Efficient Policy Learning from Surrogate-Loss Classification Reductions

Andrew Bennett 1  Nathan Kallus 1

Abstract

Recent work on policy learning from observational data has highlighted the importance of efficient policy evaluation and has proposed reductions to weighted (cost-sensitive) classification. However, efficient policy evaluation need not yield efficient estimation of optimal policy parameters. We consider the estimation problem given by a weighted surrogate-loss classification reduction of policy learning with any score function — either direct, inverse-propensity weighted, or doubly robust — and show that, under a correct specification assumption, the weighted classification formulation need not be efficient for policy parameters. We draw a contrast to actual (possibly weighted) binary classification, where correct specification implies a parametric model, while for policy learning it only implies a semiparametric model, and we show that efficiency in optimal parameter estimation implies optimal regret. In light of this, we instead propose an estimation approach based on the generalized method of moments, which is efficient for the policy parameters. We propose a particular method based on recent developments on solving moment problems using neural networks and demonstrate the efficiency and regret benefits of this method empirically.

1. Introduction

Policy learning from observational data is an important but challenging problem because it requires reasoning about the effects of interventions not observed in the data. For example, if we wish to learn an improved policy for medical treatment assignment based on observational data from electronic health records, we must take care to consider potential confounding: since healthier patients who were already predisposed to positive outcomes were likely to have historically been assigned less invasive treatments, naïve approaches may incorrectly infer that a policy of always assigning less invasive treatments will obtain better outcomes.

Various recent work has tackled this problem, known as policy learning from observational (or, off-policy) data, by optimizing causally-grounded estimates of policy value such as inverse-propensity weighting (IPW), doubly robust (DR) estimates, or similar (Beygelzimer & Langford, 2009; Jiang et al., 2019; Kallus, 2017; 2018; Kallus & Zhou, 2018; Kitagawa & Tetenov, 2018; Qian & Murphy, 2011; Swamnathan & Joachims, 2015; Zhao et al., 2012; Zhou et al., 2017). In particular, Athey & Wager (2017); Zhou et al. (2017), among others, highlight the importance of using efficient estimates of policy value as optimization objectives, i.e., having minimal asymptotic mean-squared error (MSE). Examples of efficient estimators are direct modeling or IPW when outcome functions or propensities are sufficiently smooth (Hahn, 1998; Hirano et al., 2003), or DR leveraging cross-fitting (Chernozhukov et al., 2003) in more general non-parametric settings.

Regardless of which of these three estimates one uses, the resulting optimization problem amounts to a difficult binary optimization problem. Therefore many of the above leverage a reduction of this problem to weighted classification (for two actions; cost-sensitive classification more generally) and leverage tractable convex formulations that use classification surrogate loss functions for the zero-one loss, such as, for example, hinge loss (Zhao et al., 2012; Zhou et al., 2017, which yields a weighted SVM) and logistic loss (Jiang et al., 2019, which yields a weighted logistic regression). The recently proposed entropy learning approach of Jiang et al. (2019) is particularly appealing, since the logistic regression-based surrogate loss functions for the zero-one loss, such as, for example, hinge loss (Zhao et al., 2012; Zhou et al., 2017, which yields a weighted SVM) and logistic loss (Jiang et al., 2019, which yields a weighted logistic regression). The recently proposed entropy learning approach of Jiang et al. (2019) is particularly appealing, since the logistic regression-based surrogate loss is smooth and therefore allows for statistical inference on the estimated optimal parameters. In general however, one may consider using any surrogate loss function that is classification-calibrated (Bartlett et al., 2006), meaning that any policy that minimizes the surrogate loss is optimal.

However, as we here emphasize, even if we use policy value estimates that are efficient, this does not imply that we obtain efficient estimation/learning of the optimal policy itself, even if the surrogate-loss model is well-specified. For example, in the case of logistic loss, we demonstrate that, although logistic regression is statistically efficient for actual
binary classification when well-specified (as is well-known), in the case of policy learning via a weighted-classification reduction well-specification only implies a semi-parametric model, and therefore minimizing the empirical average of the surrogate loss is not efficient in this case.

On the other hand, the implications of correct specification can be summarized as a conditional moment problem. Such problems are amenable to efficient solution using approaches based on the generalized method of moments (GMM; Hansen, 1982). We demonstrate what an efficient such estimate would look like, in terms of the efficient instruments for our specific policy learning problem. We propose a particular implementation of solving our problem based on recent work on efficiently solving conditional moment problems using a reformulation of the efficient GMM solution as a smooth game optimization problem, which can be solved using adversarial training of neural networks (Bennett et al., 2019). In addition, we prove some results relating the efficiency of optimal policy estimation to the asymptotic regret of the surrogate loss, and also prove that under correct specification the regret of the surrogate loss upper bounds the true regret of policy learning.

We demonstrate empirically over a wide range of scenarios that our methodology indeed leads to greater efficiency, with lower MSE in estimating the optimal policy parameter estimates under correct specification. Furthermore, we demonstrate that in practice, both with and without correct specification, our methodology tends to learn policies with lower regret, particularly in the low-data regime.

1.1. Setting and Assumptions

Let $X$ denote the context of an individual, $T \in \{-1, 1\}$ the treatment assigned to that individual, and $Y$ the resultant outcome. In addition let $Y(t)$ denote the counterfactual outcome that would have been obtained for the corresponding individual if, possibly contrary to fact, treatment $t$ had been assigned instead. We assume throughout that we have access to logged data consisting of $n$ iid observations, $S_n = \{(X_i, T_i, Y_i) : i \leq n\}$, of triplets $(X, T, Y)$ generated by some behavior policy.

We make standard causal assumptions of consistency and non-interference, which can be summarized by assuming that $Y = Y(T)$. Furthermore, as is standard in the above policy learning literature, we assume that $X$ encapsulates all possible confounders, that is, $Y(t) \perp T \mid X \forall t \in \{-1, 1\}$, as would for example be guaranteed if the logging policy is a function of the observed individual context.

A policy $\pi$ denotes a mapping from individual context to treatment to be assigned. Concretely, given individual context $x$, let $\pi(x) \in \{-1, +1\}$ denote the treatment assigned by policy $\pi$ (we may also consider stochastic policies but since optimal policies are deterministic we focus on these).

Let

$$J(\pi) = \mathbb{E}[Y(\pi(X))] - \frac{1}{2}\mathbb{E}[Y(+1) + Y(-1)]$$

$$= \mathbb{E}[\pi(X)(Y(+1) - Y(-1))]$$

denote the expected value of following policy $\pi$, relative to complete randomization. Given the logged data and some policy class $\Pi$, our task is to learn an optimal policy from the class, defined by $\pi^* \in \arg \max_{\pi \in \Pi} J(\pi)$ (notice that offsetting by the complete randomization policy does not affect this optimization problem). In particular we consider policy classes where each policy $\pi$ is indexed by some utility function $g$ and is defined by $\pi(x) = \text{sign}(g(x))$, where in turn the utility functions are parametrized by $\theta \in \Theta \subseteq \mathbb{R}^d$ as $\mathcal{G} = \{g_\theta : \theta \in \Theta\}$, so that

$$\Pi = \{\text{sign}(g_\theta(x)) : \theta \in \Theta\}.$$ Correspondingly, we define

$$J(\theta) = J(\text{sign}(g_\theta(\cdot))) = \mathbb{E}[\text{sign}(g_\theta(X))(Y(+1) - Y(-1))]$$

and $\theta^* \in \arg \max_{\theta \in \Theta} J(\theta)$. A prominent example is linear decision rules, where $g_\theta(x) = \theta^T x$. Other examples include decision trees of bounded depth and neural networks.

Unlike some past work that has considered non-parametric policy classes (e.g. Zhao et al. (2012)), which have advantages in terms of regret without having to rely on “correct specification” assumptions, we make the decision to focus on parametric policy classes only. This is because these classes are more amenable to efficiency analysis, and are very relevant in practice due to reasons such as interpretability and implementability.

1.2. Efficiency

We briefly review what it means to estimate the optimal policy parameters, $\theta^*$, efficiently. For simplicity, suppose that $\theta^*$ is unique. A model $\mathcal{M}$ is some set of distributions for the data-generating process (DGP), i.e., a set of probability distributions for the triplet $(X, T, Y)$. A model is generally non-parametric in the sense that this set of distributions can be arbitrary, infinite, and infinite dimensional.

