Abstract

Instrumental variable analysis is a powerful tool for estimating causal effects when randomization or full control of confounders is not possible. The application of standard methods such as 2SLS, GMM, and more recent variants are significantly impeded when the causal effects are complex, the instruments are high-dimensional, and/or the treatment is high-dimensional. In this paper, we propose the DeepGMM algorithm to overcome this. Our algorithm is based on a new variational reformulation of GMM with optimal inverse-covariance weighting that allows us to efficiently control very many moment conditions. We further develop practical techniques for optimization and model selection that make it particularly successful in practice. Our algorithm is also computationally tractable and can handle large-scale datasets. Numerical results show our algorithm matches the performance of the best tuned methods in standard settings and continues to work in high-dimensional settings where even recent methods break.

1 Introduction

Unlike standard supervised learning that models correlations, causal inference seeks to predict the effect of counterfactual interventions not seen in the data. For example, when wanting to estimate the effect of adherence to a prescription of \( \beta \)-blockers on the prevention of heart disease, supervised

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*Alphabetical order.
learning may overestimate the true effect because good adherence is also strongly correlated with health consciousness and therefore with good heart health [9]. Figure 1 shows a simple example of this type and demonstrates how a standard neural network (in blue) fails to correctly estimate the true treatment response curve (in orange) in a toy example. The issue is that standard supervised learning assumes that the residual in the response from the prediction of interest is independent of the features.

One approach to account for this is by adjusting for all confounding factors that cause the dependence, such as via matching [26, 27] or regression, potentially using neural networks [19, 28]. However, this requires that we actually observe all confounders so that treatment is as-if random after conditioning on observables. This would mean that in the β-blocker example, we would need to perfectly measure all latent factors that determine both an individual’s adherence decision and their general healthfulness which is often not possible in practice.

Instrumental variables (IVs) provide an alternative approach to causal-effect identification. If we can find a latent experiment in another variable (the instrument) that influences the treatment (i.e., is relevant) and does not directly affect the outcome (i.e., satisfies exclusion), then we can use this to infer causal effects [2]. In the β-blocker example [9], the authors used co-pay cost as an IV. Because they enable analyzing natural experiments under mild assumptions, IVs have been one of the most widely used tools for empirical research in a variety of fields [1]. An important direction of research for IV analysis is to develop methods that can effectively handle complex causal relationships and complex variables like images that necessitate more flexible
models like neural networks [17]. In this paper, we tackle this through a new method called DeepGMM that builds upon the optimally-weighted Generalized Method of Moments (GMM) [13], a widely popular method in econometrics that uses the moment conditions implied by the IV model to efficiently estimate causal parameters. Leveraging a new variational reformulation of the efficient GMM with optimal weights, we develop a flexible framework, DeepGMM, for doing IV estimation with neural networks. In contrast to existing approaches, DeepGMM is suited for high-dimensional treatments $X$ and instruments $Z$, as well as for complex causal and interaction effects. DeepGMM is given by the solution to a smooth game between a prediction function and critic function. We prove that approximate equilibria provide consistent estimates of the true causal parameters. We find these equilibria using optimistic gradient descent algorithms for smooth game play [11], and give practical guidance on how to choose the parameters of our algorithm and do model validation. In our empirical evaluation, we demonstrate that DeepGMM’s performance is on par or superior to a large number of existing approaches in standard benchmarks and continues to work in high-dimensional settings where other methods fail.

2 Setup and Notation

We assume that our data is generated by

$$Y = g_0(X) + \epsilon, \quad (1)$$

where the residual $\epsilon$ has zero mean and finite variance, i.e., $E[\epsilon] = 0$ and $E[\epsilon^2] < \infty$. However, different to standard supervised learning, we allow for the residual $\epsilon$ and $X$ to be correlated, $E[\epsilon | X] \neq 0$, i.e., $X$ can be endogenous, and therefore $g_0(X) \neq E[Y | X]$. We also assume that we have access to an instrument $Z$ satisfying

$$E[\epsilon | Z] = 0. \quad (2)$$

Moreover, $Z$ should be relevant, i.e. $\mathbb{P}(X | Z) \neq \mathbb{P}(X)$. Our goal is to identify the causal response function $g_0(\cdot)$ from a parametrized family of functions $G = \{g(\cdot; \theta) : \theta \in \Theta\}$. Examples are linear functions $g(x; \theta) = \theta^T \phi(x)$, neural networks where $\theta$ represent weights, and non-parametric classes with infinite-dimensional $\theta$. For convenience, let $\theta_0 \in \Theta$ be such that $g_0(\cdot) = g(\cdot; \theta_0)$. Throughout, we measure the performance of an estimated response function $\hat{g}$ by its MSE against the true $g_0$.

Note that if we additionally have some exogenous context variables $L$, the standard way to model this using Eq. (1) is to include them both in $X$ and
in $Z$ as $X = (X', L)$ and $Z = (Z', L)$, where $X'$ is the endogenous variable and $Z'$ is an IV for it. In the $\beta$-blocker example, if we were interested in the heterogeneity of the effect of adherence over demographics, $X$ would include both adherence and demographics whereas $Z$ would include both co-payment and demographics.

2.1 Existing methods for IV estimation

Two-stage methods. One strategy to identifying $g_0$ is based on noting that Eq. (2) implies

$$E[Y | Z] = E[g_0(X) | Z] = \int g_0(x) d\mathbb{P}(X = x | Z).$$

If we let $g(x; \theta) = \theta^T \phi(x)$ this becomes $E[Y | Z] = \theta_0^T \mathbb{E}[\phi(X) | Z]$. The two-stage least squares (2SLS) method \cite{2} first fits $\mathbb{E}[\phi(X) | Z]$ by least-squares regression of $\phi(X)$ on $Z$ (with $Z$ possibly transformed) and then estimates $\hat{\theta}_0^{2SLS}$ as the coefficient in the regression of $Y$ on $\mathbb{E}[\phi(X) | Z]$. This, however, fails when one does not know a sufficient basis $\phi(x)$ for $g(x, \theta_0)$.

