TARGETED VAE: STRUCTURED INFERENCE AND TAR-GETED LEARNING FOR CAUSAL PARAMETER ESTIMA-TION

Anonymous authors

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ABSTRACT

Undertaking causal inference with observational data is extremely useful across a wide range of domains including the development of medical treatments, advertisements and marketing, and policy making. There are two main challenges associated with undertaking causal inference using observational data: treatment assignment heterogeneity (i.e., differences between the treated and untreated groups), and an absence of counterfactual data (i.e. not knowing what would have happened if an individual who did get treatment, were instead to have not been treated). We address these two challenges by combining structured inference and targeted learning. To our knowledge, Targeted Variational AutoEncoder (TVAE) is the first method to incorporate targeted learning into deep latent variable models. Results demonstrate competitive and state of the art performance.

1 INTRODUCTION

The estimation of the causal effects of interventions or treatments on outcomes is of the upmost importance across a range of decision making processes and scientific endeavours, such as policy making (Kreif & DiazOrdaz, 2019), advertisement (Bottou et al., 2013), the development of medical treatments (Petersen et al., 2017), the evaluation of evidence within legal frameworks (Pearl, 2009; Siegerink et al., 2016) and social science (Vowels, 2020; Hernan, 2018; Grosz et al., 2020). Despite the common preference for Randomized Controlled Trial (RCT) data over observational data, this preference is not always justified. Besides the lower cost and fewer ethical concerns, observational data may provide a number of statistical advantages including greater statistical power and increased generalizability (Deaton & Cartwright, 2018). However, there are two main challenges when dealing with observational data. Firstly, the group that receives treatment is usually not equivalent to the group that does not (treatment assignment heterogeneity), resulting in selection bias and confounding due to associated covariates. For example, young people may prefer surgery, older people may prefer medication. Secondly, we are unable to directly estimate the causal effect of treatment, because only the factual outcome for a given treatment assignment is available. In other words, we do not have the counterfactual associated with the outcome for a different treatment assignment to that which was given. Treatment effect inference with observational data is concerned with finding ways to estimate the causal effect by considering the expected differences between factual and counterfactual outcomes.

We seek to address the two challenges by proposing a method that incorporates targeted learning techniques into a disentangled variational latent model, trained according to the approximate maximum likelihood paradigm. Doing so enables us to infer hidden confounders from proxy variables in the dataset, and to estimate the expected treatment effects, as well as individual-level treatment effects. Estimating the latter is especially important for treatments that interact with patient attributes, whilst also being crucial for enabling individualized treatment assignment. Thus, we propose the Targeted Variational AutoEncoder (TVAE), undertake an ablation study, and compare our method's performance against current alternatives on two benchmark datasets.



Figure 1: Directed Acyclic Graphs (DAGs) for (a) the problem of estimating the effect of treatment t on outcome y with confounders x. DAG (b) reflects an RCT. DAG (c) illustrates TVAE and is an extension of the DAG by Zhang et al. (2020), where the structure is *a priori* assumed to factorize into into risk z_y , instrumental z_t , and confounding factors z_c . We extend their model with z_o to account for the scenario whereby not all covariates will be related to treatment and/or outcome.

2 BACKGROUND

Problem Formulation: A characterization of the problem of causal inference with no unobserved confounders is depicted in the Directed Acyclic Graphs (DAGs) shown in Figs. 1(a) and 1(b). Fig. 1(a) is characteristic of observational data, where the assignment of treatment is related to the covariates. Fig. 1(b) is characteristic of the ideal RCT, where the treatment is unrelated to the covariates. Here, $\mathbf{x}_i \sim p(\mathbf{x}) \in \mathbb{R}^m$ represents the *m*-dimensional, pre-treatment covariates for individual *i* assigned factual treatment $t_i \sim p(\mathbf{t}|\mathbf{x})$ resulting in factual outcome $y_i^t \sim p(\mathbf{y}|\mathbf{x}, \mathbf{t})$. Together, these constitute dataset $\mathcal{D} = \{[y_i, t_i, \mathbf{x}_i]\}_{i=1}^N$ where *N* is the sample size.

The conditional average treatment effect for an individual with covariates \mathbf{x}_i may be estimated as $\hat{\tau}_i(\mathbf{x}_i) = \mathbb{E}[y_i|\mathbf{x}_i, do(\mathbf{t} = 1) - y_i|\mathbf{x}_i, do(\mathbf{t} = 0)]$, where the expectation accounts for the nondeterminism of the outcome (Jesson et al., 2020). Alternatively, by comparing the post-intervention distributions when we intervene on treatment \mathbf{t} , the Average Treatment Effect (ATE) is $\hat{\tau}(\mathbf{x}) = \mathbb{E}_{\mathbf{x}}[\mathbb{E}[\mathbf{y}|\mathbf{x}, do(\mathbf{t} = 1)] - \mathbb{E}[\mathbf{y}|\mathbf{x}, do(\mathbf{t} = 0)]]$. Here, $do(\mathbf{t})$ indicates the intervention on \mathbf{t} , setting all instances to a static value, dynamic value, or distribution and therefore removing any dependencies it originally had (Pearl, 2009; van der Laan & Rose, 2018; 2011). This scenario corresponds with the DAG in Fig. 1(b), where treatment \mathbf{t} is no longer a function of the covariates \mathbf{x} .

Using an estimator for the conditional mean $Q(\mathbf{t}, \mathbf{x}) = \mathbb{E}(\mathbf{y}|\mathbf{t}, \mathbf{x})$, we can calculate the Average Treatment Effect (ATE) and the empirical error for estimation of the ATE (eATE).¹ In order to estimate eATE we assume access to the ground truth treatment effect τ , which is only possible with synthetic or semi-synthetic datasets. The Conditional Average Treatment Effect (CATE) may also be calculated and the Precision in Estimating Heterogeneous Effect (PEHE) is one way to evaluate a model's efficacy in estimating this quantity. See the appendix for the complete definitions of these terms.

The Naive Approach: The DAG in Fig. 1(a) highlights the problem with taking a naive approach to modeling the joint distribution $p(\mathbf{y}, \mathbf{t}, \mathbf{x})$. The structural relationship $\mathbf{t} \leftarrow \mathbf{x} \rightarrow \mathbf{y}$ indicates both that the assignment of treatment \mathbf{t} is dependent on the covariates \mathbf{x} , and that a backdoor path exists through \mathbf{x} to \mathbf{y} . In addition to our previous assumptions, if we also assume linearity, adjusting for this backdoor path is a simple matter of adjusting for \mathbf{x} by including it in a logistic regression. The naive method is an example of the uppermost methods depicted in Fig. 2, and leads to the largest bias. The problem with the approach is (a) that the graph is likely misspecified such that the true relationships between covariates as well as the relationships between covariates and the outcome may be more complex. There is also problem (b), that linearity is not sufficient to 'let the data speak' (van der Laan & Rose, 2011) or to avoid biased parameter estimates. Using powerful nonparametric models (e.g., neural networks) may solve the limitations associated with linearity and interactions to yield a consistent estimator for $p(\mathbf{y}|\mathbf{X})$, and such a model is an example of the middlemost methods depicted in Fig. 2. However, this estimator is not targeted to the estimation of the causal effect parameter $\boldsymbol{\tau}$, only predicting the outcome, and we require a means to reduce residual bias.

Targeted Learning: Targeted Maximum Likelihood Estimation (TMLE) (Schuler & Rose, 2016; van der Laan & Rose, 2011; 2018; van der Laan & Starmans, 2014) falls under the lowermost meth-

¹For a binary outcome variable $\mathbf{y} \in \{0, 1\}$, $\mathbb{E}(\mathbf{y}|\mathbf{t}, \mathbf{x})$ is the same as the conditional probability distribution $p(\mathbf{y}|\mathbf{t}, \mathbf{x})$.