Consider any learned policy parameters $\hat{\theta}$, that is, a function of the data $\mathcal{S}_n$ with values in $\Theta$. Roughly speaking, we say that $\hat{\theta}$ is regular if, whenever the data is generated from $(X_i, T_i, Y_i) \sim p \in \mathcal{M}$, we have that $\sqrt{n}(\hat{\theta} - \theta^*)$ converges in distribution to some limit as $n \to \infty$ and this limit holds in a particular locally uniform sense in $\mathcal{M}$ (see Van der Vaart, 2000, Chapter 25 for a precise definition). Semiparametric efficiency theory (see ibid.) then establishes that there exists a covariance matrix $V$ such that for any cost function $c : \mathbb{R}^d \to \mathbb{R}$ for which the sublevel sets are
\[ \{ v : c(v) \leq c_0 \} \text{ are convex, symmetric about the origin, and closed, we have that} \]
\[ \liminf_{n \to \infty} E[c(\sqrt{n}(\hat{\theta} - \theta^*))] \geq E_{\nu \sim \mathcal{N}(0, V)}[c(v)] \]  
(1)

for any estimator \( \hat{\theta} \) that is regular in \( \mathcal{M} \). An important example is MSE, given by \( c(v) = ||v||^2_2 \).

**Efficient estimators** are those for which Eq. (1) holds with equality for all such functions \( c \), which, by the portmanteau lemma, would be implied if the estimator has the limiting law \( \sqrt{n}(\hat{\theta} - \theta^*) \Rightarrow \mathcal{N}(0, V) \). Regular estimators is a very general class of estimators so the bound in Eq. (1) is rather strong. So much so that, in fact, Eq. (1) holds in a local asymptotic minimax sense for all estimators (see ibid., Theorem 25.21).

Efficiency is important for practitioners because, in observational data, we only have the data that we have and cannot experiment or simulate to generate more data. So we should use the data optimally. Amongst other things, efficiency implies we can construct optimally-tight confidence intervals for the estimated optimal parameter values, and in Section 4.4 we show that efficiency implies asymptotically-optimal regret.

### 1.3. Related Work

There has been a variety of past work on the problem of policy learning from observational data. Much of this work considers formulating the objective of policy learning as a weighted classification problem (Beygelzimer & Langford, 2009; Dudik et al., 2011), and either minimizing the 0-1 loss directly using combinatorial optimization (Athey & Wager, 2017; Kitagawa & Tetenov, 2018; Zhou et al., 2018), using smooth stochastic policies to obtain a nonconvex but smooth loss surface (Swaminathan & Joachims, 2015), or replacing the 0-1 objective with a convex surrogate to be minimized instead (Beygelzimer & Langford, 2009; Dudik et al., 2011; Jiang et al., 2019; Zhao et al., 2012; Zhou et al., 2017).

In addition there has been a variety of past work on solving conditional moment problems (see Bennett et al. (2019); Khosravi et al. (2019) and citations therein). Our paper builds on this work as it reformulates the problem of policy learning as a conditional moment problem, which we propose to solve using optimally weighted GMM (Hansen, 1982) and DeepGMM (Bennett et al., 2019).

### 2. The Surrogate-Loss Reduction and Its Fisher Consistency

In this section, we present the surrogate-loss reduction of policy learning and the implications of correct specification.

Many policy learning methods start by recognizing that the policy value can be re-written as
\[ J(\theta) = E[\psi \text{sign}(g_\theta(X))] \]  
(2)

where \( \psi \) is any of the following score variables, which all depend on observables:
\[ \psi_{PS} = \frac{TY}{\varepsilon_T(X)}, \quad \psi_{DM} = \mu_1(X) - \mu_{-1}(X), \quad \psi_{DR} = \psi_{DM} + \psi_{PS} - \frac{T\mu_T(X)}{\varepsilon_T(X)}, \]  
(3)

where \( e_t(x) = P(T = t | X = x) \) and \( \mu_t(x) = E[Y | t, X = x] \). Equation (2) arises once we recognize that all of these satisfy \( E[\psi | X] = E[Y(1) - Y(-1)] | X] \).

Then we can approximate Eq. (2) using its empirical version:
\[ J_n(\theta) = \frac{1}{n} \sum_{i=1}^n \psi_i \text{sign}(g_\theta(X_i)). \]  
(4)

In particular, Athey & Wager (2017); Kitagawa & Tetenov (2018); Zhou et al. (2018) prove bounds of the form \( \sup_{\theta \in \Theta} | J_n(\theta) - J(\theta) | = O_p(1/\sqrt{n}) \) given that the policy class has bounded complexity. This shows that optimizing \( \theta \in \arg \max_{\theta \in \Theta} J_n(\theta) \) provides near-optimal solutions in the original policy learning problem, since \( J(\theta^*) - J(\theta) \leq J(\theta) - J_n(\theta) - J_n(\theta^*) \leq 2 \sup_{\theta \in \Theta} | J_n(\theta) - J(\theta) | \).

Given that in practice the nuisance functions \( e_t \) and \( \mu_t \) are estimated from data, we denote the corresponding score variable when such estimates are plugged in as \( \tilde{\psi} \) to differentiate it from the variable \( \psi \) that uses the true nuisance functions. We correspondingly let \( \tilde{J}_n(\theta) = \frac{1}{n} \sum_{i=1}^n \tilde{\psi}_i \text{sign}(g_\theta(X_i)) \). When \( \tilde{J}_n(\theta) \) is efficient for \( J(\theta) \) one can generally additionally prove that \( \sup_{\theta \in \Theta} | \tilde{J}_n(\theta) - J_n(\theta) | = O_p(1/\sqrt{n}) \).

Given the non-convexity and non-smoothness of the empirical objective function Eq. (4), it is not necessarily clear how to actually optimize it. Many works (Beygelzimer & Langford, 2009; Jiang et al., 2019; Zhao et al., 2012) recognize that this optimization problem is actually equivalent to weighted binary classification (in our two-action case), since \( \psi_i \text{sign}(g_\theta(X_i)) = |\psi_i|(1 - 2\text{sign}(g_\theta(X_i)) \neq \psi_i) \), so any classification algorithm that accepts instance weights can
Theorem 1 establishes that, under correct specification, if we minimize the population surrogate loss, \( L(\theta) \), then we obtain the optimal policy. Therefore, a natural strategy for policy learning would be to directly minimize the empirical loss \( L_n(\theta) \), as was done by the above. Although the above arguments indicate that this approach would be computationally tractable, and also consistent under mild regularity conditions that ensure that optimizers of \( L_n(\theta) \) would converge to optimizers of \( L(\theta) \), it is not clear that it is statistically efficient, even if we use an efficient score variable for policy value estimation.

We provide a cautionary note that, as discussed e.g. in Qiu et al. (2019); Wager (2020), hoping for correct specification to hold for a simple low-dimensional policy class such as linear policies could be unreasonable. In such a case, it is possible that using this surrogate objective could systematically lead to incorrect policy decisions (Wager, 2020). On the other hand, if we use a flexible policy class such as neural networks of a given architecture, it is reasonable to assume that correct specification holds (at least approximately), so the surrogate objective may be better justified.

3. The Conditional-Moment Reformulation of the Surrogate-Loss Reduction

In this section we establish a new interpretation of the surrogate-loss reduction as a conditional moment problem and we discuss the implications of this in terms of the model implied by correct specification. This will enable us to conduct efficiency analysis and to design algorithms with improved efficiency in the next section.

3.1. The Conditional Moment Problem

First, we define \( l' \) as the derivative of \( l \) with respect to its first argument. In the case of logistic regression loss this is

\[
l'(g, s) = 2\sigma(g) - (s + 1)g.
\]

where \( \sigma(g) = \exp(g) / (1 + \exp(g)) \) is the logistic function.

Theorem 2 (Conditional Moment Problem Under Correct Specification). Suppose Assumption 2 holds and the policy class \( \Pi \) is correctly specified for the surrogate loss in the sense that

\[
m(X; \theta) = \mathbb{E}[\psi|l'(g_\theta(x), \text{sign}(\psi)) | X].
\]

Then we have that

\[
\theta^* \in \arg\min_{\theta \in \Theta} L(\theta) \iff m(X; \theta^*) = 0 \text{ almost surely. (8)}
\]

Theorem 2 arises straightforwardly from the observation that, under correct specification, \( g_\theta(x) \) minimizes
\[ \mathbb{E}[\psi_i l(g_\theta(X), \text{sign}(\psi)) \mid X = x] \text{ for almost every } x. \] Using smoothness and convexity, this latter observation is restated using first-order optimality conditions. Dominated convergence theorem allows us to exchange differentiation and expectation and we obtain the result. Theorem 2 provides an alternative characterization of \( \theta^* \) as solving a conditional moment problem.

Notice that Eq. (8) is equivalent to the statement that, for any square integrable function \( f \) of \( X \), we have the moment restriction
\[ \mathbb{E}[m(X; \theta)f(X)] = 0. \tag{9} \]
This alternative characterization makes the problem amenable to efficiency analysis.

Notice that by first-order optimality, if \( \theta^* \in \text{interior}(\Theta) \), optimizing \( L(\theta) \) in Eq. (6) exactly corresponds to solving the set of \( d \) moment equations given by \( \mathbb{E}[m(X; \theta)\nabla_\theta g_\theta(x)] = 0 \). Similarly, optimizing the empirical loss \( L_n(\theta) \) in Eq. (5) corresponds to solving these \( d \) equations with population averages (\( \mathbb{E} \)) replaced with empirical sample averages.