\cite{10, 23} propose non-parametric methods for expanding such a basis but such approaches are limited to low-dimensional settings. \cite{17} instead propose DeepIV, which estimates the conditional density $\mathbb{P}(X = x | Z)$ by flexible neural-network-parametrized Gaussian mixtures. This may be limited in settings with high-dimensional $X$ and can suffer from the non-orthogonality of MLE under any misspecification, known as the “forbidden regression” issue \cite[§4.6.1]{2} (see Section 5 for discussion).

Moment methods. The generalized method of moments (GMM) instead leverages the moment conditions satisfied by $\theta_0$. Given functions $f_1, \ldots, f_m$, Eq. (2) implies $E[f_j(Z)c] = 0$, giving us

$$\psi(f_1; \theta_0) = \cdots = \psi(f_m; \theta_0) = 0, \quad \text{where} \quad \psi(f; \theta) = E[f(Z)(Y - g(X; \theta))].$$

A usual assumption when using GMM is that the $m$ moment conditions in Eq. (4) are sufficient to uniquely pin down (identify) $\theta_0$. To estimate $\hat{\theta}_0$, GMM considers these moments’ empirical counterparts, $\psi_n(f; \theta) = \frac{1}{n} \sum_{i=1}^n f(Z_i)(Y_i - g(X_i; \theta))$, and seeks to make all of them small simultaneously, measured by their Euclidean norm $\|v\|^2 = v^T v$:

$$\hat{\theta}_{GMM} = \arg\min_{\theta} \|\psi_n(f_1; \theta), \ldots, \psi_n(f_m; \theta)\|^2.$$
Other vector norms are possible. \cite{22} propose using $\|v\|_\infty$ and solving the optimization with no-regret learning along with an intermittent jitter to moment conditions in a framework they call AGMM (see Section 5 for discussion).

However, when there are many moments (many $f_j$), using any unweighted vector norm can lead to significant inefficiencies, as we may be wasting modeling resources to make less relevant or duplicate moment conditions small. The optimal combination of moment conditions, yielding minimal variance estimates is in fact given by weighting them by their inverse covariance, and it is sufficient to consistently estimate this covariance. In particular, a celebrated result \cite{13} shows that (with finitely-many moments), using the following norm in Eq. (5) will yield minimal asymptotic variance (efficiency) for any consistent estimate $\tilde{\theta}$ of $\theta_0$:

$$
\|v\|^2 = v^T C^{-1}_\tilde{\theta} v, \quad \text{where} \quad [C_{\tilde{\theta}}]_{jk} = \frac{1}{n} \sum_{i=1}^{n} f_j(Z_i)f_k(Z_i)(Y_i - g(X_i; \theta))^2.
$$

Examples of this are the two-step, iterative, and continuously updating GMM estimators \cite{16}. We generically refer to the GMM estimator given in Eq. (5) using the norm given in Eq. (6) as optimally-weighted GMM (OWGMM), or $\hat{\theta}_{\text{OWGMM}}$.

**Failure of (OW)GMM with Many Moment Conditions.** When $g(x; \theta)$ is a flexible model such as a high-capacity neural network, many – possibly infinitely many – moment conditions may be needed to identify $\theta_0$. However, GMM and OWGMM algorithms fail when we use too many moment conditions. On the one hand, one-step GMM (i.e., Eq. (5) with $\|v\| = \|v\|_p$, $p \in [1, \infty]$) is saddled with the inefficiency of trying to impossibly control many equally-weighted moments: at the extreme, if we let $f_1, \ldots$ be all functions of $Z$ with unit square integral, one-step GMM is simply equivalent to the non-causal least-squares regression of $Y$ on $X$. On the other hand, we also cannot hope to learn the optimal weighting: the matrix $C_{\tilde{\theta}}$ in Eq. (6) will necessarily be singular and using its pseudoinverse would mean deleting all but $n$ moment conditions. Therefore, we cannot simply use infinite or even too many moment conditions in GMM or OWGMM.

**3 Methodology**

We next present our approach. We start by motivating it using a new reformulation of OWGMM.
3.1 Reformulating OWGMM

Let us start by reinterpreting OWGMM. Consider the vector space $V$ of real-valued functions $f$ of $Z$ under the usual operations. Note that, for any $\theta, \psi_n(f; \theta)$ is a linear operator on $V$ and

$$C_\theta(f, h) = \frac{1}{n} \sum_{i=1}^{n} f(Z_i)h(Z_i)(Y_i - g(X_i; \theta))^2$$

is a bilinear form on $V$. Now, given any subset $F \subseteq V$, consider the following objective function:

$$\Psi_n(\theta; F, \tilde{\theta}) = \sup_{f \in F} \psi_n(f; \theta) - \frac{1}{4} C_{\tilde{\theta}}(f, f). \quad (7)$$

**Lemma 1.** Let $\|v\|$ be the optimally-weighted norm as in Eq. (6) and let $F = \text{span}(f_1, \ldots, f_m)$. Then

$$\|\psi_n(f_1; \theta), \ldots, \psi_n(f_m; \theta)\|^2 = \Psi_n(\theta; F, \tilde{\theta}).$$

**Corollary 1.** An equivalent formulation of OWGMM is

$$\hat{\theta}_{\text{OWGMM}} \in \arg\min_{\theta \in \Theta} \Psi_n(\theta; F, \tilde{\theta}). \quad (8)$$

In other words, Lemma 1 provides a variational formulation of the objective function of OWGMM and Corollary 1 provides a saddle-point formulation of the OWGMM estimate.

3.2 DeepGMM

In this section, we outline the details of our DeepGMM framework. Given our reformulation above in Eq. (8), our approach is to simply replace the set $F$ with a more flexible set of functions. Namely we let $F = \{f(z; \tau) : \tau \in \mathcal{T}\}$ be the class of all neural networks of a given architecture with varying weights $\tau$ (but not their span). Using a rich class of moment conditions allows us to learn correspondingly a rich $g_0$. We therefore similarly let $G = \{g(x; \theta) : \theta \in \Theta\}$ be the class of all neural networks of a given architecture with varying weights $\theta$.