Figure 2: Different methods for estimating the causal parameter τ yield different levels of bias. Adapted from (van der Laan & Rose, 2011).

ods depicted in Fig. 2 and follows an approach involving three main steps: (1) estimation of the conditional mean $\mathbb{E}(\mathbf{y}|\mathbf{t},\mathbf{x})$ with estimator $Q^0(\mathbf{t},\mathbf{x})$, (2) estimation of the propensity scores with estimator $g(\mathbf{t}|\mathbf{x})$, and (3) updating the conditional mean estimator Q^0 to get Q^* using the propensity scores to attain an estimate for the causal parameter τ . The propensity score for individual i is defined as the conditional probability of being assigned treatment $g(t_i, \mathbf{x}_i) = p(\mathbf{t} = t_i | \mathbf{x} = \mathbf{x}_i)$, $\in [0, 1]$ (Rosenbaum & Rubin, 1983). The scores can be used to compensate for the relationship between the covariates and the treatment assignment using Inverse Probability of Treatment Weights (IPTWs), reweighting each sample according to its propensity score. Step (3) is undertaken using 'clever covariates' which are similar to the IPTWs. They form an additional covariate variable $H(1, \mathbf{x}_i) = g(1|\mathbf{x}_i)^{-1}$ for individual i assigned treatment, and $H(0, \mathbf{x}_i) = -g(1|\mathbf{x}_i)^{-1}$ for individual i not assigned treatment. Note that when we condition on a single numeric value we imply an intervention (e.g. $g(1|\mathbf{x}_i) \equiv g(do(\mathbf{t} = 1)|\mathbf{x}_i)$). A logistic regression is then undertaken as $\mathbf{y} = \sigma^{-1}[Q^0(\mathbf{t}, \mathbf{x})] + \epsilon H(\mathbf{t}, \mathbf{x})$ where σ^{-1} is the logit/inverse sigmoid function, $Q^0(\mathbf{t}, \mathbf{x})$ is set to be a constant, suppressed offset and ϵ represents a fluctuation parameter. This yields an updated estimator:

$$Q^{1}(do(\mathbf{t}=t),\mathbf{x}) = \sigma \left[\sigma^{-1}[Q^{0}(do(\mathbf{t}=t),\mathbf{x})] + \epsilon H(do(\mathbf{t}=t),\mathbf{x})\right]$$
(1)

 Q^1 can then be plugged into the estimator for $\hat{\tau}(Q^1; \mathbf{x})$. The update procedure can be undertaken multiple times, although following the procedure above should yield the optimal solution after one update. It can be seen from Eq. 1 that, following the optimal solution, $\epsilon = 0$, which implies that the clever covariates no longer exert influence on the estimator. When $\epsilon = 0$, the estimator also satisfies what is known as the efficient Influence Curve (IC), or canonical gradient equation (Hampel, 1974; van der Laan & Rose, 2011):

$$\sum_{i=1}^{N} IC^{*}(y_{i}, t_{i}, \mathbf{x}_{i}) = 0 = \sum_{i=1}^{N} \left[H(t_{i}, \mathbf{x}_{i})(y_{i} - Q(t_{i}, \mathbf{x}_{i})) + Q(1, \mathbf{x}_{i}) - Q(0, \mathbf{x}_{i}) - \boldsymbol{\tau}(Q; \mathbf{x}) \right]$$
(2)

where $IC(y_i, t_i, \mathbf{x}_i)$ represents the IC, and $IC^*(y_i, t_i, \mathbf{x}_i)$ represents the efficient IC for consistent Q and g. It can be seen from Eq. 2 that at convergence, the estimator and its estimand are equal, such that over the whole dataset, all terms 'cancel' resulting in the mean IC = 0. As such, the logistic regression in Eq. 1 represents a solution to the IC via a parametric submodel. This notable result yields standard errors and confidence intervals for p-values. The TMLE method provides a doubly robust, asymptotically efficient estimate of the causal or 'target' parameter, and these theoretical guarantees make it attractive for inclusion into neural networks for causal effect estimation.

3 Methodology

In this section we present the Targeted Variational AutoEncoder (TVAE), a deep generative latent variable model that enables estimation of the average and conditional average treatment effects (ATE and CATE resp.) via the use of amortized variational inference techniques and Targeted Maximum Likelihood Estimation (TMLE). For a review on the relevant VAE theory, see the appendix. A top-level diagram for TVAE is shown in Fig. 3 and follows the structure implied by the DAG in Fig. 1(c). A more detailed architectural block-diagram is also presented in the appendix.²

Assumptions: As is common (Yao et al., 2020; Guo et al., 2020; Rubin, 2005; Imbens & Rubin, 2015) when undertaking causal inference with observational data, we make three assumptions: (1) Stable Unit Treatment Value Assumption (SUTVA): the potential outcomes for each individual or data unit are independent of the treatments assigned to all other individuals, such that there are no

²Source code is also made available in supplementary material.



Figure 3: The block-diagram for Targeted VAE. Dashed boxes indicate the variationally inferred latent variables \mathbf{z}_o , \mathbf{z}_c , \mathbf{z}_t , and \mathbf{z}_y . Arrows indicate functions, and colors distinguish treatment (green), outcome (blue), covariates (grey), and confounder (purple) related entities.

interactions between individuals. (2) Positivity: the assignment of treatment probabilities are all non-zero and non-deterministic $p(\mathbf{t} = t_i | \mathbf{x} = \mathbf{x}_i) > 0$, $\forall \mathbf{t}$ and \mathbf{x} . (3) Ignorability: all confounders are observed such that the likelihood of treatment for two individuals with the same covariates is equal, and the potential outcomes for two individuals with the same covariates are also equal $s.t. (\mathbf{y}^{t=1}, \mathbf{y}^{t=0}) \perp \mathbf{t} | \mathbf{x} \text{ and } \mathbf{t} \perp (\mathbf{y}^{t=1}, \mathbf{y}^{t=0}) | \mathbf{x}.^3$

TVAE: We seek to factorize and disentangle the latent distribution into $\{\mathbf{z}_t, \mathbf{z}_y, \mathbf{z}_c, \mathbf{z}_o\}$, which represent the instrumental factors on \mathbf{t} , the risk factors on \mathbf{y} , the confounders on both \mathbf{t} and \mathbf{y} , and factors solely related to \mathbf{x} , respectively. Without inductive bias, consistently disentangling the latent variables into these factors would be impossible (Locatello et al., 2019). In TVAE this inductive bias is incorporated in a number of ways: firstly, by incorporating supervision and predicting \mathbf{t} and \mathbf{y} from \mathbf{z}_t and \mathbf{z}_y respectively; secondly, by structuring the dependencies according to the desired factorization; and finally, by employing diagonal-covariance priors to encourage disentanglement between latent variables. A further challenge exists relating to causal identifiability and ignorability. In general, it is impossible to isolate the effect of $\mathbf{t} \rightarrow \mathbf{y}$ due to unobserved confounding (D'Amour, 2019). However, the use of deep latent variable techniques enables us to attempt to infer these hidden confounders from what are known as *noisy proxy* variables present in the dataset (see e.g., Montgomery et al. 2000). The assumption of ignorability then shifts from 'all confounders are observed', to 'all unobserved confounders have been inferred from proxies'.

Inference:

$$\begin{split} q(\mathbf{z}_{t}|\mathbf{x}) &= \prod_{d=1}^{D_{z_{t}}} \mathcal{N}(\mu_{d} = f_{1d}(\mathbf{x}), \sigma_{d}^{2} = f_{2d}(\mathbf{x})); \ q(\mathbf{z}_{y}|\mathbf{x}) = \prod_{d=1}^{D_{z_{y}}} \mathcal{N}(\mu_{d} = f_{3d}(\mathbf{x}), \sigma_{d}^{2} = f_{4d}(\mathbf{x})) \\ q(\mathbf{z}_{c}|\mathbf{x}) &= \prod_{d=1}^{D_{z_{c}}} \mathcal{N}(\mu_{d} = f_{5d}(\mathbf{x}), \sigma_{d}^{2} = f_{6d}(\mathbf{x})); q(\mathbf{z}_{o}|\mathbf{x}) = \prod_{d=1}^{D_{z_{o}}} \mathcal{N}(\mu_{d} = f_{7d}(\mathbf{x}), \sigma_{d}^{2} = f_{8d}(\mathbf{x})) \\ p(\hat{\mathbf{t}}|\mathbf{z}_{t}, \mathbf{z}_{c}) &= \operatorname{Bern}(f_{9}(\mathbf{z}_{t}, \mathbf{z}_{c})) = \operatorname{Bern}(g_{q}(\mathbf{z}_{t}, \mathbf{z}_{c})) \\ p(\hat{\mathbf{y}}|\mathbf{z}_{y}, \mathbf{z}_{c}, \mathbf{t}) &= \operatorname{Bern}(\hat{\mathbf{t}} \cdot f_{10}(\mathbf{z}_{y}, \mathbf{z}_{c}) + (1 - \hat{\mathbf{t}}) \cdot f_{11}(\mathbf{z}_{y}, \mathbf{z}_{c})) = \operatorname{Bern}(Q_{q}(.)) \\ \mathbf{Generation:} \end{split}$$