However, Eq. (9) gives a much broader set of equations. Leveraging this fact will be crucial to achieving efficiency. Indeed, it is well-known that even if a small number of moment equations are sufficient to identify a parameter (e.g., in the above, the \( d \) equations identify \( \theta^* \) via first-order optimality), taking into consideration additional moment equations that are known to hold can increase efficiency in semiparametric settings (Carrasco & Florens, 2014).

### 3.2. The Semiparametric Model Implied by Specification

In order to reason about efficiency, we need to reason about the model implied by Eq. (8). To do so, we first establish the following lemma.

**Lemma 1.** Assume Assumption 2, and that we are using logistic regression loss. Then given a policy class \( \Pi \), the model of DGPs (distributions on \( (X, T, Y) \)) where \( \Pi \) is correctly specified for the surrogate loss (in the sense of Eq. (7)) is given by all distributions on \( (X, T, Y) \) for which there exists \( \theta^* \in \Theta \) satisfying
\[ \frac{\mathbb{E}[\psi \mathbb{I}\{\psi > 0\} \mid X = x]}{\mathbb{E}[\psi \mid X = x]} = \sigma(g_{\theta^*}(X)) \text{ almost surely.} \tag{10} \]

This model is generally a semiparametric model. That is, while Eq. (10) is a parametric restriction on the function \( \mathbb{E}[\psi \mathbb{I}\{\psi > 0\} \mid X = x]/\mathbb{E}[\psi \mid X = x] \), the set of corresponding distributions on \( (X, T, Y) \) that satisfy this restriction is still infinite-dimensional and non-parametric.

---

1. This is because in an inner product space \( V \), \( v = 0 \) if and only if \( \langle v, v' \rangle = 0 \) for every \( v' \in V \). Here \( V = L_2 \).

---

### 3.3. Comparison with Logistic Regression for Classification

One question the reader might have at this point is why an approach different than empirical loss minimization is necessary for efficiency, given that the surrogate loss formulation seems mathematically identical to binary classification using logistic regression, which is known to be efficient.\(^2\) The difference between the problems is that for actual classification we have that \( \psi \) is a binary class label, i.e., \( \psi \in \{-1, 1\} \). If we assume the policy class is well-specified and \( \psi \in \{-1, 1\} \), the characterization of our semiparametric model from Lemma 1 reduces to
\[ P(\psi = 1 \mid X) = \sigma(g_{\theta^*}(X)), \]
which implies that our model is parametric, since the choice of \( \theta^* \) now fully characterizes the distribution of the label \( \psi \) given \( X \). E.g., usually for logistic regression we get \( g_{\theta}(x) = \theta^T x \) so that the above says that the logit of \( P(\psi = 1 \mid X) \) is linear. Therefore, performing logistic regression corresponds to MLE for this parametric model, which is efficient.

However in our more general setting this is not the case and there is a non-trivial nuisance space, since there is a complex, infinite-dimensional space of conditional distributions for \( \psi \) given \( X = x \) that could result in the same function \( \mathbb{E}||\psi||(\psi > 0) \mid X = x)/\mathbb{E}||\psi|| \mid X = x \). This suggests that we may need to be more careful in order to obtain efficiency and that there may exist estimators that are more efficient than empirical loss minimization.

### 4. Efficient Policy Learning Reductions

In this section we propose some efficient methods for policy learning based on the above conditional-moment formulation. In addition, we provide some analysis of these methods in terms of efficiency and regret.

#### 4.1. FiniteGMM Policy Learner

We begin by proposing an approach based on using multi-step GMM to solve the conditional moment problem, which we will call **FiniteGMM**. This approach works by optimally enforcing for the moment conditions given by Eq. (9) for a finite collection of critic functions \( F = \{f_1, \ldots, f_k\} \).

Specifically, given some initial estimate \( \theta_0 \), define:
\[ m(\theta) = \frac{1}{n} \sum_{i=1}^n \psi_i l'(g_\theta(X_i), \text{sign}(\psi_i)) f_i(X_i) \]
\[ C(\tilde{\theta}_n)_{jk} = \frac{1}{n} \sum_{i=1}^n \psi_i^2 l''(g_{\theta}(X_i), \text{sign}(\psi_i))^2 f_j(X_i) f_k(X_i) \]
\[ O(\theta; \tilde{\theta}_n) = m(\theta)^T C(\tilde{\theta}_n) m(\theta). \]

\(^2\)This is because logistic regression performs maximum likelihood estimation (MLE), which is statistically efficient for well-specified parametric models.
We then estimate \( \theta \) by \( \hat{\theta}_n = \arg \min_{\theta} O(\theta; \hat{\theta}_n) \). We can repeat this multiple times, plugging in \( \hat{\theta}_n \) as \( \hat{\theta}_n \) and resolving.

An important issue with this estimator, however, is how to choose the critic functions. Standard GMM theory requires that the \( k \) moment conditions are sufficient to identify \( \theta^* \). And even then, the above is only the most efficient among estimators of the form \( \arg \min_{\theta} \| (m_1(\theta), \ldots, m_k(\theta)) \| \) for any norm \( \| \cdot \| \), but there may still be more efficient choices of critic functions.

4.2. The Efficient Instruments for Policy Learning

One nice result from the theory of conditional moment problems is the existence of a finite set of critic functions ensuring efficiency in the sense of Section 1.2. Define:

\[
\begin{align*}
\Omega(x) &= E[\psi^2 l'(g_\theta(X), \text{sign}(\psi))^2 \mid X = x] \\
h_{\theta^*}(x) &= \nabla g_\theta(x) \big|_{\theta = \theta^*} \\
D(x) &= E[\nabla_\theta (|\psi| l'(g_\theta(X), \text{sign}(\psi)))]_{\theta = \theta^*} \mid X = x \\
f^*_i(x) &= \frac{D(x)}{\Omega(x)}.
\end{align*}
\]

We call \( F^* = \{f^*_1, \ldots, f^*_d\} \) the efficient instruments, and as long as the span of \( F \) contains these instruments then \( \text{FINITEGMM} \) is guaranteed efficiency (Newey, 1993). We can observe that these equations correspond to linearizing the moment equation at \( \theta^* \), and \( \Omega \) is akin to the Fisher information matrix. We refer readers to Newey (1993) for more details on the derivation of these efficient instruments.

Given this, one approach would be to let \( F \) be flexible with the hope of approximately containing \( F^* \). Letting, for example, \( F \) be the first \( k(n) \) functions in a basis for \( L_2 \) such as a polynomial basis and letting \( k(n) \to \infty \) can be shown to be efficient under certain conditions (Newey, 1993). This, however, can perform very badly in practice, especially when the number of features is high, due to known curse of dimensionality issues with such classical nonparametric regression methods (Bauer et al., 2019; Geenens et al., 2011; Nagler & Czado, 2016).

4.3. ESPRM Policy Learner

Motivated by the above concerns, we now present our proposed approach: ESPRM (efficient surrogate policy risk minimization). This is based on the extension of Bennett et al. (2019) to our conditional moment problem. In the setting of instrumental variable regression, Bennett et al. (2019) proposes an adversarial reformulation of optimally-weighted GMM, which allows us to consider critic functions given by flexible classes such as neural networks. Then if this class provides a good approximation for the efficient instruments, this approach should be approximately efficient.

Specifically, we define:

\[
\begin{align*}
&u(X, \psi; \theta, f) = |\psi| l'(g_\theta(X), \text{sign}(\psi)) f(X) \\
&U(\theta, f; \hat{\theta}) = \frac{1}{n} \sum_{i=1}^{n} u(X_i, \hat{\psi}_i; \hat{\theta}, f) \\
&\hat{\theta}_{\text{ESPRM}} = \arg \min_{\theta} \sup_{f \in \mathcal{F}} U(\theta, f; \hat{\theta}),
\end{align*}
\]

where as above \( \hat{\theta}_n \) is some initial consistent estimate of \( \theta^* \). Then following Bennett et al. (2019), the ESPRM estimator is defined as

\[
\hat{\theta}_{\text{ESPRM}} = \arg \min_{\theta} \sup_{f \in \mathcal{F}} U(\theta, f; \hat{\theta}),
\]

where \( \mathcal{F} \) is our flexible function class (henceforth assumed to be a class of neural networks). In brief, the motivation of this objective is as follows. First, if we let \( \mathcal{F} = \text{span}\{f_1, \ldots, f_k\} \) in this objective, it follows from a generalization of Bennett et al. (2019, Lemma 1) that ESPRM is identical to \( \text{FINITEGMM} \) using critic functions \( \{f_1, \ldots, f_k\} \). It follows that ESPRM is equivalent to replacing the span of the critic functions in \( \text{FINITEGMM} \) with a generic function space. Therefore, instead of trying to approximate the efficient instrument using a growing basis for \( L_2 \) — an approach which is known to suffer from curse of dimensionality issues — we can instead use a flexible function space, such as a space of neural networks, that is designed to handle flexible function approximation without such issues (Bauer et al., 2019). We describe the theory behind this estimator in more detail in the appendix.