Given these choices, we let $\hat{\theta}_{\text{DeepGMM}}$ be the minimizer in $\Theta$ of $\Psi_n(\theta; F, \tilde{\theta})$ for any, potentially data-driven, choice $\tilde{\theta}$. We discuss choosing $\tilde{\theta}$ in Section 4. Since this is no longer closed form, we formulate our algorithm in terms of solving a smooth zero-sum game. Formally, our estimator is defined as:

$$\hat{\theta}_{\text{DeepGMM}} \in \arg\min_{\theta \in \Theta} \sup_{\tau \in \mathcal{T}} U_\theta(\theta, \tau) \quad (9)$$
where \[ U_{\tilde{\theta}}(\theta, \tau) = \frac{1}{n} \sum_{i=1}^{n} f(Z_i; \tau)(Y_i - g(X_i; \theta)) \]

\[ - \frac{1}{4n} \sum_{i=1}^{n} f^2(Z_i; \tau)(Y_i - g(X_i; \tilde{\theta}))^2. \]

Since evaluation is linear, for any \( \tilde{\theta} \), the game’s payoff function \( U_{\tilde{\theta}}(\theta, \tau) \) is convex-concave in the functions \( g(\cdot; \theta) \) and \( f(\cdot; \tau) \), although it may not be convex-concave in \( \theta \) and \( \tau \) as is usually the case when we parametrize functions using neural networks. Solving Eq. (9) can be done with any of a variety of smooth game playing algorithms; we discuss our choice in Section 4.

Notably, our approach has very few tuning parameters: only the models \( F \) and \( G \) (i.e., the neural network architectures) and whatever parameters the optimization method uses. In Section 3, we discuss how to select these.

Finally, we highlight that unlike the case for OWGMM as in Lemma 1, our choice of \( F \) is not a linear subspace of \( V \). Indeed, per Lemma 1, replacing \( F \) with a high- or infinite-dimensional linear subspace simply corresponds to GMM with many or infinite moments, which fails as discussed in Section 2.1 (in particular, we would generically have \( \Psi_n(\theta; F, \tilde{\theta}) = \infty \) unhelpfully). Similarly, enumerating many moment conditions as generated by, say, many neural networks \( f \) and plugging these into GMM, whether one-step or optimally weighted, will fail for the same reasons. Instead, our approach is to leverage our variational reformulation in Lemma 1 and replace the class of functions \( F \) with a rich (non-subspace) set in this new formulation, which is distinct from GMM and avoids these issues. In particular, as long as \( F \) has bounded complexity, even if its ambient dimension may be infinite, we can guarantee the consistency of our approach. Since the last layer in a network is a linear combination of the penultimate one, our choice of \( F \) can in some sense be thought of as a union over neural network weights of subspaces spanned by the penultimate layer of nodes.

### 3.3 Consistency

Before discussing practical considerations in implementating DeepGMM, we first turn to the theoretical question of what consistency guarantees we can provide about our method if we were to approximately solve Eq. (9). We phrase our results for generic bounded-complexity functional classes \( F, G \); not necessarily neural networks.

Our main result depends on the following assumptions, which we discuss after stating the result.
**Assumption 1** (Identification). $\theta_0$ is the unique $\theta \in \Theta$ satisfying $\psi(f; \theta) = 0$ for all $f \in \mathcal{F}$.

**Assumption 2** (Bounded complexity). $\mathcal{F}$ and $\mathcal{G}$ have vanishing Rademacher complexities:

$$
\frac{1}{2^n} \sum_{\xi \in \{-1, +1\}^n} \mathbb{E} \sup_{\tau \in \mathcal{T}} \frac{1}{n} \sum_{i=1}^{n} \xi_i f(Z_i; \tau) \to 0,
$$

$$
\frac{1}{2^n} \sum_{\xi \in \{-1, +1\}^n} \mathbb{E} \sup_{\theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \xi_i g(X_i; \theta) \to 0.
$$

**Assumption 3** (Absolutely star shaped). For every $f \in \mathcal{F}$ and $|\lambda| \leq 1$, we have $\lambda f \in \mathcal{F}$.

**Assumption 4** (Continuity). For any $x$, $g(x; \theta), f(x; \tau)$ are continuous in $\theta, \tau$, respectively.

**Assumption 5** (Boundedness). $Y, \sup_{\theta \in \Theta} |g(X; \theta)|, \sup_{\tau \in \mathcal{T}} |f(Z; \tau)|$ are all bounded random variables.

**Theorem 2.** Suppose Assumptions 1 to 5 hold. Let $\hat{\theta}_n$ be any data-dependent sequence with a limit in probability. Let $\hat{\theta}_n, \hat{\tau}_n$ be any approximate equilibrium in the game Eq. (9), i.e.,

$$
\sup_{\tau \in \mathcal{T}} U_{\hat{\theta}_n}(\hat{\theta}_n, \tau) - o_p(1) \leq U_{\hat{\theta}_n}(\hat{\theta}_n, \hat{\tau}_n) \leq \inf_{\theta} U_{\hat{\theta}_n}(\theta, \hat{\tau}_n) + o_p(1).
$$

Then $\hat{\theta}_n \to \theta_0$ in probability.

Theorem 2 proves that approximately solving Eq. (9) (with eventually vanishing approximation error) guarantees the consistency of our method. We next discuss the assumptions we made.

Assumption 1 stipulates that the moment conditions given by $\mathcal{F}$ are sufficient to identify $\theta_0$. Note that, by linearity, the moment conditions given by $\mathcal{F}$ are the same as those given by the subspace $\text{span}(\mathcal{F})$ so we are actually successfully controlling many or infinite moment conditions, perhaps making the assumption defensible. If we do not assume Assumption 1, the arguments in Theorem 2 easily extend to showing instead that $\min_{\theta: \psi(f; \theta) = 0} \forall f \in \mathcal{F} \|\theta - \hat{\theta}_n\| \to 0$, i.e., we approach some identified $\theta$ that satisfies all moment conditions.