$$p(\mathbf{z}_{\{o,t,c,y\}}) = \prod_{d}^{D_{\{\mathbf{z}_{o,t,c,y}\}}} \mathcal{N}(z_{\{o,t,c,y\}d}|0,1); \quad p(\hat{\mathbf{t}}|\mathbf{z}_{t},\mathbf{z}_{c}) = \operatorname{Bern}(h_{1}(\mathbf{z}_{t},\mathbf{z}_{c})) = \operatorname{Bern}(g_{p}(.))$$

$$p(\hat{\mathbf{y}}|\mathbf{z}_{y},\mathbf{z}_{c},\hat{\mathbf{t}}) = \operatorname{Bern}(\hat{\mathbf{t}} \cdot h_{2}(\mathbf{z}_{y},\mathbf{z}_{c}) + (1-\hat{\mathbf{t}}) \cdot h_{3}(\mathbf{z}_{y},\mathbf{z}_{c})) = \operatorname{Bern}(Q_{p}(.))$$

$$p(\hat{\mathbf{x}}_{bin}|\mathbf{z}_{c},\mathbf{z}_{o},\mathbf{z}_{t},\mathbf{z}_{y}) = \operatorname{Bern}(h_{6}(\mathbf{z}_{c},\mathbf{z}_{o},\mathbf{z}_{t},\mathbf{z}_{y}))$$

$$p(\hat{\mathbf{x}}_{cont}|\mathbf{z}_{c},\mathbf{z}_{o},\mathbf{z}_{t},\mathbf{z}_{y}) = \prod_{d=1}^{D_{x_{cont}}} \mathcal{N}(x_{cont,d}|\mu_{d} = h_{4}(\mathbf{z}_{c},\mathbf{z}_{o},\mathbf{z}_{t},\mathbf{z}_{y}), \sigma_{d}^{2} = h_{5}(\mathbf{z}_{c},\mathbf{z}_{o},\mathbf{z}_{t},\mathbf{z}_{y}))$$
(3)

³Taken together, assumptions (2) and (3) constitute *strong ignorability*.

The structural inductive bias on the model is such that \mathbf{z}_y , and \mathbf{z}_t , and \mathbf{z}_c learn factors relevant to outcome and treatment, for which we provide explicit supervision, thereby leaving \mathbf{z}_o for all remaining factors. The combination of KL regularization and supervision helps prevent \mathbf{z}_o from learning information in $\mathbf{z}_{t,c,y}$, although whether information in \mathbf{z}_o is duplicated in other factors is not a problem in itself because \mathbf{z}_o is d-separated and does not participate in the calculation of the causal effect. Identifiability then rests on the assumption that $\mathbf{z}_{\{o,t,c,y\}}$ have been inferred from the proxies. The proof for identifiability in the case where all unoberved confounders have been inferred from proxy variables has been derived previously by Louizos et al. (2017) and Zhang et al. (2020). The factor \mathbf{z}_o is *d*-separated from \mathbf{t} and \mathbf{y} given \mathbf{x} , and does not affect the identification of the causal effect. i.e., $p(y|do(t), \mathbf{x}) = p(y|do(t), \mathbf{z}_{\{t,o,y,c\}}) = p(y|t, \mathbf{z}_y, \mathbf{z}_c)$ (see Zhang et al. 2020 and Louizos et al. 2017). We impose the priors and parameterizations denoted in Equation 3, where $D_{(.)}$ is the number of dimensions in the respective variable (latent or otherwise), and f_{1-11} and h_{1-6} represent fully connected neural network functions. The parameters for these neural networks are learnt via variational Bayesian approximate inference (Kingma & Welling, 2014) according to the following objective:

$$\mathcal{L}_{i}^{\text{ELBO}} = \sum_{i}^{N} \mathbb{E}_{q_{c}q_{t}q_{y}q_{o}} \left[\log p\left(\hat{\mathbf{x}}_{i} | \mathbf{z}_{t}, \mathbf{z}_{c}, \mathbf{z}_{y}, \mathbf{z}_{o}\right) + \log p\left(\hat{t}_{i} | \mathbf{z}_{t}, \mathbf{z}_{c}\right) + \log p\left(\hat{y}_{i} | t_{i}, \mathbf{z}_{y}, \mathbf{z}_{c}\right) \right] - \left[D_{KL} \left(q\left(\mathbf{z}_{t} | \mathbf{x}_{i}\right) | | p\left(\mathbf{z}_{t}\right)\right) + D_{KL} \left(q\left(\mathbf{z}_{c} | \mathbf{x}, \right) | | p\left(\mathbf{z}_{c}\right)\right) + D_{KL} \left(q\left(\mathbf{z}_{c} | \mathbf{x}, \right) | | p\left(\mathbf{z}_{c}\right)\right) \right] \right]$$

$$(4)$$

$$+ D_{KL} \left(q\left(\mathbf{z}_{y} | \mathbf{x}_{i}\right) | | p\left(\mathbf{z}_{y}\right)\right) + D_{KL} \left(q\left(\mathbf{z}_{c} | \mathbf{x}, \right) | | p\left(\mathbf{z}_{c}\right)\right) \right]$$

Note that all Gaussian variance parameterizations are diagonal. In cases where prior knowledge dictates a discrete rather than continuous outcome, equivalent parameterizations to those in Eqs. 3 may be employed. For example, in the IHDP dataset, the outcome data are standardized to have a variance of 1, and the outcome generation model becomes a Gaussian with variance also equal to 1. Note that separate treatment and outcome classifiers are used both during inference and generation $(Q_q, g_q \text{ and } Q_p, g_p \text{ resp.})$. The classifiers for inference have separate parameters to those use during generation. Predictors or classifiers of outcome incorporate the two-headed approach of (Shalit et al., 2017), and use ground-truth t are used for Q_q whereas samples \hat{t} are used for Q_p . For unseen test cases, either the ground-truth t or an sampled treatment \hat{t} from treatment classifier g_p may be used to simulate an outcome. During training \hat{t} is used.

We now introduce the targeted regularization, the purpose of which is to encourage the outcome to be independent of the treatment assignment. Following Eq. 1, we define the fluctuation sub-model and corresponding logistic loss for finding ϵ as:

$$\hat{Q}(g, t_i, \mathbf{z}_i^y, \mathbf{z}_i^c, \epsilon) = \sigma \left[\sigma^{-1} [Q(t_i, \mathbf{z}_i^y, \mathbf{z}_i^c)] + \epsilon \left(\frac{I(t_i = 1)}{g(t_i = 1; \mathbf{z}_i^t, \mathbf{z}_i^c)} - \frac{I(t_i = 0)}{g(t_i = 0; \mathbf{z}_i^t, \mathbf{z}_i^c)} \right) \right]$$
(5)

$$\xi_i(\hat{Q};\epsilon) = -y_i \log(\hat{Q}(g,t_i, \mathbf{z}_i^y, \mathbf{z}_i^c, \epsilon)) - (1 - y_i) \log(1 - \hat{Q}(g,t_i, \mathbf{z}_i^y, \mathbf{z}_i^c, \epsilon))$$
(6)

In Eq. 14, I is the indicator function. For an unbounded regression loss, mean squared error loss may be used (see appendix). Note that the logistic loss is suitable for continuous outcomes bounded between 0 and 1 (see van der Laan & Rose 2011, pp.121:132 for proof). Putting it all together, we then optimize to find generative parameters for functions h_{1-6} , inference parameters for functions f_{1-12} , and fluctuation parameter ϵ as follows:

$$\mathcal{L} = \min\left[\sum_{i}^{N} \left(\mathcal{L}_{i}^{\text{ELBO}} + \lambda_{TL}\xi_{i}(Q, g, \epsilon)\right)\right]; \quad \frac{\partial}{\partial\epsilon}\mathcal{L}^{*}\Big|_{\epsilon=0} = \bar{IC}^{*} = 0 \tag{7}$$

Where λ_{TL} represents a hyperparameter loss weight for the targeted regularization. At convergence, $\epsilon = 0$ when Q and g become consistent estimators, satisfying the conditions for the EIC (see Eq. 2 and reference van der Laan & Rose 2011, pp125:128).

A further element that differentiates our work from one other recent contribution (Shi et al., 2019) that uses targeted regularization is that the gradients resulting from ξ are *not* taken with respect to g_p or g_q . Targeted learning is concerned with de-biasing the outcome classifier Q using propensity scores from g. It therefore only makes sense to regularize Q using ξ , not g, which we assume to be consistent and unbiased. During training, the outcome and treatment predictions/classifications are

made during inference *and* generation. The targeted regularization is applied to the outcome classifications made during the generation stage, but gradients are taken with respect to all parameters that influence the outcome classifier, which include upstream parameters for Q_q as well as the more direct parameters Q_p . Note also that the gradients are not taken with respect to the general inference and generation parameters, only the parameters of the classification arms themselves (in the case of a binary outcome, these arms are represented by functions f_{10} and f_{11} for Q_q and h_1 and h_2 for Q_p).