It remains to describe how this adversarial game is to be solved, and how to define \( \hat{\theta}_n \). As in Bennett et al. (2019) we optimize the objective by performing alternating first-order optimization steps using the OAdam algorithm (Daskalakis et al., 2017), which was designed for solving smooth game problems such as generative adversarial networks (GANs). In addition, we continuously update \( \hat{\theta}_n \) during optimization, where at each step of alternating first order optimization we set \( \hat{\theta}_n \) equal to the previous iterate of \( \hat{\theta}_n \).

4.4. Efficient Learning implies Optimal Regret

Finally we prove that efficiency not only ensures minimal MSE in estimating \( \theta^* \) but also implies regret bounds. Let

\[
\begin{align*}
\text{Regret}_L(\theta) &= \arg \max_{\pi \text{ unconstrained}} J(\pi) - J(\theta) \\
\text{Regret}_L(\theta) &= L(\theta) - \inf_{\delta \in \Theta} L(\theta).
\end{align*}
\]

Theorem 3 (Regret Upper Bound). Suppose Assumptions 1 and 2 hold and that the policy class II is correctly specified for the surrogate loss in the sense that Eq. (7) holds. Then, for any \( \theta \in \Theta \) we have

\[
\varphi(\text{Regret}_L(\theta)) \leq \text{Regret}_L(\theta),
\]

for some continuous \( \varphi \) that depends only on \( l \), and satisfies

\[
\varphi(0) = 0 \text{ and } \varphi(\alpha) > 0 \text{ for } \alpha > 0.
\]

Furthermore, in the
We note that this assumption is very strong, and may be unrealistic for classes such as neural networks where multiple different parameter values can correspond to the same function (for example by permuting the units in hidden layers). However, we argue that in practice this is not a major issue, and can be addressed for example by using symmetry-breaking constraints.

Given Assumptions 1 and 3, a Taylor’s theorem expansion yields \( \text{Regret}_L(\hat{\theta}_n) = (\hat{\theta}_n - \theta^*)^T H(\theta^*) (\hat{\theta}_n - \theta^*) + o(\|\hat{\theta}_n - \theta^*\|^2) \), where \( H(\theta^*) \) is the Hessian of \( L \) at \( \theta^* \). For any regular estimator \( \hat{\theta}_n \), we can also define the asymptotic regret \( \text{AR}_L \) as the limiting distribution:

\[
\frac{n\text{Regret}_L(\hat{\theta}_n)}{d} \rightarrow_d \text{AR}_L(\hat{\theta}_n),
\]

which exists since regularity implies that \( \sqrt{n}(\hat{\theta}_n - \theta^*) \) has a limiting distribution. Given this we can prove the following optimality result of our efficient estimators in terms of asymptotic regret:

**Theorem 4 (Optimal Asymptotic Regret).** Given Assumption 3 and any non-negative, non-decreasing \( \phi \), we define the risk \( R_\phi(\hat{\theta}_n) = \mathbb{E}[\phi(\text{AR}_L(\hat{\theta}_n))] \). Given this, there exists a risk bound \( B_\phi \) such that \( R_\phi(\hat{\theta}_n) \geq B_\phi \) for every regular \( \hat{\theta}_n \) with equality if \( \hat{\theta}_n \) is semi-parametrically efficient.

Together with Theorem 3, this means that both the actual regret (\( \text{Regret}_f \)) and the surrogate regret (\( \text{Regret}_L \)) of policies given by efficient estimators \( \hat{\theta} \) are \( O_p(1/n) \), and the surrogate regret has an optimal constant.

Note that this is not the first regret result for policy learning based on convex surrogate losses. Of particular interest, Zhao et al. (2012) previously provided optimal regret results for nonparametric policy learning using hinge loss. However, unlike us they showed that their regret obtains an optimal rate but not optimal constants, and their result depends on various additional technical assumptions.

5. Experiments

5.1. Synthetic Scenarios

First we investigate the performance of our algorithms on a variety of synthetic scenarios, using logistic regression loss for \( l \). In all these scenarios \( X \) is 2-dimensional, and \( X \) and \( Y(t) - \mu_t(X) \) are standard Gaussian distributed for each \( t \); the scenarios only differ in the functions \( \mu_t \) and \( e_t \). In none of the scenarios is our policy class actually well-specified in the sense of Eq. (7).

We consider the following kinds of synthetic scenarios:

- **LINEAR:** \( \mu_t(x) = a^T x + a_0 \) and \( e_1(x) = \text{sigmoid}(b^T x + b_0) \) for some vectors \( a_{-1}, a_1, b \).
- **QUADRATIC:** \( \mu_t(x) = x^T A_t x + a_1^T x + a_0 \) and \( e_1(x) = \text{sigmoid}(x^T B x + b_1 x + b_0) \) for some symmetric matrices \( A \) and \( B \), and vectors \( a_{-1}, a_1, b \).

In addition we experiment with the following policy classes: a linear policy class where \( g_0(x) = \theta^T x + \theta_0 \), and a flexible policy class where \( g_0(x) \) is given by a fully-connected neural network with a single hidden layer of size 50, and leaky ReLU activations.

In all cases we use the surrogate loss method of Jiang et al. (2019) described in Section 2 as a benchmark, which we henceforth refer to as ERM. We note that although in the prior work they used \( \hat{\psi}_{PIS} \), we instead use \( \hat{\psi}_{PIS} \), both because it is theoretically better grounded given its double robustness property (Athey & Wager, 2017; Zhou et al., 2017) and we found that it gives stronger results for all methods. For our ESPR method we let \( F \) be the same neural network function class as for flexible policies, and perform alternating first-order optimization as described in Section 4.3 for a fixed number of epochs. For \( \text{FiNITEGMM} \) we experimented with function sets \( F \) based on various polynomial basis expansions, and also various finite-dimensional approximations of Gaussian kernel basis expansions using the method of Random Kitchen Sinks (Rahimi & Recht, 2009). We provide details of these function sets in the appendix.

For all methods, except where otherwise specified, we use the \( \hat{\psi}_{DR} \) weights described in Eq. (3), with nuisance functions fit using correctly specified linear regression or logistic regression algorithms on a separately sampled tuning dataset of the same size as the training dataset.\(^3\) We provide some additional results in the appendix where nuisances were instead fit via flexible neural networks, which show that this has little effect on our results. In all cases except for ESPRM we perform optimization using LBFGS. Additional optimization details are given in the appendix.

For all configurations of scenario kind and policy we ran our experiments by sampling random scenarios of the respective kind, by setting all scenario parameters to be independent standard Gaussian variables. Specifically, for each

\(^3\)By correctly specified we mean that for LINEAR we fit using linear/logistic regression on \( X \), whereas for QUADRATIC we fit on a quadratic feature expansion of \( X \).
Efficient Policy Learning from Surrogate-Loss Classification Reductions

Figure 1. Difference in performance between ESPRM and ERM. We plot RMRR against training set size for each combination of policy class and scenario kind. Shaded regions are 95% confidence intervals, constructed from bootstrapping using the 64 replications.

$n \in \{100, 200, 500, 1000, 2000, 5000, 10000\}$ we sample 64 random scenarios of the respective kind, and for each random scenario we sample $n$ training data points and run all methods on this data. Results for $\text{FINITEGMM}$, which generally did badly as predicted for all the basis function sets described above, are given in the appendix.

Define Relative Mean Regret Reduction (RMRR), given by:

$$RMRR(\hat{\theta}_n) = \left( 1 - \frac{\mathbb{E}[\text{Regret}(\hat{\theta}_n)]}{\mathbb{E}[\text{Regret}(\hat{\theta}_{\text{ERM}})]} \right) \times 100\%,$$

where each expectation in the fraction is taken over the joint distribution of randomly sampled scenarios, and the corresponding random estimates $\hat{\theta}$. Then for each scenario kind and policy class, we plot predicted RMRR against number of training data based on our ESPRM estimates in Fig. 1. In addition, we provide plots for this experiment on a raw utility scale in the appendix. Note that the confidence intervals in all these plots for each data point are with respect to the joint distribution over randomly generated scenarios and corresponding random estimate $\hat{\theta}$.

We see that ESPRM consistently obtains policies on average that are lower regret or on-par than those obtained by ERM (typically with around 10% to 20% RMRR), with the 95% confidence intervals indicating clearly better performance in almost every case, except for in the case of training flexible policies in the quadratic scenario where there are a couple of outlier points, although even there the two methods seem at worst roughly on-par. We can also observe that the most significant regret benefits tend to occur with smaller training set sizes (since the same RMRR implies a larger absolute decrease in regret), indicating that the statistical efficiency of our method is leading to improved finite sample behavior.