Assumption 2 provides that $\mathcal{F}$ and $\mathcal{G}$, although potentially infinite and even of infinite ambient dimension, have limited complexity. Rademacher
complexity is one way to measure function class complexity [4]. Given a bound (envelope) as in Assumption 5, this complexity can also be reduced to other combinatorial complexity measures such as VC- or pseudo-dimension via chaining [25]. [5] studied such combinatorial complexity measures of neural networks.

Assumption 3 is needed to ensure that, for any \( \theta \) with \( \psi(f; \theta) > 0 \) for some \( f \), there also exists an \( f \) such that \( \psi(f; \theta) > \frac{1}{4}C_\theta(f, f) \). It trivially holds for neural networks by considering their last layer. Assumption 4 similarly holds trivially and helps ensure that the moment conditions cannot simultaneously arbitrarily approach zero far from their true zero point at \( \theta_0 \). Assumption 5 is a purely technical assumption that can likely be relaxed to require only nice (sub-Gaussian) tail behavior. Its latter two requirements can nonetheless be guaranteed by either bounding weights (equivalently, using weight decay) or applying a bounded activation at the output. We do not find doing this is necessary in practice.

4 Practical Considerations in Implementing Deep-GMM

Solving the Smooth Zero-Sum Game. In order to solve Eq. (9), we turn to the literature on solving smooth games, which has grown significantly with the recent surge of interest in generative adversarial networks (GANs). In our experiments, we use the OAdam algorithm of [11]. For our game objective, we found this algorithm to be more stable than standard alternating descent steps using SGD or Adam.

Using first-order iterative algorithms for solving Eq. (9) enables us to effectively handle very large datasets. In particular, we implement DeepGMM using PyTorch, which efficiently provides gradients for use in our descent algorithms [24]. As we see in Section 5, this allows us to handle very large datasets with high-dimensional features and instruments where other methods fail.

Choosing \( \tilde{\theta} \). In Eq. (9), we let \( \tilde{\theta} \) be any potentially data-driven choice. Since the hope is that \( \tilde{\theta} \approx \theta_0 \), one possible choice is just the solution \( \tilde{\theta}\text{DeepGMM} \) for another choice of \( \tilde{\theta} \). We can recurse this many times over. In practice, to simulate many such iterations on \( \tilde{\theta} \), we continually update \( \tilde{\theta} \) as the previous \( \theta \) iterate over steps of our game-playing algorithm. Note that \( \tilde{\theta} \) is nonetheless treated as “constant” and does not enter into the gradient of \( \theta \). Given this
approach we can interpret $\hat{\theta}$ in the premise of Theorem 2 as the final $\hat{\theta}$ at convergence, since Theorem 2 allows $\theta$ to be fully data-driven.

**Hyperparameter Optimization.** The only parameters of our algorithm are the neural network architectures for $F$ and $G$ and the optimization algorithm parameters (e.g., learning rate). To tune these parameters, we suggest the following general approach. We form a validation surrogate $\Psi_n$ for our variational objective in Eq. (7) by taking instead averages on a validation data set and by replacing $F$ with the pool of all iterates $f$ encountered in the learning algorithm for all hyperparameter choice. We then choose the parameters that maximize this validation surrogate $\Psi_n$. We discuss this process in more detail in Appendix B.1.

**Early Stopping.** We further suggest to use $\hat{\Psi}_n$ to facilitate early stopping for the learning algorithm. Specifically, we periodically evaluate our iterate $\theta$ using $\hat{\Psi}_n$ and return the best evaluated iterate.

5 Experiments

In this section, we compare DeepGMM against a wide set of baselines for IV estimation. Our implementation of DeepGMM is publicly available at [https://github.com/CausalML/DeepGMM](https://github.com/CausalML/DeepGMM).

We evaluate the various methods on two groups of scenarios: one where $X,Z$ are both low-dimensional and one where $X$, $Z$, or both are high-dimensional images. In the high-dimensional scenarios, we use a convolutional architecture in all methods that employ a neural network to accommodate the images. We evaluate performance of an estimated $\hat{g}$ by MSE against the true $g_0$.

More specifically, we use the following baselines:

1. DirectNN: Predicts $Y$ from $X$ using a neural network with standard least squares loss.
2. Vanilla2SLS: Standard two-stage least squares on raw $X,Z$.
3. Poly2SLS: Both $X$ and $Z$ are expanded via polynomial features, and then 2SLS is done via ridge regressions at each stage. The regularization parameters as well polynomial degrees are picked via cross-validation at each stage.
4. GMM+NN: Here, we combine OWGMM with a neural network $g(x; \theta)$. We solve Eq. (5) over network weights $\theta$ using Adam. We employ optimal weighting, Eq. (6), by iterated GMM [16]. We are not aware of any prior work that uses OWGMM to train neural networks.

5. AGMM [22]: Uses the publicly available implementation\(^1\) of the Adversarial Generalized Method of Moments, which performs no-regret learning on the one-step GMM objective Eq. (5) with norm $\| \cdot \|_\infty$ and an additional jitter step on the moment conditions after each epoch.

6. DeepIV [17]: We use the latest implementation that was released as part of the econML package\(^2\).

Note that GMM+NN relies on being provided moment conditions. When $Z$ is low-dimensional, we follow AGMM [22] and expand $Z$ via RBF kernels around 10 centroids returned from a Gaussian Mixture model applied to the $Z$ data. When $Z$ is high-dimensional, we use the moment conditions given by each of its components.