In summary, the notable aspects of our model are as follows: the introduction of a new latent variable z_o for factors unrelated to outcome and/or treatment to help avoid misspecification and aid the recovery of the true underlying structure; the ability to estimate both individual level and average treatment effects; and, as far as we are aware, the first incorporation of targeted learning in a deep latent variable approach, and one which backpropagates the regularization gradients to specific outcome-related parameters in the network.

4 RELATED WORK

There are a number of ways to mitigate the problems associated with the confounding between the covariates and the treatment. For a review on such methods, readers are pointed to the recent survey by (Yao et al., 2020). Here we consider methods that utilize neural networks as part of their models, but note that many non-neural network methods exist (Chernozhukov et al., 2017; van der Laan & Rose, 2011; van der Laan & Starmans, 2014; Rubin, 2005; Hill, 2011; Athey & Imbens, 2016).

Perhaps the most similar works to ours are those of Dragonnet (Shi et al., 2019) and TEDVAE (Zhang et al., 2020). We discuss the differences between these and TVAE in turn. Dragonnet incorporates a similar TMLE regularization process which allows for the simultaneous optimization of Q and ϵ . However, the method sacrifices the ability to estimate individual treatment effect in preference to achieving good estimation of the average treatment effect across the sample. Indeed, they do not report PEHE. Finally, Dragonnet applies regularization to the entire network, whereas we specifically 'target' the regularization to the outcome prediction arm.

TEDVAE, on the other hand, builds on CEVAE (Louizos et al., 2017) and seeks inference and disentanglement of the latent instrumental, risk, and confounding factors from proxy variables with a variational approach. However, it has no means to allocate latent variables that are unrelated to treatment and/or outcome (i.e., TVAE's z_o). The advantage of including factors z_o with a variational penalty is that the model has the option to use them, or not to use them, depending on whether they are necessary (i.e. KL is pushed to zero). It is important not to force factors unrelated to treatment and outcome into $z_{\{c,y,t\}}$ because doing so restricts the overlap between the class of models that can be represented using TEDVAE, and the class of models describing the true data generating process.

Other methods include GANITE (Yoon et al., 2018) which requires adversarial training, and may therefore be more difficult to optimise. PM (Schwab et al., 2019), SITE (Yao et al., 2018), and MultiMBNN (Sharma et al., 2020) incorporate propensity score matching. TARNET (Shalit et al., 2017) inspired the two-headed outcome arm in our TVAE, as well as the three-headed architecture in (Shi et al., 2019). RSB (Zhang et al., 2019) incorporates a regularization penalty, based on the Pearson Correlation Coefficient, intended to reduce the association between latent variables predictive of treatment assignment and those predictive of outcome.

5 **EXPERIMENTS**

We perform an ablation study, beginning with (a) TVAE (base) which is equivalent to TEDVAE (b) TVAE with z_o , and (c) TVAE with both with z_o and targeted regularization ξ during training. In order to fairly evaluate the benefits of introducing z_o , we ensure that the total number of latent dimensions remains constant. We undertake the ablation study on a synthetic dataset which we call TVAEsynth, before comparing against methods on both the IHDP dataset (Hill, 2011; Gross, 1993) and the Jobs dataset (LaLonde, 1986; Smith & Todd, 2005; Dehejia & Wahba, 2002). In particular, the synthetic dataset was intentionally created such that not all covariates are exogenous and so that there exist some latent factors not related to outcome or treatment. Thus, we should expect a significant improvement in performance to occur with the introduction of z_o , demonstrating the importance of incorporating inductive bias that closely matches the true structure of the data. Note that while these datasets vary in whether the outcome variable is continuous (IHDP, TVAESynth) or binary (Jobs), the treatment variable is always binary. Whilst it is possible to undertake Targeted Learning on continuous treatment effects, we leave this to future work.

For the IHDP dataset, we evaluate our network on the Average Treatment Effect estimation error (eATE), and the Precision in Estimation of Heretogeneous Effect (PEHE). As per (Louizos et al., 2017; Shalit et al., 2017; Yao et al., 2018), for the Jobs dataset (for which we have only partial effect supervision) we evaluate our network on the Average Treatment effect on the Treated error (eATT) to approximate the eATE, and the policy risk R_{pol} to approximate the error on the CATE. See the appendix for definitions of these metrics. When estimating treatment effects, 100 samples are drawn for each set of input covariates x_i . We compare against Dragonnet (Shi et al., 2019), TEDVAE (Zhang et al., 2020), CEVAE (Louizos et al., 2017), GANITE (Yoon et al., 2018), Targeted Maximum Likelihood Estimation (TMLE) (van der Laan & Rose, 2018), and TARNET + variants (Shalit et al., 2017). We provide results for both within sample and out-of-sample performance. It is worth noting that within sample and out-of-sample results are equally valid for treatment effect estimation, because the network is never supervised on treatment effect (Shi et al., 2019).

For model selection we follow Louizos et al. (2017) and Zhang et al. (2020) and use the minimum validation loss on the total objective function. Whilst some model selection heuristics exist that serve as surrogates for the eATE itself (e.g., see Hassanpour & Greiner 2019 or Athey & Imbens 2016) we take the same view as Zhang et al. (2020), namely that the development of our model 'should be self-sufficient and not rely on others'. Furthermore, we undertake minimal hyperparameter tuning for the simple reason that, in real-world applications, the supervision required for effective tuning would not be available. For all experiments, we undertake 100 replications and provide mean and standard error. See the appendix for details on these datasets and the architecture, as well as training and testing details.

Table 1: Means and standard errors for the ablation study using the TVAESynth dataset. Here, 'oos' means out-of-sample, 'ws' means within sample. '+ z_o ' indicates the introduction of the miscellaneous factors, and '+ ξ ' indicates targeted regularization with subscript indicating its weight in the loss.

Method	$\sqrt{\epsilon_{PEHE}}$ ws	$\sqrt{\epsilon_{PEHE}}$ oos	ϵ_{ATE} ws	ϵ_{ATE} oos
TVAE (base/TEDVAE)	.179±.003	.178±.003	$.128 \pm .005$.128±.005
TVAE + \mathbf{z}_o	$.166 \pm .003$	$.166 {\pm} .003$	$.069 {\pm} .004$	$.069 {\pm} .004$
TVAE + \mathbf{z}_o + $\xi_{\lambda=0.1}$	$.150 {\pm} .002$.150±.003	$.048 {\pm} .004$	$.048 {\pm} .004$

Ablation Study - TVAESynth: The results for the ablation study on the synthetic dataset are shown in Table 1. The results demonstrate that both eATE and PEHE are significantly improved by the incorporation of z_o , and further improved with the introduction of targeted regularization for both within sample and out of sample testing.

IHDP Results: Results on IHDP are shown in Table 2 and indicate state of the art performance for both within sample and out-of-sample eATE and PEHE. The results corroborate the ablation results in Table 1, in that the incorporation of z_o and targeted regularization result in monotonic improvement above TEDVAE. TVAE is outperformed only by Dragonnet on within-sample eATE performance. However, this method does not provide estimations for indvidual CATE, and is limited to the estimation of average treatment effects.

Jobs Results: The results for the Jobs data are shown in Table 3. GANITE was found to perform the best across most metrics, although this method has been argued to be more reliant on larger sample sizes than others, on the basis that it performs relatively poorly on the smaller IHDP dataset (Yoon et al., 2018). Furthermore, GANITE relies on potentially unstable/unreliable adversarial training (Moyer et al., 2018; Lezama, 2019; Gabbay & Hosen, 2019). Finally, TVAE outperforms GANITE on eATT, is consistent (beyond 2 decimal places) across out-of-sample and within-sample evaluations and has a lower standard err, and is competitive across all metrics. On this dataset, the concomitant improvements associated with the additional latent factors and targeted learning were negligible.

Table 2: Means and standard errors for evaluation on the semi-synthetic IHDP dataset Hill (2011). Results from: (Louizos et al., 2017; Shalit et al., 2017; Zhang et al., 2020; Yoon et al., 2018). Here, 'oos' means out-of-sample and 'ws' means within sample. '+ z_o ' indicates the introduction of the miscellaneous factors, and '+ ξ ' indicates targeted regularization with subscript indicating its weight in the loss.

