \notin \text{span}(P'')$. This implies $OPT(\frac{1}{2}P + \frac{1}{2}P') = OPT(P'') > 0 = \frac{1}{2}OPT(P) + \frac{1}{2}OPT(P')$.

Thus $OPT(P)$ is not globally convex in the feasible domain. \qed

**Theorem 1.** If $f(\cdot, \theta)$ is quasiconvex, then $OPT(\theta, P) = \min_{y \text{ feasible}} g_{P}(y, \theta)$ is quasiconvex in $P_i$ for any $1 \leq i \leq m$, where $P = [P_1, P_2, \ldots, P_m] \succeq 0$.  

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Proof. Let us assume $P = [p_1, p_2, \ldots, p_m]$ and $P' = [p'_1, p'_2, \ldots, p'_m]$, where $p_i = p'_i \forall i \neq 1$ with only the first column different. In the optimization problem parameterized by $P$, there is an optimal solution $x = \sum_{i=1}^m p_i y_i$, $y_i \geq 0 \forall i$. Similarly, there is an optimal solution $x' = \sum_{i=1}^m p'_i y'_i$, $y'_i \geq 0 \forall i$ for the optimization problem parameterized by $P'$. We know that $f(x) = h(P)$, $f(x') = h(P')$. Denote $P'' = cP + (1-c)P' = [p''_1, p''_2, \ldots, p''_m]$ to be a convex combination of $P$ and $P'$. Clearly, $p''_1 = cp_1 + (1-c)p'_1$ and $p''_i = p_i = p'_i \forall i \neq 1$. Then we can construct a solution
\[
x'' = \frac{1}{c} + \frac{1-c}{y_1} \left( \frac{c}{y_1} x + \frac{1-c}{y'_1} x' \right)
= \frac{1}{y_1} \left( cp_1 + (1-c)p'_1 \right) + \frac{1}{y_1} \sum_{i=2}^m \frac{p_i (y_i - y'_i)}{y_i y'_i}
\in \text{Span}(P'')
\]
Thus, $x''$ is a feasible solution in the optimization problem parameterized by $P''$. By the convexity of $f$, we also know that
\[
h(cP + (1-c)P') = h(P'') \leq f(x'')
= f(x'')
\leq \max(f(x), f(x'))
= \max(h(P), h(P'))
\]
When one of $y_1$, $y'_1$ is 0, without loss of generality we assume $y_1 = 0$. Then we can construct a solution $x'' = x$ which is still feasible in the optimization problem parameterized by $P'' = cP + (1-c)P'$. Then we have the following:
\[
h(P'') \leq f(x'') = f(x) = h(P) \leq \max(h(P), h(P'))
\]
which concludes the proof. \hfill \square

11 Sample Complexity of Learning Predictive Model in Surrogate Problem

**Theorem 2.** Let $\mathcal{H}_{\text{lin}}$ be the hypothesis class of all linear function mappings from $\xi \in \Xi \subset \mathbb{R}^p$ to $\theta \in \Theta \subset \mathbb{R}^n$, and let $P \in \mathbb{R}^{n \times m}$ be a linear reparameterization used to construct the surrogate. The expected Rademacher complexity over $t$ i.i.d. random samples drawn from $\mathcal{D}$ can be bounded by:
\[
\text{Rad}^t(\mathcal{H}_{\text{lin}}) \leq 2mC \sqrt{\frac{2p \log(2n^t || P^+ || \rho_2(S))}{t}} + O\left(\frac{1}{t}\right)
\]
where $C$ is the gap between the optimal solution quality and the worst solution quality, $\rho_2(S)$ is the diameter of the set $S$, and $P^+$ is the pseudoinverse.

The proof of Theorem 2 relies on the results given by Balghiti et al. Balghiti et al. analyzed the sample complexity of predict-then-optimize framework when the optimization problem is a constrained linear optimization problem.