5.1 Low-dimensional scenarios

In this first group of scenarios, we study the case when both the instrument as well as treatment is low-dimensional. Similar to [22], we generated data via the following process:

$$
Y = g_0(X) + e + \delta \quad \quad X = 0.5 \, Z_1 + 0.5 \, e + \gamma \\
Z \sim \text{Uniform}([-3, 3]^2) \quad \quad e \sim \mathcal{N}(0, 1), \quad \gamma, \delta \sim \mathcal{N}(0, 0.1)
$$

In other words, only the first instrument has an effect on $X$, and $e$ is the confounder breaking independence of $X$ and the residual $Y - g_0(X)$. We keep this data generating process fixed, but vary the true response function $g_0$ between the following cases:

- **sin**: \( g_0(x) = \sin(x) \)
- **step**: \( g_0(x) = 1_{\{x \geq 0\}} \)
- **abs**: \( g_0(x) = |x| \)
- **linear**: \( g_0(x) = x \)

We sample $n = 2000$ points for train, validation, and test sets each. To avoid numerical issues, we standardize the observed $Y$ values by removing the mean and scaling to unit variance. We plot the results in Fig. 2. The left column shows the sampled $Y$ plotted against $X$, with the true $g_0$ in orange.

\(^1\)https://github.com/vsyrgkanis/adversarial_gmm
\(^2\)https://github.com/microsoft/EconML
Figure 2: Low-dimensional scenarios (Section 5.1). Estimated \( \hat{g} \) in blue; true response \( g_0 \) in orange.
other columns show in blue the estimated $\hat{g}$ using various methods. Table 1 shows the corresponding MSE over the test set.

First we note that in each case there is sufficient confounding that the DirectNN regression fails badly and a method that can use the IV information to remove confounding is necessary.

Our next substantive observation is that our method performs competitively across scenarios, attaining the lowest MSE in each (except linear where are beat just slightly and only by methods that use a linear model). At the same time, other methods employing neural networks perform well in some scenarios and less well in others. Therefore we conclude that in the low dimensional setting, our method is able to adapt to the scenario and compete with best tuned methods for the scenario.

Overall, we also found that GMM+NN performed well (but not as well as our method). In some sense GMM+NN is a novel method; we are not aware of previous work using (OW)GMM to train a neural network. Whereas GMM+NN needs to be provided moment conditions, our method can be understood as improving further on this by learning the best moment condition over a large class using optimal weighting. Moreover, we found that GMM+NN outperformed AGMM, which uses the same moment conditions. Aside from the jitter step implemented in the AGMM code, it is equivalent to one-step GMM, Eq. (5), with $\| \cdot \|_\infty$ vector norm in place of the standard $\| \cdot \|_2$ norm. Its worse performance can perhaps be explained by this change and by its lack of optimal weighting.

In the experiments, the other NN-based method, DeepIV, was consistently outperformed by Poly2SLS across scenarios. This can potentially be attributed to the infamous “forbidden regression” issue. DeepIV relies on learning $\mathbb{P}(X = x | Z)$ using a neural-network-parametrized Gaussian mixture model and plugging this into Eq. (3). However, without exactly correct specification, only least-squares is guaranteed to produce orthogonal estimates (with uncorrelated residuals) and ensure Eq. (3) can be used downstream to remove confounding in fitting $\theta$; see [2, §4.6.1]. Other MLE estimates (such as Gaussian mixture), unless perfectly specified, may introduce spurious biases. In addition, the second stage regression in DeepIV is computationally heavy. In the next section, we also discuss its limitations with high-dimensional $X$.

5.2 High-dimensional scenarios

We now move on to scenarios based on the MNIST dataset [20] in order to test our method’s ability to deal with structured, high-dimensional $X$ and $Z$ variables. For this group of scenarios, we use same data generating process as
We sampled 20000 points for the training, validation, and test sets and ran each method 10 times with different random seeds. We report the averaged MSEs in Table 2. We failed to run the AGMM code on any of these
scenarios, as it crashed and returned overflow errors. Similarly, the DeepIV code produced nan outcomes on any scenario with a high-dimensional $X$. Furthermore, because of the size of the examples, we were similarly not able to run Poly2SLS. Instead, we present Vanilla2SLS and Ridge2SLS, where the latter is Poly2SLS with fixed linear degree. Vanilla2SLS failed to produce reasonable numbers for high-dimensional $X$ because the first-stage regression is ill-posed.

Again, we found that our method performed competitively across scenarios, achieving the lowest MSE in each scenario. In the MNIST$_Z$ setting, our method had better MSE than DeepIV. In the MNIST$_X$ and MNIST$_{X,Z}$ scenarios, it handily outperformed all other methods. Even if DeepIV had run on these scenarios, it would be at great disadvantage since it models the conditional distribution over images using a Gaussian mixture. This can perhaps be improved using richer conditional density models like [8, 18], but the forbidden regression issue remains nonetheless. Overall, these results highlights our method’s ability to adept not only to each low-dimensional scenario but also to high-dimensional scenarios, whether the features, instrument, or both are high-dimensional, where other methods break. Aside from our method’s competitive performance, our algorithm was tractable and was able to run on these large-scale examples where other algorithms broke computationally.

6 Conclusions

Other related literature and future work. We believe that our approach can also benefit other applications where moment-based models and GMM is used [6, 14, 15, 15]. Moreover, notice that while DeepGMM is related to GANs [12], the adversarial game that we play is structurally quite different. In some senses, the linear part of our payoff function is similar to that of the Wasserstein GAN [3]; therefore our optimization problem might benefit from a similar approaches to approximating the sup player as employed by WGANs. Another important direction is further investigation into the possible efficiency of DeepGMM or efficient modifications thereof [7].

Conclusions. We presented DeepGMM as a way to deal with IV analysis with high-dimensional variables and complex relationships. The method was based on a new variational reformulation of GMM with optimal weights with the aim of handling many moments and was formulated as the solution to a smooth zero-sum game. Our empirical experiments showed that the method
is able to adapt to a variety of scenarios, competing with the best tuned method in low dimensional settings and performing well in high dimensional settings where even recent methods break.