Method	$\sqrt{\epsilon_{PEHE}}$ ws	$\sqrt{\epsilon_{PEHE}}$ oos	ϵ_{ATE} ws	ϵ_{ATE} oos
TMLE (van der Laan & Rose, 2018)	$5.0 \pm .20$	-	$.30 {\pm} .01$	-
CEVAE (Louizos et al., 2017)	$2.7 \pm .10$	$2.6 \pm .10$	$.34 {\pm} .01$	$.46 {\pm} .020$
TARNet (Shalit et al., 2017)	$.88 {\pm} .00$	$.95 {\pm} .00$	$.26 {\pm} .01$	$.28 {\pm} .01$
CFR-MMD (Shalit et al., 2017)	$.73 {\pm} .00$	$.78 {\pm} .00$	$.30 {\pm} .01$	$.31 {\pm} .01$
CFR-Wass (Shalit et al., 2017)	$.71 {\pm} .00$	$.76 {\pm} .00$	$.25 {\pm} .01$	$.27 {\pm} .01$
TEDVAE (Zhang et al., 2020)	$.62 \pm .11$.63±.12	-	$.20 {\pm} .05$
GANITE (Yoon et al., 2018)	$1.9 \pm .40$	$2.4 \pm .40$	$.43 {\pm} .05$	$.49 {\pm} .05$
Dragonnet (Shi et al., 2019)	-	-	.14±.01	$.20 {\pm} .01$
TVAE (w/ $\mathbf{z}_0, \xi_{\lambda=0.0}$)	.57±.03	.57±.03	.16±.01	.16±.01
TVAE (w/ $\mathbf{z}_0, \xi_{\lambda=0.4}$)	.52±.02	.54±.02	$.15 {\pm} .01$.16±.01

Table 3: Means and standard errors for evaluation on the real-world Jobs dataset (LaLonde, 1986; Smith & Todd, 2005; Shalit et al., 2017; Yoon et al., 2018). Results taken from: (Louizos et al., 2017; Zhang et al., 2020). Here, 'oos' means out-of-sample and 'ws' means within sample. '+ z_o ' indicates the introduction of the miscellaneous factors, and '+ ξ ' indicates targeted regularization with subscript indicating its weight in the loss.

Method	R_{pol} ws	R_{pol} oos	ϵ_{ATT} ws	ϵ_{ATT} oos
TMLE (van der Laan & Rose, 2018)	$.22 \pm .00$	-	$.02 {\pm} .01$	-
CEVAE (Louizos et al., 2017)	$.15 {\pm} .00$	$.26 {\pm} .00$	$.02 {\pm} .01$	$.03 {\pm} .01$
TARNet (Shalit et al., 2017)	$.17 {\pm} .00$	$.21 \pm .00$	$.05 {\pm} .02$	$.11 \pm .04$
CFR-MMD (Shalit et al., 2017)	$.18 {\pm} .00$	$.21 {\pm} .00$	$.04 {\pm} .01$	$.08 {\pm} .03$
CFR-Wass (Shalit et al., 2017)	$.17 {\pm} .00$	$.21 {\pm} .00$	$.04 {\pm} .01$	$.09 {\pm} .03$
GANITE (Yoon et al., 2018)	.13±.00	$.14 {\pm} .00$.01±.01	$.06 {\pm} .03$
TEDVAE (Zhang et al., 2020)	-	-	$.06 {\pm} .00$	$.06 {\pm} .00$
TVAE (w/ $\mathbf{z}_0, \xi_{\lambda=0}$)	$.16 {\pm} .00$	$.16 {\pm} .00$.01±.00	$.01 {\pm} .00$
TVAE (w/ $\mathbf{z}_0, \xi_{\lambda=1}$)	$.16 {\pm} .00$	$.16 {\pm} .00$.01±.00	$.01 {\pm} .00$

6 CONCLUSION

In this work we aimed to improve existing latent variable models for causal parameter estimation in two ways: Firstly, by introducing a latent variable to model factors unrelated to treatment and outcome, thereby enabling the model to more closely reflect the data structure; and secondly, by incorporating a targeted learning regularizer with selected backpropagation to further de-bias outcome predictions. Our experiments demonstrated concomitant improvements in performance, and our comparison against other methods demonstrated TVAE's ability to compete with and/or exceed state of the art for both individual as well as average treatment effect estimation. For future work, we plan to explore the application of TVAE to sequential data with continuous or categorical treatment.

REFERENCES

- A. A. Alemi, I. FIscher, J. V. Dillon, and K. Murphy. Deep variational information bottleneck. *arXiv:1612.00410v7*, 2017.
- S. Athey and G.W. Imbens. Recursive partitioning for heterogeneous causal effects. *Proc. of the National Academy of Sciences of the USA*, 113(27), 2016.
- Eli Bingham, Jonathan P. Chen, Martin Jankowiak, Fritz Obermeyer, Neeraj Pradhan, Theofanis Karaletsos, Rohit Singh, Paul A. Szerlip, Paul Horsfall, and Noah D. Goodman. Pyro: Deep universal probabilistic programming. *J. Mach. Learn. Res.*, 20, 2019.

- D. M. Blei, A. Kucukelbir, and J. D. McAuliffe. Variational inference: a review for statisticians. arXiv:1601.00670v9, 2018.
- L. Bottou, J. Peters, Quinonero-Candela J., D. X. Charles, D. M. Chickering, E. Portugaly, D. Ray, P. Simard, and E. Snelson. Counterfactual reasoning and learning systems: the example of computational advertising. *Journal of Machine Learning Research*, 14, 2013.
- C. P. Burgess, I. Higgins, A. Pal, L. Matthey, N. Watters, G. Desjardins, and A. Lerchner. Understanding disentangling in Beta-VAE. *arXiv:1804.03599v1*, 2018.
- V. Chernozhukov, D. Chetverikov, M. Demirer, E. Duflo, C. Hansen, and W. Newey. Double/debiased/Neyman machine learning of treatment effects. *American Economic Review*, 5, 2017.
- B. Dai, Z. Wang, and D. Wipf. The usual suspects? reassessing blame for VAE posterior collapse. arXiv:1912.10702v1, 2019.
- A. D'Amour. On multi-cause causal inference with unobserved confounding: Counterexamples, impossibility and alternatives. *Proceedings of the 22nd International Conference on Artificial Intelligence and Statistics*, 89, 2019.
- A. Deaton and N. Cartwright. Understanding and misundertstanding randomized controlled trials. Social Science and Medicine, 210:2–21, 2018. doi: 10.1016/j.socscimed.2017.12.005.
- R. H. Dehejia and S. Wahba. Propensity score-matching methods for nonexperimental causal studies. *Review of Economics and Statistics*, 84(1):151–161, 2002.
- V. Dorie. Non-parametrics for causal inference. https://github.com/vdorie/npci, 2016.
- A. Gabbay and Y. Hosen. Demystifying inter-class disentanglement. arXiv:1906.11796v2, 2019.
- I. J. Goodfellow, J. Pouget-Abadie, M. Mirza, B. Xu, D. Warde-Farley, S. Ozair, A. Courville, and Y. Bengio. Generative adversarial nets. *arXiv:1406.2661*, 2014.
- R. T. Gross. Infant health and development program (IHDP): enhancing the outcomes of low birth weight, premature infants in the United States. MI: Inter-university Confortium for Political and Social Research, Ann Arbor, 1993.
- M.P. Grosz, J.M. Rohrer, and F. Thoemmes. The taboo against explicit causal inference in nonexperimental psychology. *Perspectives on Psychological Science*, pp. 1–13, 2020. doi: 10.1177/1745691620921521.
- R. Guo, J. Li, and H. Liu. Learning individual causal effects from networked observational data. *Association for Computing Machinery*, 2020.
- F. R. Hampel. The influence curve and its role in robust estimation. *Journal of the American Statistical Association*, 69(346):383–393, 1974.
- N. Hassanpour and R. Greiner. Counterfactual regression with importance sampling weights. *Proceedings of the 28th International Joint Conference on Artificial Intellgence (IJCAI-19)*, 2019.
- N. Hassanpour and R. Greiner. Learning disentangled representations for counterfactual regression. *ICLR*, 2020.
- M. Hernan. The c-word: scientific euphemisms do not improve causal inference from observational data. *American Journal of Public Health*, 108(5):625–626, 2018. doi: 10.2105/AJPH.2018. 304337.
- I. Higgins, L. Matthey, A. Pal, C. Burgess, X. Glorot, M. Botvinick, S. Mohamed, and A. Lerchner. Beta-VAE: Learning basic visual concepts with a constrained variational framework. *ICLR*, 2017.
- J. L. Hill. Bayesian nonparametric modeling for causal inference. *Journal of Computational and Graphical Statistics*, 20(1), 2011.