In Fig. 2 we plot the convergence in terms of the MSE of the estimated parameter from ESPRM and ERM, for the LIN-EAR setting and linear policy class (where parameters are low-dimensional and correctly specified). We plot both the MSE convergence, and the average difference in the squared error between the estimates, across the random scenarios. It is clear from these results that ESPRM consistently estimates optimal policy parameters with lower squared error on average compared to ERM across these random simulated scenarios. This provides strong evidence that the methodology indeed provides an improvement in statistical efficiency for solving the smooth surrogate loss problem.

5.2. Jobs Case Study

We next consider an application to a dataset derived from a large scale experiment comparing different programs offered to unemployed individuals in France (Behaghel et al., 2014). We focus our attention to two arms from the experiment: a treatment arm where individuals receive an intensive counseling program run by a public agency and a treatment arm with a similar program run by a private agency. The hypothetical application is learning a personalized allocation to counseling program, with the aim of maximizing the number of individuals who reenter employment within six months, minus costs. (The original study’s focus was not

---

4 All parameter vectors are normalized first given that the policy function is scale-invariant.
Efficient Policy Learning from Surrogate-Loss Classification Reductions

Figure 2. Above we plot the convergence in MSE of the predicted $\hat{\theta}_n$ for each method with a linear policy class, over the random scenarios of the LINEAR class. Below we plot the average difference in the squared error of ESPRM and ERM (positive numbers indicate improvement over ERM). All shaded regions are 95% confidence intervals constructed from bootstrapping.

We then consider 64 replications of the following procedure. Each time, we randomly split the data 40%/60% into train/test. We then introduce some confounding into the training dataset. We consider the following three binary variables: whether individual has 1–5 years experience in the desired job, whether they seek a skilled blue collar job, and whether their statistical risk of long-term unemployment is medium. After studentizing each variable, we segment the data by the tertiles of their sum. In the first tertile, we drop private-program units with probability $1/4$ and public-program units with probability $7/8$. In the third tertile, we drop public-program units with probability $1/4$ and private-program units with probability $7/8$. Given a policy learned on this training data, we evaluate it on the held-out test set using a Horvitz-Thompson estimator.

Of the training data, 20% was set aside for training nuisances, and an additional 20% as validation data for early stopping. We then trained both linear and flexible policies using ERM and ESPRM as in our simulation studies, with the exception that nuisances were fitted using neural networks (of the same architecture as the flexible policy class).

We summarize the mean estimated outcome for the policies from each method in Table 1. We note from these values that on average ESPRM seems to be learning higher value job-assignment policies than ERM. In addition, we conducted paired two-sided $t$-tests to test the hypothesis that the two algorithms lead to different mean policy values on this data, under the randomness in our data splitting and confounding procedures as well as the estimation algorithms. We obtained $p$-values of .0429 for the linear policy class and .0007 for the flexible policy class, clearly highlighting the benefit of our ESPRM method.

<table>
<thead>
<tr>
<th>Policy</th>
<th>ERM</th>
<th>ESPRM</th>
<th>Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear</td>
<td>$-0.96 \pm 4.32$</td>
<td>$4.42 \pm 3.78$</td>
<td>$5.38 \pm 5.06$</td>
</tr>
<tr>
<td>Flexible</td>
<td>$-1.75 \pm 4.64$</td>
<td>$7.68 \pm 3.16$</td>
<td>$9.42 \pm 5.17$</td>
</tr>
</tbody>
</table>

Table 1. Average predicted policy value (multiplied by 1000) for the Jobs case study for ERM versus ESPRM over 64 repetitions. The $\pm$ interval provides the 95% confidence intervals.

6. Conclusion

We considered a common reduction of learning individualized treatment rules from observational data to weighted surrogate risk minimization. We showed that, quite differently from actual classification problems, assuming correct specification in the policy learning case actually suggests more efficient solutions to this reduction. In particular, even if we use efficient policy evaluation, this may not necessarily lead to efficient policy learning. Specifically, under correct specification, the problem becomes a conditional moment problem in a semiparametric model and efficiency here translates to both better MSE in estimating optimal policy parameters and improved regret bounds.

Based on this observation, we proposed an algorithm, ESPRM, for efficiently solving the surrogate loss problem. We showed that our method consistently outperformed the standard method of empirical risk minimization on the surrogate loss, both over a wide variety of synthetic scenarios and in a case study based on a real job training experiment.

personalization.) Our intervention is simply the offer of the counseling program; we therefore ignore the fact that some individuals offered one of the programs did not attend.

In order to make our policies focus on heterogeneous effects, so that a constant-treatment policy wouldn’t be optimal and policy learning would be non-trivial, we set the costs of each arm to be equal to their within-arm average outcome in the original data. That is, the outcome we consider is equal whether one reentered employment within 6 months, minus the average number of individuals who entered employment within 6 months in that arm, so therefore each arm has a mean outcome of zero. The covariates we consider personalizing on are: statistical risk of long-term unemployment, whether individual is seeking full-time employment, whether individual lives in sensitive suburban area, whether individual has a college education, the number of years of experience in the desired job, and the nature of the desired job (e.g., technician, skilled clerical worker, etc.).

We then consider 64 replications of the following procedure. Each time, we randomly split the data 40%/60% into train/test. We then introduce some confounding into the training dataset. We consider the following three binary variables: whether individual has 1–5 years experience in the desired job, whether they seek a skilled blue collar job, and whether their statistical risk of long-term unemployment is medium. After studentizing each variable, we segment the data by the tertiles of their sum. In the first tertile, we drop each unit with probability $7/8$. In the second tertile, we drop each unit with probability $1/4$ and private-program units with probability $7/8$. In the third tertile, we drop public-program units with probability $1/4$ and private-program units with probability $7/8$. Given a policy learned on this training data, we evaluate it on the held-out test set using a Horvitz-Thompson estimator.

Of the training data, 20% was set aside for training nuisances, and an additional 20% as validation data for early stopping. We then trained both linear and flexible policies using ERM and ESPRM as in our simulation studies, with the exception that nuisances were fitted using neural networks (of the same architecture as the flexible policy class).

We summarize the mean estimated outcome for the policies from each method in Table 1. We note from these values that on average ESPRM seems to be learning higher value job-assignment policies than ERM. In addition, we conducted paired two-sided $t$-tests to test the hypothesis that the two algorithms lead to different mean policy values on this data, under the randomness in our data splitting and confounding procedures as well as the estimation algorithms. We obtained $p$-values of .0429 for the linear policy class and .0007 for the flexible policy class, clearly highlighting the benefit of our ESPRM method.

6. Conclusion

We considered a common reduction of learning individualized treatment rules from observational data to weighted surrogate risk minimization. We showed that, quite differently from actual classification problems, assuming correct specification in the policy learning case actually suggests more efficient solutions to this reduction. In particular, even if we use efficient policy evaluation, this may not necessarily lead to efficient policy learning. Specifically, under correct specification, the problem becomes a conditional moment problem in a semiparametric model and efficiency here translates to both better MSE in estimating optimal policy parameters and improved regret bounds.

Based on this observation, we proposed an algorithm, ESPRM, for efficiently solving the surrogate loss problem. We showed that our method consistently outperformed the standard method of empirical risk minimization on the surrogate loss, both over a wide variety of synthetic scenarios and in a case study based on a real job training experiment.
Acknowledgements

This material is based upon work supported by the National Science Foundation under Grant No. 1846210.

References


Rahimi, A. and Recht, B. Weighted sums of random kitchen 
sinks: Replacing minimization with randomization in 
learning. In Advances in neural information processing 

Swaminathan, A. and Joachims, T. Counterfactual risk 
minimization: Learning from logged bandit feedback. 
In International Conference on Machine Learning, pp. 

Van der Vaart, A. W. Asymptotic statistics. Cambridge 

Wager, S. On regression tables for policy learning: comment 
on a paper by jiang, song, li and zeng. Statistica Sinica, 

Zhao, Y., Zeng, D., Rush, A. J., and Kosorok, M. R. Esti-
mating individualized treatment rules using outcome 
weighted learning. Journal of the American Statistical 

Zhou, X., Mayer-Hamblett, N., Khan, U., and Kosorok, 
M. R. Residual weighted learning for estimating individu-
alized treatment rules. Journal of the American Statistical 

Zhou, Z., Athey, S., and Wager, S. Offline multi-action 
policy learning: Generalization and optimization. arXiv 
A. Theory on Classification Calibration

We provide a brief background on what it means technically for a surrogate loss function to be classification-calibrated, and justify that this property holds for the logistic loss function.