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A Omitted Proofs

Proof of Lemma 1. First note that since $\|v\|_2^2 = v^T C^{-1} \tilde{\theta} v$, the associated dual norm is $\|v\|_2^* = v^T C \tilde{\theta} v$. Next, define as shorthand $\psi$ as shorthand for $(\psi_n(f_1; \theta), \ldots, \psi_n(f_m; \theta))$. It follows from the definition of the dual norm that $\|\psi\| = \sup_{\|v\| \leq 1} v^T \psi$. Therefore we have:

$$
\|\psi\|_2^2 = \sup_{\|v\| \leq 1} v^T \psi = \sup_{v^T C \tilde{\theta} v \leq \|\psi\|^2} v^T \psi
$$

The Lagrangian of this optimization problem is given by:

$$
L(v, \lambda) = v^T \psi + \lambda(v^T C \tilde{\theta} v - \|\psi\|^2)
$$

Taking the derivative of this with respect to $v$ shows us that when $\lambda < 0$, this quantity is maximized by $v = -\frac{1}{2\lambda} C^{-1} \tilde{\theta} \psi$. In addition we clearly have strong duality for this problem by Slater’s condition whenever $\|\psi\| > 0$ (since in this case $v = 0$ is a feasible interior point). This therefore gives us the following dual formulation for $\|\psi\|^2$:

$$
\|\psi\|^2 = \inf_{\lambda < 0} -\frac{1}{2\lambda} \|\psi\|^2 + \lambda(\frac{1}{4\lambda^2} \|\psi\|^2 - \|\psi\|^2)
= \inf_{\lambda < 0} -\frac{1}{4\lambda} \|\psi\|^2 - \lambda \|\psi\|^2
$$

Taking derivative with respect to $\lambda$ we can see that this is minimized by setting $\lambda = -\frac{1}{2}$. Given this and strong duality, we know it must be the case that $\|\psi\|^2 = \sup_{v} L(v, -\frac{1}{2}) = \sup_{v} v^T \psi - \frac{1}{2} v^T C \tilde{\theta} v + \frac{1}{2} \|\psi\|^2$. Rearranging terms and doing a change of variables $v \leftarrow 2v$ gives us the identity:

$$
\|\psi\|^2 = \sup_{v} v^T \psi + \frac{1}{4} v^T C \tilde{\theta} v
$$

Finally, we can note that any vector $v \in \mathbb{R}^m$ corresponds to some $f \in \text{span}(\mathcal{F})$, such that $f = \sum_i v_i f_i$, and according to this notation we have $v^T \psi = \psi_n(f; \theta)$ and $v^T C \tilde{\theta} v = C_{\tilde{\theta}}(f, f)$. Therefore our required result follows directly from the previous identity. \qed

Proof of Theorem 2. Define $m(\theta, \tau, \tilde{\theta}) = f(Z; \tau)(Y - g(X; \theta)) - \frac{1}{4} f(Z; \tau)^2(Y - g(X; \tilde{\theta}))^2$, $M(\theta) = \sup_{\tau \in \mathcal{T}} \mathbb{E}[m(\theta, \tau, \tilde{\theta})]$, and $M_n(\theta) = \sup_{\tau \in \mathcal{T}} \mathbb{E}_n[m(\theta, \tau, \tilde{\theta}_n)]$, where $\mathbb{E}_n$ refers to the empirical measure (average over the $n$ data points) and $\tilde{\theta}_n \rightarrow_p \tilde{\theta}$. We will proceed by proving the following three conditions, and then proving our results in terms of these conditions:
1. $\sup_{\theta} |M_n(\theta) - M(\theta)| \to_p 0$

2. for every $\delta > 0$ we have $\inf_{\theta \in \Theta} M(\theta) > M(\theta_0)$

3. $M_n(\hat{\theta}_n) \leq M_n(\theta_0) + o_p(1)$

We will proceed by proving these conditions one by one. For the first, we can derive the inequality:

$$\sup_{\theta} |M_n(\theta) - M(\theta)|$$

$$= \sup_{\theta} \left| \sup_{\tau} \mathbb{E}_n[m(\theta, \tau, \hat{\theta}_n)] - \mathbb{E}[m(\theta, \tau, \hat{\theta})] \right|$$

$$\leq \sup_{\theta, \tau} \left| \mathbb{E}_n[m(\theta, \tau, \hat{\theta}_n)] - \mathbb{E}[m(\theta, \tau, \hat{\theta})] \right|$$

$$\leq \sup_{\theta, \tau} \left| \mathbb{E}_n[m(\theta_1, \tau_1, \hat{\theta}_n)] - \mathbb{E}[m(\theta_1, \tau_1, \hat{\theta})] \right| + \sup_{\theta, \tau} \left| \mathbb{E}[m(\theta, \tau, \hat{\theta}_n)] - \mathbb{E}[m(\theta, \tau, \hat{\theta})] \right|$$

Next we will bound these two terms separately, which we will term $B_1$ and $B_2$. For the first term, we can derive the following bound, where $\epsilon_i$ are iid Rademacher random variables, $m_i(\theta, \tau, \hat{\theta}_n) = f(Z_i; \tau)(Y_i - g(X_i; \theta) - \frac{1}{2}f(Z_i; \tau)^2(Y_i - g(X_i; \theta))^2$, and $m'_i(\theta, \tau, \hat{\theta}_n)$ are shadow variables:

$$\mathbb{E}[B_1] = \mathbb{E} \left[ \sup_{\theta_1, \theta_2, \tau} \left| \frac{1}{n} \sum_{i} m_i(\theta_1, \tau, \theta_2) - \mathbb{E}[m'_i(\theta_1, \tau, \theta'_2)] \right| \right]$$

$$\leq \mathbb{E} \left[ \sup_{\theta_1, \theta_2, \tau} \left| \frac{1}{n} \sum_{i} m_i(\theta_1, \tau, \theta_2) - m'_i(\theta_1, \tau, \theta'_2) \right| \right]$$