- G.W. Imbens and D.B. Rubin. *Causal inference for statistics, social, and biomedical sciences. An Introduction.* Cambridge University Press, New York, 2015.
- A. Jesson, S. Mindermann, U. Shalit, and Y. Gal. Identifying causal effect inference failure with uncertainty-aware models. arXiv:2007.00163v1, 2020.
- D. P. Kingma and J. L. Ba. Adam: a method for stochastic optimization. arXiv:1412.6980v9, 2017.
- D. P. Kingma and M. Welling. Auto-encoding variational Bayes. arXiv:1312.6114v10, 2014.
- N. Kreif and K. DiazOrdaz. Machine learning in policy evaluation: new tools for causal inference. *arXiv:1903.00402v1*, 2019.
- R.J. LaLonde. Evaluating the econometric evaluations of training programs with experimental data. *The American Economic Review*, pp. 604–620, 1986.
- J. Lezama. Overcoming the disentanglement vs reconstruction trade-off via Jacobian supervision. *ICLR*, 2019.
- F. Locatello, S. Bauer, M. Lucic, G. Ratsch, S. Gelly, B. Scholkopf, and Bachem O. Challenging common assumptions in the unsupervised learning of disentangled representations. arXiv:1811.12359v3, 2019.
- C. Louizos, U. Shalit, J. Mooij, D. Sontag, R. Zemel, and M. Welling. Causal effect inference with deep latent-variable models. *31st Conference on Neural Information Processing Systems*, 2017.
- J. Lucas, G. Tucker, R. Grosse, and M Norouzi. Understanding posterior collapse in generative latent variable models. *ICLR*, 2019a.
- J. Lucas, G. Tucker, R. Grosse, and M. Norouzi. Don't blame the ELBO! a linear VAE perspective on posterior collapse. *arXiv:1911.02469v1*, 2019b.
- M. Maier, K. Marazopoulou, D. Arbour, and D. David. A sound and complete algorithm for learning causal models from relational data. *Proceedings of the 29th Conf. on Uncertainty in Artificial Intelligence*, 2013.
- I. Mayer, J. Josse, and J. P. Raimundo, F.and Vert. MissDeepCausal: causal inference from incomplete data using deep latent variable models. *arXiv:2002.10837v1*, 2020.
- M.R. Montgomery, M. Gragnolati, K.A. Burke, and E. Paredes. Measuring living standards with proxy variables. *Demography*, 37(2):155–174, 2000.
- J. M. Mooij, O. Stegle, D. Janzing, K. Zhang, and B. Scholkopf. Probabilistic latent variable models for distinguishing between cause and effect. *NIPS*, pp. 1687–1695, 2010.
- D. Moyer, S. Gao, R. Brekelmans, G. V. Steeg, and A. Galstyan. Invariant representations without adversarial training. *NeurIPS*, 2018.
- H. Oktay. Using latent variable models to improve causal estimation. PhD thesis, Doctoral Dissertation 1205, 2018.
- J. Pearl. Causality. Cambridge University Press, Cambridge, 2009.
- M. Petersen, L. Balzer, D. Kwarsiima, N. Sang, G. Chamie, J. Ayieko, J. Kabami, A. Owaraganise, T. Liegler, F. Mwangwa, and K. Kadede. Association of implementation of a universal testing and treatment intervention with HIV diagnosis, receipt of antiretroviral therapy, and viral suppression in East Africa. *Journal of American Medical Association*, 317(21):2196–2206, 2017. doi: 10. 1001/jama.2017.5705.
- M. Rolinek, D. Zietlow, and G. Martius. Variational autoencoders pursue PCA directions (by accident). arXiv:1812.06775v2, 2019.
- P. R. Rosenbaum and D. B. Rubin. The central role of the propensity score in observational studies for causal effects. *Biometrika*, 70(1):41–55, 1983.

- D. B. Rubin. Causal inference using potential outcomes: Design, modeling, decisions. *Journal of the American Statistical Association*, 100(469):322–331, 2005. doi: 10.1198/016214504000001880.
- M. S. Schuler and S. Rose. Targeted maximum likelihood estimation for causal inference in observational studies. *American Journal of Epidemiology*, 185(1):65–73, 2016.
- P. Schwab, L. Linhardt, and W. Karlen. Perfect match: a simple method for learning representations for counterfactual inference with neural networks. arXiv:1810.00656v5, 2019.
- U. Shalit, F. D. Johansson, and D. Sontag. Estimating individual treatment effect: generalization bounds and algorithms. arxiv:1606.03976v5, 2017.
- A. Sharma, G. Gupta, A. Prasad, R.and Chatterjee, L. Vig, and G. Shroff. MultiMBNN: matched and balanced causal inference with neural networks. arXiv:2004.13446v2, 2020.
- C. Shi, D. M. Blei, and V. Veitch. Adapting neural networks for the estimation of treatment effects. 33rd Conference on Neural Information Processing Systems, 2019.
- B. Siegerink, W. den Hollander, M. Zeegers, and R. Middelburg. Causal inference in law: an epidemiological perspective. *European Journal of Risk Regulation*, 7(1):175–186, 2016. doi: 10.1017/S1867299X0000547X.
- R. Silva, R. Scheine, G. Clark, and P. Spirtes. Learning the structure of linear latent variable models. *Journal of Machine Learning Research*, 7:191–246, 2006.
- J. A. Smith and P. E. Todd. Does matching overcome LaLonde's critique of nonexperimental estimators? *Journal of Econometrics*, 125(1):305–353, 2005.
- N. Tishby and N. Zaslavsky. Deep learning and the information bottleneck principle. *arXiv:1503.02406v1*, 2015.
- M. J. van der Laan and S. Rose. *Targeted Learning Causal Inference for Observational and Experimental Data*. Springer International, New York, 2011.
- M. J. van der Laan and S. Rose. *Targeted Learning in Data Science*. Springer International, Switzerland, 2018.
- M. J. van der Laan and R. J. C. M. Starmans. Entering the era of data science: targeted learning and the integration of statistics and computational data analysis. *Advances in Statistics*, 2014.
- M. J. Vowels. Limited functional form, misspecification, and unreliable interpretations in psychology and social science. arXiv:2009.10025, 2020.
- M. J. Vowels, N. C. Camgoz, and R. Bowden. Gated variational autoencoders: Incorporating weak supervision to encourage disentanglement. *arXiv:1911.06443v1*, 2019.
- M. J. Vowels, N. C. Camgoz, and R. Bowden. NestedVAE: Isolating common factors via weak supervision. *Conference on Computer Vision and Pattern Recognition*, 2020.
- L. Yao, S. Li, Y. Li, M. Huai, J. Gao, and A. Zhang. Representation learning for treatment effect estimation from observational data. 32nd Conference on Neural Information Processing Systems (NeurIPS), 2018.
- L. Yao, Z. Chu, S. Li, Y. Li, J. Gao, and A. Zhang. A survey on causal inference. *arXiv:2002.02770*, 2020.
- J. Yoon, J. Jordan, and M. van der Schaar. GANITE: Estimation of individualized treatment effects using generative adversarial nets. *ICLR*, 2018.
- W. Zhang, L. Liu, and J. Li. Treatment effect estimation with disentangled latent factors. *arXiv:2001.10652*, 2020.
- Z. Zhang, Q. Lan, Y. Wang, N. Hassanpour, and R. Greiner. Reducing selectin bias in counterfactual reasoning for individual treatment effects estimation. 33rd Conference on Neural Information Processing Systems, 2019.

A APPENDIX

This appendix includes details on the metrics used for evaluating treatment effect estimation; background on Variational AutoEncoders; a formulation of the targeted learning regularizer for continuous, unbounded outcomes (as used with the IHDP dataset evaluations); details about the datasets; and training, testing, hyperparameters, architecture, and hardware details. Source code and data (including IHDP (Hill, 2011; Gross, 1993), Jobs (LaLonde, 1986; Smith & Todd, 2005), and our own TVAESynth) is also attached as supplementary material for reproducibility purposes, in a separate folder.

A.1 METRICS

This section presents the metrics used for the experiments (Section 5 in the main paper).