The sample complexity depends on the hypothesis class $\mathcal{H}$, mapping from the feature space $\Xi$ to the parameter space $\Theta$. $x^*_S(\theta) = \arg\min_{x \in S} f(x, \theta)$ characterizes the optimal solution with given parameter $\theta \in \Theta$ and feasible region $S$. This can be obtained by solving any linear program solver with given parameters $\theta$. The optimization gap with given parameter $P$ is defined as $\omega_S(\theta) := \max_{x \in S} f(x, \theta) - \min_{x \in S} f(x, \theta)$, and $\omega_S(\Theta) := \sup_{\theta \in \Theta} \omega_S(\theta)$ is defined as the upper bound on optimization gap of all the possible parameter $\theta \in \Theta$, $x^*(\mathcal{H}) := \{ \xi \rightarrow x^*(\Phi(\xi)) | \Phi \in \mathcal{H} \}$ is the set of all function mappings from features $\xi$ to the predictive parameters $\theta = \Phi(\xi)$ and then to the optimal solution $x^*(\theta)$. 

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**Definition 1** (Natarajan dimension). Suppose that $S$ is a polyhedron and $\mathcal{S}$ is the set of its extreme points. Let $F \in \mathcal{S}^\mathcal{F}$ be a hypothesis space of function mappings from $\mathcal{X}$ to $\mathcal{S}$, and let $A \in \mathcal{X}$ to be given. We say that $F$ shatters $A$ if there exists $g_1, g_2 \in F$ such that

- $g_1(\xi) \neq g_2(\xi)$ $\forall \xi \in A$.
- For all $B \subset A$, there exists $g \in F$ such that (i) for all $\xi \in B$, $g(\xi) = g_1(\xi)$ and (ii) for all $\xi \in A \setminus B$, $g(\xi) = g_2(\xi)$.

The Natarajan dimension of $F$, denoted by $d_N(F)$, is the maximum cardinality of a set $N$-shattered by $F$.

We first state their results below:

**Theorem 3** (Balghiti et al. [11] Theorem 2). Suppose that $S$ is a polyhedron and $\mathcal{S}$ is the set of its extreme points. Let $\mathcal{H}$ be a family of functions mapping from features $\mathcal{X}$ to parameters $\Theta \in \mathbb{R}^n$ with decision variable $x \in \mathbb{R}^n$ and objective function $f(x, \theta) = \theta^T x$. Then we have that

$$\text{Rad}^d(\mathcal{H}) \leq \omega^*_S(\Theta) \sqrt{\frac{2d_N(x^*(\mathcal{H})) \log(t|\mathcal{S}|^2)}{t}}.$$  \hspace{1cm} (5)

where $\text{Rad}^d$ denotes the Rademacher complexity averaging over all the possible realization of $t$ i.i.d. samples drawn from distribution $D$.

The following corollary provided by Balghiti et al. [11] introduces a bound on Natarajan dimension of linear hypothesis class $\mathcal{H}$, mapping from $\mathcal{X} \in \mathcal{R}^p$ to $\Theta \in \mathcal{R}^n$:

**Corollary 1** (Balghiti et al. [11] Corollary 1). Suppose that $S$ is a polyhedron and $\mathcal{S}$ is the set of its extreme points. Let $\mathcal{H}_{lin}$ be the hypothesis class of all linear functions, i.e., $\mathcal{H}_{lin} = \{ \xi \rightarrow B\xi | B \in \mathcal{R}^{n \times p} \}$. Then we have

$$d_N(x^*(\mathcal{H}_{lin})) \leq np \hspace{1cm} (6)$$

Also $|\mathcal{S}|$ can be estimated by constructing an $\epsilon$-covering of the feasible region by open balls with radius $\epsilon$. Let $\hat{\mathcal{S}}_\epsilon$ be the centers of all these open balls. We can choose $\epsilon = \frac{1}{t}$ and the number of open balls required to cover $S$ can be estimated by

$$|\hat{\mathcal{S}}_\epsilon| \leq (2t \rho_2(S) \sqrt{m})^n \hspace{1cm} (7)$$

Combining Equation 6 and 7 the Rademacher complexity can be bounded by:

**Corollary 2** (Balghiti et al. [11] Corollary 2).

$$\text{Rad}^d(\mathcal{H}_{lin}) \leq 2m \omega_S(\Theta) \sqrt{\frac{2p \log(2nt\rho_2(S))}{t}} + O\left(\frac{1}{t}\right) \hspace{1cm} (8)$$

Now we are ready to prove Theorem 2.

**Proof of Theorem 2**. Now let us consider our case. We have a linear mapping from features $\xi \in X \subset \mathcal{R}^p$ to the parameters $\theta = B\xi \in \mathcal{R}^n$ with $B \in \mathcal{R}^{n \times p}$. The objective function is formed by

$$g_P(y, \theta) = f(Py, \theta) = \theta^T Py = (P^T \theta)^T y = (P^T B\xi)^T y \hspace{1cm} (9)$$

This is equivalent to have a linear mapping from $\xi \in \mathcal{X} \subset \mathcal{R}^p$ to $\theta' = P^T B\xi$ where $P^T B \in \mathcal{R}^{m \times p}$, and the objective function is just $g_P(y, \theta') = \theta'^T y$. This yields a similar bound but with a smaller dimension $m \ll n$ as in Equation 10.

$$|\text{Rad}^d(\mathcal{H}_{lin})| \leq 2m \omega_S(\Theta) \sqrt{\frac{2p \log(2nt\rho_2(S'))}{t}} + O\left(\frac{1}{t}\right) \hspace{1cm} (10)$$

where $\omega_S(\Theta)$ is unchanged because the optimality gap is not changed by the reparameterization. The only thing changed except for the substitution of $m$ is that the feasible region $S'$ is now defined in a
lower-dimensional space under reparameterization $P$. But since $\forall y \in S'$, we have $Py \in S$ too. So the diameter of the new feasible region can also be bounded by:

$$
\rho(S') = \max_{y, y' \in S'} \|y - y'\| \\
= \max_{y, y' \in S'} \|P^+P(y - y')\| \\
= \max_{y, y' \in S'} \|P^+(Py - Py')\| \\
\leq \max_{x, x' \in S'} \|P^+(x - x')\| \\
= \|P^+\| \rho(S)
$$

where $P^+ \in \mathbb{R}^{m \times n}$ is the pseudoinverse of the reparameterization matrix $P$ with $P^+P = I \in \mathbb{R}^{m \times m}$ (assuming the matrix does not collapse). Substituting the term $\rho(S')$ in Equation 10, we can get the bound on the Radamacher complexity in Equation 4, which concludes the proof of Theorem 2.

### 12 Non-linear Reparameterization

The main reason that we use a linear reparameterization is to maintain the convexity of the inequality constraints and the linearity of the equality constraints. Instead, if we apply a convex reparameterization $x = P(y)$, e.g., an input convex neural network [3], then the inequality constraints will remain convex but the equality constraints will no longer be affine anymore. So such convex reparameterization can be useful when there is no equality constraint. Lastly, we can still apply non-convex reparameterization but it can create non-convex inequality and equality constraints, which can be challenging to solve. All of these imply that the choice of reparameterization should depend on the type of optimization problem to make sure we do not lose the scalability while solving the surrogate problem.

### 13 Computing Infrastructure

All experiments were run on the computing cluster, where each node configured with 2 Intel Xeon Cascade Lake CPUs, 184 GB of RAM, and 70 GB of local scratch space. Within each experiment, we did not implement parallelization. So each experiment was purely run on a single CPU core. The main bottleneck of the computation is on solving the optimization problem, where we use Scipy [41] blackbox optimization solver. No GPU was used to train the neural network and throughout the experiments.