$$= \mathbb{E} \left[ \sup_{\theta_1, \theta_2, \tau} \left| \frac{1}{n} \sum_{i} \epsilon_i(m_i(\theta_1, \tau, \theta_2) - m'_i(\theta_1, \tau, \theta'_2)) \right| \right]$$

$$\leq 2\mathbb{E} \left[ \sup_{\theta_1, \theta_2, \tau} \left| \frac{1}{n} \sum_{i} \epsilon_i m_i(\theta_1, \tau, \theta_2) \right| \right]$$

$$\leq 2\mathbb{E} \left[ \sup_{\theta, \tau} \left| \frac{1}{n} \sum_{i} \epsilon_i f(Z_i; \tau)(Y_i - g(X_i; \theta)) \right| \right]$$

$$+ \frac{1}{2} \mathbb{E} \left[ \sup_{\theta, \tau} \left| \frac{1}{n} \sum_{i} \epsilon_i f(Z_i; \tau)^2(Y_i - g(X_i; \theta))^2 \right| \right]$$
Given this we can bound the Continuous Mapping Theorem, and Assumption 4 we have the premise of the theorem we have

\[ \mathbb{P}(c) \text{ for some constant } X \]

bounded differences inequality:

Now let Rademacher complexity of class \( L \)

Now given Assumption 5, the functions that map \( B \)

\( = \sup n \mid \epsilon_i f(Z_i; \tau) \leq 1 \)

\( = \sup \sum \mid \epsilon_i g(x; \theta) \leq Y \)

Next, define \( \| \theta, \tau, \theta, \tau \) \)

\( \leq \theta, \tau, \theta \)

\( \leq c, \theta, \tau, \theta, \theta \)

for some constant \( c \). Therefore from McDiarmid’s Inequality we have

\( P(|B_1 - \mathbb{E}[B_1]| \geq \epsilon) \leq 2 \exp(-\frac{2n \epsilon^2}{\epsilon^2}) \). Putting this and the previous result for \( \mathbb{E}[B_1] \) together we get \( B_1 \rightarrow \tau_p 0 \).

Next, define \( \omega_n = \left( (Y - g(X; \tilde{\theta}_n))^2 - (Y - g(X; \tilde{\theta}))^2 \right) \). Recall that from the premise of the theorem we have \( \tilde{\theta}_n \rightarrow \tilde{\theta} \). Then by Slutsky’s Theorem, the Continuous Mapping Theorem, and Assumption 4 we have \( \omega_n = o_p(1) \).

Given this we can bound \( B_2 \) as follows:

\[ B_2 = \sup \mathbb{E}[m(\theta, \tau; \tilde{\theta})] - \mathbb{E}[m(\hat{\theta}, \hat{\tau})] \]

\[ = \frac{1}{4} \sup \mathbb{E}[f(Z; \tau)^2(Y - g(X; \hat{\theta}))^2] - \mathbb{E}[f(Z; \tau)^2(Y - g(X; \tilde{\theta}))^2] \]

\( \leq \frac{1}{4} \sup \mathbb{E}[f(Z; \tau)^2(Y - g(X; \tilde{\theta}))^2] - \mathbb{E}[f(Z; \tau)^2(Y - g(X; \hat{\theta}))^2] + \frac{1}{4} \sup \mathbb{E}[f(Z; \tau)^2 \omega_n] \]

\( = \frac{1}{4} \sup \mathbb{E}[f(Z; \tau)^2 \omega_n] \)
Now we know from Assumption 5 that \( f(Z; \tau) \) is uniformly bounded, so it follows that \( B_2 \leq b \mathbb{E}[|\omega_n|] \) for some constant \( b \). Next we can note, again based on our boundedness assumption, that \( \omega_n \) is uniformly bounded. Therefore it follows from the Lebesgue Dominated Convergence Theorem that \( \mathbb{E}[|\omega_n|] \rightarrow 0 \). Thus we know that both \( B_1 \) and \( B_2 \) converge, so we have proven the first of the three conditions, that \( \sup_0 |M_n(\theta) - M(\theta)| \) converges in probability to zero.

For the second condition we will first prove that \( M(\theta_0) \) is the unique minimizer of \( M(\theta) \). Clearly by Assumptions 1 and 3 we have that \( \sup_\tau \mathbb{E}[f(Z; \tau)(Y - g(X; \theta))] \) is uniformly bounded. Furthermore, for any other \( \theta' \neq \theta_0 \), let \( f' \) be some function in \( \mathcal{F} \) such that \( \mathbb{E}[f(Z)(Y - g(X; \theta'))] > 0 \). If we have \( \mathbb{E}[f'(Z)^2(Y - g(X; \theta))^2] = 0 \) then it follows immediately that \( M(\theta') > 0 \). Otherwise, consider the function \( \lambda f' \) for arbitrary \( 0 < \lambda < 1 \). Since by Assumption 3 this function is also contained in \( \mathcal{F} \), it follows that:

\[
M(\theta') = \sup_{f \in \mathcal{F}} \mathbb{E}[f(Z)(Y - g(X; \theta'))] - \frac{1}{4} f(Z)^2(Y - g(X; \hat{\theta}))^2 \\
\geq \lambda \mathbb{E}[f'(Z)(Y - g(X; \theta'))] - \frac{\lambda^2}{4} \mathbb{E}[f'(Z)^2(Y - g(X; \hat{\theta})^2]
\]

This expression is a quadratic in \( \lambda \) that is clearly positive when \( \lambda \) is sufficiently small, so therefore it still follows that \( M(\theta') > 0 \).