The Average Treatment Effect (ATE) and error on the estimation of ATE (eATE) is given in Eq. 8.

$$\hat{\tau}(Q;\mathbf{x}) = \frac{1}{N} \sum_{i=1}^{N} (Q(1,\mathbf{x}_i) - Q(0,\mathbf{x}_i)), \quad \epsilon_{ATE} = \left|\frac{1}{N} \sum_{i=1}^{N} (\hat{\tau}(Q;\mathbf{x}_i) - \tau(\mathbf{x}))\right|$$
(8)

To estimate the error on the model's capacity to model the Conditional Average Treatment effect (CATE), the Precision in Estimation of Heteogenous Effects (PEHE) is given in Eq. 9.

$$\epsilon_{PEHE} = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (\hat{\boldsymbol{\tau}}(Q; \mathbf{x}_i) - \boldsymbol{\tau}(\mathbf{x}_i))^2}$$
(9)

It can be seen from Equations 8 that the ATE is essentially the expectation of the conditional treatment effect (conditioned on the covariates for each individual) over the data set (Jesson et al., 2020). For scenarios when a proportion of the dataset is from a Randomized Controlled Trial (RCT), as is the case for the Jobs dataset, we may use the error on the estimation of Average Treatment effect on the Treated (ATT), which is given in Eq. 10 (Shalit et al., 2017; Louizos et al., 2017):

$$eATT = \left|\frac{1}{|T_1|} \sum_{i \in T_1} y_i - \frac{1}{|T_0|} \sum_{j \in T_0} y_j - \frac{1}{|T_1|} \sum_{i \in T_1} (Q(1, \mathbf{x}_i) - Q(0, \mathbf{x}_i))\right|$$
(10)

where $T = T_1 \cup T_0$ constitutes all individuals in the RCT, and the subscripts denote whether or not those individuals were in the treatment (subscript 1) or control groups (subscript 0). The first two terms in Eq. 10 comprise the true ATT, and the third term the estimated ATT. In datasets where the ground-truth for the CATE is not available (as is the case with the Jobs dataset) we may use the policy risk as a proxy for PEHE:

$$\mathcal{R}_{pol} = 1 - \left(\mathbb{E}[\mathbf{y}^{t=1} | \pi(\mathbf{x}) = 1] p(\pi(\mathbf{x}) = 1) + \mathbb{E}[\mathbf{y}^{t=0} | \pi(\mathbf{x}) = 0] p(\pi(\mathbf{x}) = 0) \right)$$
(11)

where $\pi(\mathbf{x}_i) = 1$ is the policy to treat when $\hat{y}_i^{t=1} - \hat{y}_i^{t=0} > \epsilon$, and $\pi(\mathbf{x}_i) = 0$ is the policy not to treat otherwise (Yao et al., 2018; Shalit et al., 2017). ϵ is a treatment threshold. This threshold can be varied to understand how treatment inclusion rates affect the policy risk. For our experiments we set ϵ to zero, as per (Shalit et al., 2017; Louizos et al., 2017).

A.2 VARIATIONAL AUTOENCODERS

This part of the appendix is referenced in Section 3 (Methodology) of the main paper.

We now consider the theory behind a popular and powerful latent variable representation learning and density estimation method known as Variational AutoEncoders (VAEs) (Kingma & Welling, 2014). Although adversarial methods (Goodfellow et al., 2014) have been shown to be effective for density estimation, they are also troublesome to train. Hence we choose VAEs which have more stable training dynamics (Moyer et al., 2018). A further motivation for the use of VAEs relates to their ability to infer latent variables. The Ignorability assumption holds that all confounders are observed, and much causal inference is undertaken with this assumption (Mooij et al., 2010; Maier et al., 2013; Oktay, 2018; Silva et al., 2006). The use of a latent variable model allows us to infer unobserved/hidden confounders (Louizos et al., 2017), although the use of VAEs means that there is some uncertainty concerning the guarantees of the learned model (Rolinek et al., 2019; Dai et al., 2019; Lucas et al., 2019a;b). This uncertainty notwithstanding, previous implementations for latent variable models with causal inference show promising results in comparison with other methods (Louizos et al., 2017; Mayer et al., 2020; Hassanpour & Greiner, 2020).

We wish to encode the *m*-dimensional covariates \mathbf{x} on a manifold via a stochastic mapping $p(\mathbf{z}|\mathbf{x})$, where latent variables \mathbf{z} provide a compact representation. The distribution $p_{\theta}(\mathbf{z}|\mathbf{x})$ also serves as the posterior to the generative model $p_{\theta}(\mathbf{x}) = \int p_{\theta}(\mathbf{x}|\mathbf{z})p(\mathbf{z})d\mathbf{z}$ having parameters θ . Marginalizing out \mathbf{z} is usually intractable, and the true posterior $p(\mathbf{z}|\mathbf{x})$ is unknown, so a simpler approximating posterior $q_{\phi}(\mathbf{z}|\mathbf{x})$ having parameters ϕ is introduced (Blei et al., 2018) such that:

$$\log p_{\theta}(\mathbf{x}) = \mathbb{E}_{q_{\phi}(\mathbf{z}|\mathbf{x})} \left[\log \frac{p_{\theta}(\mathbf{x}|\mathbf{z})p(\mathbf{z})}{q_{\phi}(\mathbf{z}|\mathbf{x})} + \log \frac{q_{\phi}(\mathbf{z}|\mathbf{x})}{p_{\theta}(\mathbf{z}|\mathbf{x})} \right]$$
(12)

Taken together with the expectation, the second term on the right hand side Eq. 12 represents the Kullback-Liebler Divergence (KLD) between the approximating posterior and the true posterior. The KLD is always positive, and by ignoring this term, we are left with a lower-bound on the log-likelihood known as the variational lower bound, or Evidence Lower BOund (ELBO). The VAE provides the means to scale this amortized variational inference to intractable, high-dimensional problems, and minimizes the negative log likelihood over a dataset of N samples by adjusting the parameters of neural networks $\{\theta, \phi\}$ according to the ELBO.

$$\frac{1}{N}\sum_{i=1}^{n} -\log p_{\theta}(\mathbf{x}_{i}) \leq \frac{1}{N}\sum_{i=1}^{N} \left(-\mathbb{E}_{q_{\phi}(\mathbf{z}|\mathbf{x}_{i})}\left[\log p_{\theta}\left(\mathbf{x}_{i}|\mathbf{z}\right)\right] + \beta \mathbb{D}_{KL}\left[q_{\phi}(\mathbf{z}|\mathbf{x}_{i})\|p(\mathbf{z})\right]\right), \quad (13)$$

where $\beta = 1$ is used for the standard variational approximation procedure, but may be set empirically (Higgins et al., 2017), annealed (Burgess et al., 2018) or optimized according to the Information Bottleneck principle (Alemi et al., 2017; Tishby & Zaslavsky, 2015). The first term in Eq. 13 is the negative log-likelihood and is calculated in the form of a reconstruction error. The second term is the KLD between the approximating posterior and the prior, and therefore acts as a prior regularizer. Typically, the family of isotropic Gaussian distributions is chosen for the posterior $q_{\phi}(.)$, and an isotropic Gaussian with unit variance for the prior $p(\mathbf{z})$ which helps to encourage disentanglement (Vowels et al., 2020; 2019; Higgins et al., 2017).

A.3 TARGETED REGULARIZATION FOR BOUNDED AND UNBOUNDED OUTCOMES

This part of the appendix is referenced in Section 3 (Methodology) of the main paper. In contrast with Eq. 6 in the main paper, this formulation of targeted regularization is for an unbounded, continuous outcome (as is the case in the IHDP experiments). A mean-squared error version of ξ , similar to the one found in (Shi et al., 2019), is given as follows:

$$\hat{Q}(t_i, \mathbf{z}_i^y, \mathbf{z}_i^c, \epsilon) = Q(t_i, \mathbf{z}_i^y, \mathbf{z}_i^c) + \epsilon \left(\frac{I(t_i = 1)}{g(t_i = 1; \mathbf{z}_i^t, \mathbf{z}_i^c)} - \frac{I(t_i = 0)}{g(t_i = 0; \mathbf{z}_i^t, \mathbf{z}_i^c)} \right)$$
(14)

$$\xi_i(\hat{Q}, g; \phi_{c,t,y}, \epsilon) = (y_i - Q(t_i, \mathbf{z}_i^y, \mathbf{z}_i^c, \epsilon))^2$$
(15)

However, we stress that this formulation is only appropriate for unbounded outcomes, and not continuous outcomes in general (van der Laan & Rose, 2011). For continuous, bounded outcomes, as well as binary outcomes, the NLL formulation (as in the main paper) has been shown to be more appropriate, and theoretically sound.



Figure 4: The DAG for TVAESynth dataset.

A.4 DATASETS

This part of the appendix is referenced in Section 5 (Experiments) of the main paper.

We utilize 100 replications of the semi-synthetic *Infant Health and Development Program (IHDP)* dataset (Hill, 2011; Gross, 1993)⁴ The linked version (see footnote) corresponds with setting A of the NPCI data generating package (Dorie, 2016), which is the version that is used most frequently in other comparisons (e.g., see Shi et al. 2019; Shalit et al. 2017; Yao et al. 2018) and comprises 608 untreated and 139 treated samples (747 in total). There are 25 covariates, 19 of which are discrete/binary, and the rest are continuous. The outcome for the IHDP data is continuous and unbounded. Similarly to (Louizos et al., 2017; Shalit et al., 2017) and others, we utilize a 60/30/10 train/validation/test split.