First, define $S = \text{sign}(\psi)$, and let $p$ denote the joint probability measure on $S, X$. In addition, let $p^*$ be the corresponding measure re-weighted by $\mathbb{E}[|\psi| | S = s, X = x]$. Formally, if we let $\sigma(S, X)$ be the sigma algebra of $S, X$, then for any $A \in \sigma(S, X)$ this re-weighted measure takes value

$$p^*(A) = \frac{1}{\mathbb{E}[|\psi|]} \int 1\{s, x \in A\} \mathbb{E}[|\psi| | S = s, X = x] dp(s, x),$$

which is well-defined since $\mathbb{E}[|\psi|] < \infty$ by Assumption 2, is clearly non-negative, and has total measure one since $\mathbb{E}[|\psi| | S, X] = \mathbb{E}[|\psi|]$. Now let $J^*(\pi) = (J(\pi)/\mathbb{E}[|\psi|] + 1)/2$, let $L^*(\theta) = L(\theta)/\mathbb{E}[|\psi|]$, and let $\mathbb{E}^*$ denote expectation with respect to $p^*$. Then clearly we have

$$J^*(\pi) = \mathbb{E}^*[1\{S = \pi(X)\}]$$

$$L^*(\theta) = \mathbb{E}^*[l(g_0(X), S)].$$

These re-scaled definitions allow us to analyze regret in terms of theory for unweighted binary classification. Many surrogate loss functions $l$ can be expressed as $l(s, g) = \phi(sg)$, for some convex $\phi$. It is easy to verify that in the case of the logistic regression loss, the definition given in Section 2 is equivalent to $l(g, s) = \phi(sg)$, where $\phi(\alpha) = 2 \log(1 + \exp(\alpha)) - 2 \alpha$.

We first note that given $\mathbb{E}[|\psi|] < \infty$ from Assumption 2, we have that $J$ can be re-scaled by a factor of $\mathbb{E}[|\psi|]$ and expressed as $\mathbb{E}[1\{S = \text{sign}(g_0(X))\}]$. Furthermore, for many classification-based loss functions such as logistic regression loss, the same re-scaling of $L$ allows it to be expressed as $\mathbb{E}[\phi(S_g(X))]$ for some non-negative function $\phi$. In the case of logistic regression loss this function is given by $\phi(\alpha) = 2 \log(1 + \exp(\alpha)) - 2 \alpha$. Now, for any such loss function and corresponding $\phi$, define

$$H(\eta) = \inf_{\alpha \in \mathbb{R}} \eta \phi(\alpha) + (1 - \eta) \phi(-\alpha)$$

$$H^-(\eta) = \inf_{\alpha: -\alpha(2\eta - 1) \leq 0} \eta \phi(\alpha) + (1 - \eta) \phi(-\alpha)$$

$$\tilde{w}(\theta) = H^-(\frac{1 + \theta}{2}) - H\left(\frac{1 + \theta}{2}\right),$$

and let $w$ be the Fenchel-Legendre biconjugate of $\tilde{w}$ (which as discussed in Bartlett et al. (2006), is equal to $\tilde{w}$ if and only if $\tilde{w}$ is convex). Then it follows from Bartlett et al. (2006, Theorem 1) that for any $\theta \in \Theta$ we have

$$w \left( \sup_{\pi \text{ unconstrained}} J^*(\pi) - J^*(g_0) \right) \leq \mathbb{E}^*[l(g_0(X), S)] - \inf_{g \text{ unconstrained}} \mathbb{E}^*[l(g(X), S)].$$

Now, in general, for any non-negative $\phi$ it follows from Bartlett et al. (2006, Lemma 2) that $w$ is non-negative and continuous, and satisfies $w(0) = 0$. Then they define classification-calibrated losses as those where $w(\alpha) > 0$ for all $\alpha \in (0, 1]$, which ensures that surrogate-loss regret converging to zero implies true regret converging to zero.

Finally, Bartlett et al. (2006, Theorem 2) states that for convex, non-negative $\phi$, the corresponding loss is classification-calibrated if and only if it is differentiable at zero and $\phi'(0) < 0$. For logistic regression loss we have $\phi'(\alpha) = 2\sigma(\alpha) - 2$, where $\sigma$ is the logistic function, so therefore $\phi'(0) = -1$ which justifies that this loss is classification-calibrated.

B. Omitted Proofs

Before presenting our omitted proofs, we first establish the following two useful lemmas.

Define

$$\mathcal{G}^* = \arg \min_{g \text{ unconstrained}} \mathbb{E}[|\psi| l(g(X), \text{sign}(\psi))],$$

$$\mathcal{G}^* = \arg \min_{g \in \mathcal{G}} \mathbb{E}[|\psi| l(g(X), \text{sign}(\psi))].$$
Correct specification, Eq. (7), is the assumption that \( G \cap \overline{G}^* \neq \emptyset \).

**Lemma 2.** Suppose \( G \cap \overline{G}^* \neq \emptyset \). Then

\[
G \cap \overline{G}^* = G^*.
\]

**Proof.** For brevity, let \( c(g) = \mathbb{E}[|\psi|l(g(X), \text{sign}(\psi))]. \)

Let any \( g \in G \cap \overline{G}^* \) be given. Now let any \( g' \in G \) be given. Since \( g \in \overline{G}^* \), we have \( c(g) \leq c(g') \). Since \( g' \in G \) was arbitrary, we conclude that \( g \in G^* \).

Now, let any \( g \in G^* \) be given. By assumption, \( \exists g^* \in G \cap \overline{G}^* \). Since \( g^* \in G^* \) and \( g^* \in G \), we obtain that \( c(g) \leq c(g^*) \). Now let any \( g' \) unconstrained be given. Since \( g^* \in \overline{G}^* \), we have \( c(g^*) \leq c(g') \), whence \( c(g) \leq c(g') \). Since \( g' \) unconstrained was arbitrary, we conclude that \( g \in \overline{G}^* \). Since \( g \in G \) by definition of \( G^* \), we conclude that \( g \in G \cap \overline{G}^* \). \( \square \)

**Lemma 3.** Suppose \( \mathbb{E}[|\psi| \mid X] < \infty \) almost surely. Then

\[
\overline{G}^* = \{ g(\cdot) : \mathbb{E}[|\psi|l'(g(X), \text{sign}(\psi)) \mid X] = 0 \text{ almost surely} \}.
\]

**Proof.** Notice that because \( g \) is an unconstrained function of \( X \) it must minimize the conditional expectation. That is,

\[
\overline{G}^* = \{ g : g(x) \in \arg \min_{z \in \mathbb{R}} \mathbb{E}[|\psi|l(z, \text{sign}(\psi)) \mid X = x] \text{ for a.e. } x \}.
\]

Since \( |\psi|l(z, \text{sign}(\psi)) \) is convex in \( z \), so is \( \mathbb{E}[|\psi|l(z, \text{sign}(\psi)) \mid X = x] \). Next, note that by mean value theorem, we have

\[
\frac{\partial}{\partial z} \mathbb{E}[|\psi|l(z, \text{sign}(\psi)) \mid X = x] = \lim_{h \to 0} \mathbb{E} \left[ |\psi|l(z + h, \text{sign}(\psi)) - |\psi|l(z, \text{sign}(\psi)) \middle| X = x \right] = \lim_{h \to 0} \mathbb{E} \left[ |\psi|l'(z(h), \text{sign}(\psi)) \mid X = x \right]
\]

for some \( z(h) \in [z, z+h] \). Since \( |l'(z(h), \text{sign}(\psi))| \leq 4 \) and \( \mathbb{E}[|\psi| \mid X = x] < \infty \), dominated convergence theorem yields

\[
\frac{\partial}{\partial z} \mathbb{E}[|\psi|l(z, \text{sign}(\psi)) \mid X = x] = \mathbb{E} \left[ |\psi| \lim_{h \to 0} l'(z(h), \text{sign}(\psi)) \mid X = x \right] = \mathbb{E} \left[ |\psi|l'(z, \text{sign}(\psi)) \mid X = x \right].
\]

We conclude via first-order conditions for unconstrained optimization over \( z \) that

\[
\overline{G}^* = \{ g : \mathbb{E}[|\psi|l'(g(x), \text{sign}(\psi)) \mid X = x] = 0 \text{ for a.e. } x \},
\]

which is a restatement of the lemma’s result. \( \square \)

We are now prepared to present our omitted proofs.

**Proof of Theorem 1.** Let \( L^*, J^*, \) and \( \mathbb{E}^* \) be defined as in Appendix A, and let any \( \theta^* \in \arg \min_{\theta} L(\theta) \) be given. By our correct-specification assumption, it must be the case that \( L^*(\theta^*) = \inf_{g \text{ unconstrained}} \mathbb{E}^*[l(g(X), \text{sign}(\psi))] \), since any minimizer of \( \mathbb{E}[|\psi|l(g(X), \text{sign}(\psi))] = \mathbb{E}[|\psi|\mathbb{E}^*[l(g(X), \text{sign}(\psi))]] \) is also a minimizer of \( \mathbb{E}^*[l(g(X), \text{sign}(\psi))] \). Therefore the regret bound in Appendix A becomes

\[
w \left( \sup_{\pi \text{ unconstrained}} J^*(\pi) - J^*(\theta^*) \right) \leq L^*(\theta) - L^*(\theta^*),
\]

where \( w \) is a non-negative function that depends on \( l \), and satisfies \( w(0) = 0 \).