Given this, we will prove the second condition by contradiction. If this were false, then for some \( \delta > 0 \) we would have that \( \inf_{\theta \in B(\theta_0, \delta)} M(\theta) = M(\theta_0) \), where \( B(\theta_0, \delta) = \{\theta \mid d(\theta, \theta_0) \geq \delta\} \). This is because from Assumption 1 we know \( \theta_0 \) is the unique minimizer of \( M(\theta) \). Given this there must exist some sequence \( (\theta_1, \theta_2, \ldots) \) in \( B(\theta_0, \delta) \) satisfying \( M(\theta_n) \rightarrow M(\theta_0) \). Now by construction \( B(\theta_0, \delta) \) is closed, and the corresponding limit parameters \( \theta^* = \lim_{n \rightarrow \infty} \theta_n \in B(\theta_0, \delta) \) must satisfy \( M(\theta^*) = M(\theta_0) \), since given Assumption 4 \( M(\theta) \) is clearly a continuous function of \( \theta \) so we can swap function application and limit. However \( d(\theta^*, \theta_0) \geq \delta > 0 \), so \( \theta^* \neq \theta_0 \). This contradicts the fact that \( \theta_0 \) is the unique minimizer of \( M(\theta) \), so we have proven the second condition.

Finally, for the third condition we will use the fact that by assumption
\( \hat{\theta}_n \) satisfies the approximate equilibrium condition:

\[
\sup_{\tau \in \mathcal{T}} \mathbb{E}_n m(\hat{\theta}_n, \tau, \hat{\theta}_n) - o_p(1) \leq \mathbb{E}_n m(\hat{\theta}_n, \hat{\tau}_n, \hat{\theta}_n) \leq \inf_{\theta} \mathbb{E}_n m(\theta, \hat{\tau}_n, \hat{\theta}_n) + o_p(1)
\]

Now by definition \( M_n(\hat{\theta}_n) = \sup_{\tau \in \mathcal{T}} \mathbb{E}_n m(\hat{\theta}_n, \tau, \hat{\theta}_n) \). Therefore,

\[
\inf_{\theta} \mathbb{E}_n m(\theta, \hat{\tau}_n, \hat{\theta}_n) \leq \inf_{\theta} \sup_{\tau} \mathbb{E}_n m(\theta, \tau, \hat{\theta}_n) = \inf_{\theta} M_n(\theta) \leq M_n(\theta_0).
\]

Thus we have

\[
M_n(\hat{\theta}_n) - o_p(1) \leq \mathbb{E}_n m(\hat{\theta}_n, \hat{\tau}_n, \hat{\theta}_n) \leq M_n(\theta_0) + o_p(1).
\]

At this point we have proven all three conditions stated at the start of the proof. For the final part we can first note that from the first and third conditions it easily follows that \( M_n(\hat{\theta}_n) \leq M(\theta_0) + o_p(1) \), since \( |M_n(\theta_0) - M(\theta_0)| \to 0 \). Therefore we have:

\[
M(\hat{\theta}_n) - M(\theta_0) \leq M(\hat{\theta}_n) - M_n(\hat{\theta}_n) + o_p(1)
\]

\[
\leq \sup_{\theta} \left| M(\hat{\theta}) - M_n(\hat{\theta}) \right| + o_p(1)
\]

\[
\leq o_p(1)
\]

Next, define \( \eta(\delta) = \inf_{d(\theta_n, \theta_0) \geq \delta} M(\theta) - M(\theta_0) \). Now by definition of \( \eta \) we know that whenever \( d(\theta_n, \theta) \geq \delta \) we have \( M(\hat{\theta}_n) - M(\theta_0) \geq \eta(\delta) \). Therefore \( \mathbb{P}[d(\theta_n, \theta) \geq \delta] \leq \mathbb{P}[M(\hat{\theta}_n) - M(\theta_0) \geq \eta(\delta)] \). Now since for every \( \delta > 0 \) we have \( \eta(\delta) > 0 \) from the second condition, and we know \( M(\theta_0) - M(\theta_0) = o_p(1) \), we have that for every \( \delta > 0 \) the RHS probability converges to zero. Thus \( d(\theta_n, \theta_0) = o_p(1) \), so we can conclude that \( \hat{\theta}_n \to_p \theta_0 \)

\[\square\]

B Additional Methodology Details

B.1 Hyperparameter Optimization Procedure

We provide more details here about the hyperparameter optimization procedure described in Section 4. Let \( m \) be the total number of hyperparameter choices under consider. Then for each candidate set of hyperparameters \( \gamma_i \in \{\gamma_1, \ldots, \gamma_m\} \) we run our learning algorithm for a fixed number of epochs using \( \gamma_i \), training it on our train partition. Every \( k_{eval} \) epochs we save the current parameters \( \hat{\tau} \) and \( \hat{\theta} \) at that epoch. This gives, for each hyperparameter choice \( \gamma_i \), a finite set of \( f \) functions \( \hat{F}_i \), and a finite set of \( \theta \) values \( \hat{\Theta}_i \).
Now, define the set of functions $\tilde{F} = \cup_{i=1}^{m} \tilde{F}_i$. We define our approximation of our variational objective as

$$\hat{\Psi}_n(\theta) = \Psi_n(\theta; \tilde{F}, \theta),$$

where $\Psi_n$ is as defined in Eq. (7). Note that this means for every $\theta$ we wish to evaluate we choose to approximate $\tilde{\theta}$ using that value of $\theta$.

Given this, we finally choose the set of hyperparameters $\gamma_i$ whose corresponding trajectory of parameter values $\hat{\Theta}_i$ minimizes the objective function $\min_{\theta \in \hat{\Theta}_i} \hat{\Psi}_n(\theta)$, calculated on the validation data. Note that this objective function is meant to approximate the value of the variational objective we would have obtained if we performed learning with that set of hyperparameters using early stopping.

Note that in practice, since we only ever calculate $\hat{\Psi}_n$ on the validation data, and we only ever use $\Theta_i$ in optimizing the above objective function on the validation data, instead of saving the actual parameter values $\tilde{\tau}$ and $\tilde{\theta}$ we can instead save vectors $f(Z_{val}, \tau)$ and $g(X_{val}, \theta)$, where $Z_{val}$ and $X_{val}$ are the vectors of $Z$ and $X$ values respectively in our validation data. This makes our methodology tractable when working with very complex deep neural networks.