We also utilize the job outcomes dataset which we refer to as *Jobs* (LaLonde, 1986; Smith & Todd, 2005).⁵ Unlike the IHDP datset, Jobs is real-world data with a binary outcome. We follow a similar procedure to (Shalit et al., 2017) who indicate that they used the Dehejia and Wahba (Dehejia & Wahba, 2002) and PSID comaprison samples. The Dehejia-Wahba sample comprises 260 treated samples and 185 control samples, along with the PSID comparison group comprising 2490 samples'. The dataset contains a mixture of observational and Randomized Controlled Trial data. Similarly to (Louizos et al., 2017; Shalit et al., 2017) and others, we utilize a 56/24/20 train/validation/test split, and undertake 100 runs with varying random split allocations in order to acquire an estimate of average performance and standard error. Note that, between models, the same random seed is used both for intialization as well as dataset splitting, and therefore the variance due to these factors is equivalent across experiments.

Finally, we introduce a new synthetic dataset named *TVAESynth* which follows the structure shown in Figure 4. While the weightings are chosen relatively arbitrarily, the structure is intentionally designed such that there are a mixture of exogenous and endogenous covariates. This enables us to compare the performance of TVAE with and without z_o (while keeping the total number of latent dimensions constant). These data comprised 1000 samples, a continuous outcome and binary treatment, 8 covariates, and generative/structural equations as follows:

$$\mathbf{U}_{z_o, z_c, z_t, z_y, y} \sim \mathcal{N}(\mathbf{0}, \mathbf{1}) \quad \mathbf{U}_{x_1, x_4, t} \sim \text{Bernoulli}(0.5) \quad \mathbf{U}_{x_{2:3}, x_{5:8}} \sim \mathcal{N}(\mathbf{0}, \mathbf{1})$$
(16)

$$\mathbf{z}_o = \mathbf{U}_{z_o} \quad \mathbf{z}_y = \mathbf{U}_{z_y} \quad \mathbf{z}_t = \mathbf{U}_{z_t} \quad \mathbf{z}_c = \mathbf{U}_{z_c} \tag{17}$$

 $\mathbf{x}_1 \sim \text{Bernoulli}(\sigma(\mathbf{z}_t + 0.1(\mathbf{U}_{x_1} - 0.5))) \ \mathbf{x}_2 \sim \mathcal{N}(0.4\mathbf{z}_o + 0.3\mathbf{z}_c + 0.5\mathbf{z}_y + 0.1\mathbf{U}_{x_2}, 0.2)$ (18)

⁴Available from https://www.fredjo.com/, https://github.com/WeijiaZhang24/ TEDVAE/ and elsewhere, and included in supplementary folder with source code.

⁵Available from https://users.nber.org/~rdehejia/data/.nswdata2.html.



Figure 5: Block architectural diagram for TVAE's inference and generation models (number of layers in Pred y 1 & 2 and Pred x blocks varies by experiment, as does the number of neurons in each layer (see details in text).

$$\mathbf{x}_{3} \sim \mathcal{N}(0.2\mathbf{z}_{o} + 0.2\mathbf{z}_{c} + 1.2\mathbf{z}_{t} + 0.1\mathbf{U}_{x_{3}}, 0.2) \quad \mathbf{x}_{4} \sim \text{Bernoulli}(\sigma(0.6\mathbf{z}_{o} + 0.1(\mathbf{U}_{x_{4}} - 0.5))) \quad (19)$$

$$\mathbf{x}_5 \sim \mathcal{N}(0.6\mathbf{z}_t + 0.1\mathbf{U}_{x_5}, 0.1) \quad \mathbf{x}_6 \sim \mathcal{N}(0.9\mathbf{z}_y + 0.1\mathbf{U}_{x_6}, 0.1)$$
 (20)

$$\mathbf{x}_7 \sim \mathcal{N}(0.5\mathbf{z}_o + 0.1\mathbf{U}_{x_7}, 0.1) \quad \mathbf{x}_8 \sim \mathcal{N}(0.5\mathbf{z}_o + 0.1\mathbf{U}_{x_8}, 0.1)$$
 (21)

$$\mathbf{t}_p = \sigma(0.2\mathbf{z}_c + 0.8\mathbf{z}_t + 0.1\mathbf{U}_t) \quad \mathbf{t} \sim \text{Bernoulli}(\mathbf{t}_p)$$
(22)

$$y := 0.2\mathbf{z}_c + 0.5\mathbf{z}_u \mathbf{t} + 0.2\mathbf{t} + 0.1\mathbf{U}_{\mathbf{v}}$$
(23)

Interventional distributions were generated by setting t equal to 1 and 0 thereby yielding ground truth ATE (≈ 0.8) and individual effects for evaluation purposes. The number of treated individuals is $\approx 70\%$. Note that for every replication, the 1000 sample evaluation set is regenerated, and so may have an empirical statistics that vary. For experiments, we use a 80/20 train/test split.

A.5 MODEL AND HYPERPARAMETERS

This part of the appendix is referenced in Section 3 (Methodology), as well as in Section 5 (Experiments) of the main paper. A block diagram of the TVAE architecture is shown in Fig. 5. For continuous outcomes (as in the IHDP dataset) we standardize the values and model as a Gaussian with a fixed variance of 1, and a mean determined by the outcome arm. All binary outcomes in the model (e.g. treatment or the relevant covariates) are modelled as Bernoulli distributed with a probability determined by the associated neural network function.

We now list the hyperparameters that were explored as part of model training. There may be room to improve on our figures with further hyperparameter tuning. However, given that the tuning of hyperparameters in a causal inference paradigm is problematic in general, we intentionally limited the space of hyperparameters (similarly to Zhang et al. 2020). Bold font indicates the settings that were used in the presented results.

Hyperparameter settings for IHDP dataset experiments were: hidden layers: **3**; the weight on targeted regularization $\lambda_{TL} = \{0.0, 0.1, 0.2, 0.4, 0.6, 0.8, 1.0\}$; an Adam (Kingma & Ba, 2017) optimizer with learning rate $LR = \{1e-3, 1e-4, 5e-5\}$; number of hidden neurons was **300**; number of

layers = 4; dimensionality of the latent factors was $D_{z_t} = D_{z_t} = 10, D_{z_c} = 15, D_{z_o} = 5$; batch size of 200; number of epochs 200; weight regularization 1e-4; and learning rate decay 5e-4.

Hyperparameter settings for the Jobs dataset experiments were: hidden layers: **3**; the weight on targeted regularization $\lambda_{TL} = \{0.0, 0.1, 0.2, 0.4, 0.6, 0.8, 1.0\}$; an Adam (Kingma & Ba, 2017) optimizer with learning rate $LR = \{5e-5, 1e-5\}$; number of hidden neurons was **200**; number of layers = **2**; dimensionality of the latent factors was $D_{z_t} = D_{z_t} = 6$, $D_{z_c} = 8$, $D_{z_o} = 4$; batch size of **200**; number of epochs **200**; weight regularization 1e-4; and learning rate decay **5e-4=3**.

Hyperparameter settings for the TVAESynth dataset experiments were: hidden layers: 2; the weight on targeted regularization $\lambda_{TL} = \{0.0, 0.1, 0.2, 0.4, 0.6, 0.8, 1.0\}$; an Adam (Kingma & Ba, 2017) optimizer with learning rate LR = 5e-5; number of hidden neurons was 20; number of layers = 2; dimensionality of the latent factors was $D_{z_t} = D_{z_t} = D_{z_c} = 2$, $D_{z_o} = 1$; batch size of 200; number of epochs 40; weight regularization 1e-4; and learning rate decay 5e-3.

As described in the main paper, wherever a range of hyperaparameters was explored, the validation loss on the total objective function was used as the model selection criterion (i.e., not the causal effect estimation performance, which is not available in real-world scenarios).

A.6 SOFTWARE AND HARDWARE

The network is coded using Pyro (Bingham et al., 2019) and is an extension of the code by (Zhang et al., 2020) available here: https://github.com/WeijiaZhang24/TEDVAE. We train on a GPU (e.g. NVIDIA 2080Ti) driven by a 3.6GHz Intel I9-9900K CPU running Ubuntu 18.04. Training 200 epochs of the IHDP dataset (training split of 450 samples) takes approx. 35 seconds (0.175s per epoch).