Now as discussed in Appendix A, for any classification-calibrated loss we have \( w(\alpha) > 0 \) for \( \alpha \in (0, 1] \), so therefore \( w(\alpha) = 0 \implies \alpha = 0 \). Thus \( J^*(\theta^*) = \sup_{\pi \text{ unconstrained}} J^*(\pi) \), and the final result then follows from the fact that \( J(\pi) = \mathbb{E}[|\psi|](2J^*(\pi) - 1) \) for every policy \( \pi \). \( \square \)
Proof of Theorem 2. First note that Assumption 2 implies that $\mathbb{E}[|\psi| | X = x] < \infty$ almost everywhere, so the conditions of Lemma 3 apply. Now suppose $\theta^* \in \operatorname{arg\,min}_{\theta \in \Theta} L(\theta)$. By Lemma 2, $g_{\theta^*} \in \mathcal{G}^*$. Then, by Lemma 3, $\mathbb{E}[|\psi| | \psi(x), \operatorname{sign}(\psi)) | X = x] = 0$ for a.e. $x$, which is a restatement of $m(X; \theta^*) = 0$ almost surely.

Now suppose $m(X; \theta^*) = 0$ almost surely. Then, by Lemma 3, $g_{\theta^*} \in \mathcal{G}^*$. By definition, $g_{\theta^*} \in \mathcal{G}$. Therefore, by Lemma 2, $g_{\theta^*} \in \mathcal{G}^*$, which is a restatement of $\theta^* \in \operatorname{arg\,min}_{\theta \in \Theta} L(\theta)$.

Proof of Lemma 1. Suppose that II is correctly specified for the surrogate loss. Then given Assumption 2, there exists $\theta^*$ such that, for each $x$ almost everywhere, we have:

$$
\mathbb{E}[|\psi|(2\sigma(g_{\theta^*}(X)) - (\operatorname{sign}(\psi) + 1)) | X = x] = 0
\iff \mathbb{E}[|\psi|(2\sigma(g_{\theta^*}(x)) - (\operatorname{sign}(\psi) + 1)) | X = x] = 0
\iff 2\sigma(g_{\theta^*}(x))\mathbb{E}[|\psi| | X = x] = 2\mathbb{E}[|\psi| \{\psi > 0\} | X = x]
\iff \sigma(g_{\theta^*}(x)) = \frac{\mathbb{E}[|\psi| \{\psi > 0\} | X = x]}{\mathbb{E}[|\psi| | X = x]}.
$$

Proof of Theorem 3. Let $J^*$ and $L^*$ be defined as in Appendix A. Then under our correct-specification assumption, the regret bound presented in Appendix A becomes

$$
w \left( \sup_{\pi \text{ unconstrained}} J^*(\pi) - J^*(\theta^*) \right) \leq L^*(\theta) - \inf_{\theta \in \Theta} L^*(\theta).
$$

Now, by definition we have $L(\theta) = \mathbb{E}[|\psi|] L^*(\theta)$ for every $\theta \in \Theta$, and $J(\pi) = \mathbb{E}[|\psi|](2J^*(\pi) - 1)$ for any policy $\pi$. Therefore we have $\varphi(\text{Regret}_J(\theta)) \leq \text{Regret}_L(\theta)$, where

$$
\varphi(\alpha) = \mathbb{E}[|\psi|] w \left( \frac{\alpha}{2\mathbb{E}[|\psi|]} \right).
$$

As discussed in Appendix A, given Assumption 1 $w$ is continuous and satisfies $w(0) = 0$, and $w(\alpha) > 0$ for $\alpha > 0$. Therefore $\varphi$ clearly must also satisfy these properties.

Finally, in the case of logistic regression loss, it is trivial to show from the definitions in Appendix A that $w(\alpha) = |\alpha|$, so therefore $\varphi(\alpha) = |\alpha|/2$. The result that $\text{Regret}_J(\theta) \leq 2\text{Regret}_L(\theta)$ then immediately follows.

Proof of Theorem 4. Given Assumption 3, from the Taylor expansion from Section 4.4 we have:

$$n\text{Regret}_L(\hat{\theta}_n) = (\sqrt{n}(\hat{\theta}_n - \theta^*))^T H(\theta^*)(\sqrt{n}(\hat{\theta}_n - \theta^*)) + \|\sqrt{n}(\hat{\theta}_n - \theta^*)\|^2 o(1)
$$

Thus assuming that $\hat{\theta}_n$ is regular, we let $W$ be the limiting distribution of $\sqrt{n}(\hat{\theta}_n - \theta^*)$, which by Slutsky’s and the continuous mapping theorem gives us that

$$A R_L(\hat{\theta}_n) = W^T H(\theta^*) W.$$

Now, by Van der Vaart (2000, Theorem 25.20) we have that $W = N(0, V) * M$, where $M$ is given by some arbitrary distribution, $V$ is the covariance matrix of the semi-parametrically efficient estimator, and $*$ denotes convolution. In addition $M = 0$ a.s. $\iff \hat{\theta}_n$ is semi-parametrically efficient. Now let $W^* = N(0, V)$ Then it follows from Van der Vaart (2000, Lemma 8.5) that $\mathbb{E}[\phi(W^T H(\theta^*) W^*)] \geq \mathbb{E}[\phi(W^T H(\theta^*) W^*)]$ for any $W = W^* * M$, since $l(w) = \phi(w^T H(\theta^*) w)$ is a bowl-shaped loss in the sense of Van der Vaart (2000) given that $\phi$ is non-negative and non-decreasing. Thus we can conclude by noting that the efficiency bound is given by $B_\phi = \mathbb{E}[\phi(W^T H(\theta^*) W^*)]$, which is clearly realized for any semi-parametrically efficient $\theta_n$.
C. Extension to Other Surrogate Losses

We provide a brief discussion here about how our theoretical results extend to surrogate losses that satisfy the properties of Assumption 1 except for the assumption of differentiability. A common example of such a loss is hinge loss, which gives a surrogate-loss formulation equivalent to weighted SVM classification.

For such losses, Theorems 1 and 3 both still apply, since the proofs of these do not depend on the assumption of differentiability. Therefore these losses are still Fisher-consistent, and it is still the case that optimal surrogate regret implies an optimal upper bound on true policy learning regret.

Clearly the conditional moment formulation of Theorem 2 cannot apply, since this formulation depends on the derivative of \(l(g, s)\). Instead, define \(D(z', x)\) as the subdifferential of \(E[\psi l(z, \text{sign}(\psi)) \mid X = x] \) in \(z = z'\). Then following an almost identical argument as in the proof of Lemma 3, the class of unconstrained optimizers of \(J\) is given by those where \(0 \in D(g(x), x)\) for almost everywhere \(x\). Then following the same argument as in the proof of Theorem 2, under correct specification optimal policy parameters are those that satisfy \(0 \in D(g\theta, x)\) for almost everywhere \(x\). Alternatively, one may come up with other alternative formulations of the learning problem using for example KKT conditions.

In addition, Theorem 4 clearly cannot apply given that with such losses \(L\) is non-differentiable. In these settings, it is unclear what the implications of the above formulation are in terms of efficiency of estimating \(\theta^*\), and given non-differentiability of \(L\), much of standard efficiency theory breaks down. Therefore we leave the questions of how to achieve optimal regret with such losses, and whether ERP methods are efficient, to future work.

Finally, we note that despite these theoretical gaps, one could still try applying the FiniteGMM algorithm with such losses. In the case of hinge loss, we could use

\[
l'(g, s) = \begin{cases} -1 & \text{if } gs < 1 \\ 0 & \text{if } gs > 1 \end{cases}
\]

and define \(l'\) arbitrarily at the hinge point \(gs = 1\) (which is unlikely to ever be occurred in practice using floating point numbers in the numeric algorithm). It is unclear how such an approach would work both theoretically and practically, and we leave this question to future work.

D. Theory Behind ESPRM Algorithm

In Bennett et al. (2019), they study the instrumental variable problem, which is defined by the conditional moment restriction

\[
E[Y - g(X; \theta^*) \mid Z] = 0,
\]

where \(X\) is the treatment, \(Y\) is the outcome, and \(Z\) is the instrumental variable. As in our setting, this is equivalent to the infinite set of moment restrictions \(E[f(Z)(Y - g(X; \theta^*))] = 0\) for every square integrable function \(f\).

Now, let \(\mathcal{F}_k = \{f_1, \ldots, f_k\}\) be a given finite set of critic functions, and let \(\hat{\theta}_n\) be some prior estimate of \(\theta^*\). Consider the finite set of moment conditions \(E[f_i(Z)(Y - g(X; \theta^*))] = 0\) for \(i \in [k]\) given by \(\mathcal{F}_k\). Then Bennett et al. (2019, Lemma 1) states that the efficiently weighted GMM objective corresponding to these moment conditions and using \(\theta_n\) as the prior parameter estimate is given by

\[
L_n(\theta) = \sup_{f \in \text{span}(\mathcal{F}_k)} \left( \frac{1}{n} \sum_{i=1}^{n} f(Z_i)(Y_i - g(X_i; \theta)) - \frac{1}{4n} \sum_{i=1}^{n} f(Z_i)^2(Y_i - g(X_i; \hat{\theta}_n))^2 \right)
\]

We notice that the proof of this lemma is not specific to the particular structure of this problem, and trivially generalizes to the following lemma, from which the structure of our ESPRM algorithm naturally follows.

**Lemma 4.** Let \((U, W)\) denote a given pair of variables from some fixed joint distribution, and suppose we have \(n\) iid pairs \([(U_1, W_1), (U_2, W_2), \ldots, (U_n, W_n)]\) sampled from this distribution. Consider the moment matching problem given by the \(k\) regular moment conditions \(E[f_i(W)h(U, W; \theta^*)] = 0\), where \(\theta^* \in \Theta \subseteq \mathbb{R}^d\) is the unique solution to this problem (for some integer \(d\)), \(\mathcal{F}_k = \{f_1, \ldots, f_k\}\) are a fixed set of critic functions, and \(h\) is a known function parameterized by \(\theta\). Then the efficiently weighted GMM objective on these moment conditions using \(\hat{\theta}_n\) as the prior estimate is given by

\[
L_n(\theta) = \sup_{f \in \text{span}(\mathcal{F}_k)} \left( \frac{1}{n} \sum_{i=1}^{n} f(W_i)h(U_i, W_i; \theta) - \frac{1}{4n} \sum_{i=1}^{n} f(W_i)^2h(U_i, W_i; \hat{\theta}_n)^2 \right)
\]
We provide here some additional results for our simulation study on Quadratic functions. We include here results for our finite-dimensional GMM method. As mentioned in Section 5, the results for these methods were poor as expected. In particular the results seem to be very unstable, with extremely poor policy learning in a small percentage of cases, leading to extremely negative RMRR values in all cases except with Quadratic scenario and linear policy network. However even in the majority of cases where these estimators don’t have unstable behavior, they seem to perform par with or at best only marginally better than ERM, with the one exception of Quadratic scenario and linear policy network.

In our experiments with finite-dimensional GMM we experimented with two different kinds of choices for the set of critic functions $F$: (1) polynomial expansion of $X$ of degree $d$; and (2) Random Kitchen Sink (RKS; Rahimi & Recht (2009)) expansion of $X$ of length $n$ using the Gaussian kernel with $\sigma = 0.5$. Note that the Random Kitchen Sink expansion is designed such that $\phi_n(x_1)\phi_n(x_2) \approx K(x_1, x_2)$ for some given kernel, with approximation error vanishing as $n \to \infty$. In both cases, the function $f_i$ is given by the $i$'th coordinate of the corresponding feature map. We calculated $\theta_n^{\text{finiteGMM}}$ using 3 stages, with the guess of $\theta_0$ in the first stage chosen at random.

In Fig. 3 we plot the performance of both finite-dimensional GMM and ESPRM in terms of the RMRR metric, plotting both mean and median values across different values of $n$. Although we experimented with multiple choices of polynomial degree / RKS expansion length, we only plot results for degree 3 polynomials (Poly(3)), and length 64 expansions (RBF(64)) for clarity, as we found these gave the least-worst results. We see that in general these methods result in poor performance, and even when they don’t fail they are still inferior to ESPRM.

E.3. Additional Results for Flexible Nuisance Fitting

We provide here some additional results for our simulation study on Quadratic where the nuisances were fit using flexible neural network training (using the same neural network architecture as for the flexible policy class) instead of using a correctly specified model. We show results for the ESPRM and ERM methods in Fig. 4 both with nuisance fit using correctly specified model and using flexible neural network model. We note that results are about the same in both cases: for linear policies we have clearly superior results with our method versus ERM, while for flexible policies in both cases the methods are roughly on par with each other, with slight performance increase in favor of ESPRM for some values of $n$.

E.4. Additional Results on Utility Scale

We provide here some additional results for our simulation study, where we plot the results on a utility scale. Specifically, we plot the same results that were presented in Fig. 1, except that instead of plotting the RMRR metric, we plot the mean difference between the value of the ESPRM policy and the value of the ERM policy. 95% confidence intervals are presented based on the empirical distribution of these differences, from the 64 replications. We display these results in Fig. 5. As with our results on the RMRR scale, we see that our method generally leads to improved policies over the ERM baseline.

Proof. The proof of this lemma is identical to that of Bennett et al. (2019, Lemma 1), replacing $Z$ everywhere with $W$, and replacing $Y - g(X; \theta)$ terms everywhere with corresponding $h(U; W; \theta)$ terms. \qed

E. Additional Experiment Details

E.1. Additional Optimization Details

Solving ESPRM Smooth Game As mentioned in Section 4.3, we solve the smooth game by running alternating first-order optimization using the OAdam algorithm. We tuned this procedure manually by experimenting on a couple of hand selected synthetic scenarios, one linear and one Quadratic, prior to running our main experiments. We found generally good results using a learning rate of 0.001 for linear policy networks, and 0.0002 for flexible policy networks, with the learning rate of the critic $f$ network set to 5 times that of the policy network. Furthermore we found good results using a number of epochs given by the fixed rule of $min(8000000/n, 8000)$, where $n$ is the number of training data points used.

Optimizing Neural Networks for Nuisance Functions and finiteGMM In all cases where we optimized neural networks in these problems we used the LBFGS algorithm. Furthermore we performed some additional first-order optimization using Adam to deal with potential cases of poor convergence, using a learning rate of 0.001, and stopping once performance on a held-out validation set (of same size as training set) failed to improve for 5 consecutive epochs.

E.2. Results for finiteGMM Methods

We include here results for our finiteGMM method. As mentioned in Section 5, the results for these methods were poor as expected. In particular the results seem to be very unstable, with extremely poor policy learning in a small percentage of cases, leading to extremely negative RMRR values in all cases except with Quadratic scenario and linear policy network. However even in the majority of cases where these estimators don’t have unstable behavior, they seem to perform par with or at best only marginally better than ERM, with the one exception of Quadratic scenario and linear policy network.

In our experiments with finiteGMM we experimented with two different kinds of choices for the set of critic functions $F$: (1) polynomial expansion of $X$ of degree $d$; and (2) Random Kitchen Sink (RKS; Rahimi & Recht (2009)) expansion of $X$ of length $n$ using the Gaussian kernel with $\sigma = 0.5$. Note that the Random Kitchen Sink expansion is designed such that $\phi_n(x_1)\phi_n(x_2) \approx K(x_1, x_2)$ for some given kernel, with approximation error vanishing as $n \to \infty$. In both cases, the function $f_i$ is given by the $i$'th coordinate of the corresponding feature map. We calculated $\theta_n^{\text{finiteGMM}}$ using 3 stages, with the guess of $\theta_0$ in the first stage chosen at random.

In Fig. 3 we plot the performance of both finiteGMM and ESPRM in terms of the RMRR metric, plotting both mean and median values across different values of $n$. Although we experimented with multiple choices of polynomial degree / RKS expansion length, we only plot results for degree 3 polynomials (Poly(3)), and length 64 expansions (RBF(64)) for clarity, as we found these gave the least-worst results. We see that in general these methods result in poor performance, and even when they don’t fail they are still inferior to ESPRM.

E.3. Additional Results for Flexible Nuisance Fitting

We provide here some additional results for our simulation study on Quadratic where the nuisances were fit using flexible neural network training (using the same neural network architecture as for the flexible policy class) instead of using a correctly specified model. We show results for the ESPRM and ERM methods in Fig. 4 both with nuisance fit using correctly specified model and using flexible neural network model. We note that results are about the same in both cases: for linear policies we have clearly superior results with our method versus ERM, while for flexible policies in both cases the methods are roughly on par with each other, with slight performance increase in favor of ESPRM for some values of $n$.

E.4. Additional Results on Utility Scale

We provide here some additional results for our simulation study, where we plot the results on a utility scale. Specifically, we plot the same results that were presented in Fig. 1, except that instead of plotting the RMRR metric, we plot the mean difference between the value of the ESPRM policy and the value of the ERM policy. 95% confidence intervals are presented based on the empirical distribution of these differences, from the 64 replications. We display these results in Fig. 5. As with our results on the RMRR scale, we see that our method generally leads to improved policies over the ERM baseline.
Figure 3. Results for ESPRM, ERM, and FINITEGMM methods for all scenarios and policy network types, where nuisances are fit using linear/logistic regression as in main experiments. Shaded regions correspond to 95% confidence intervals, constructed from bootstrapping using the 64 replications.
Figure 4. Results for both ESPRM and ERM methods for QUADRATIC, where in top row results obtained by fitting correctly specified nuisance model, while in bottom row results fit using flexible neural network nuisance model. Shaded regions are 95% confidence intervals, constructed from bootstrapping using the 64 replications.
Figure 5. Results for ESPRM and ERM methods as in Fig. 1, but plotted on a utility scale. We plot the mean difference between the policy values of our method and the ERM baseline, with positive values indicating our method learnt a better policy on average. Shaded regions are 95% confidence intervals based on the 64 replications.