
A SURVEY OF DEEP CAUSAL MODELS AND THEIR INDUSTRIAL APPLICATIONS

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ABSTRACT

The concept of causality plays a significant role in human cognition. In the past few decades, causal effect estimation has been well developed in many fields, such as computer science, medicine, economics, and other industrial applications. With the advancement of deep learning, it has been increasingly applied in causal effect estimation against counterfactual data. Typically, deep causal models map the characteristics of covariates to a representation space and then design various objective functions to estimate counterfactual data unbiasedly. Different from the existing surveys on causal models in machine learning, this paper mainly focuses on the overview of the deep causal models, and its core contributions are as follows: 1) we summarize the popularly adopted relevant metrics under multiple treatment, continuous-dose treatment and times series treatment; 2) we cast insight on a comprehensive overview of deep causal models from both timeline of development and method classification perspectives; 3) we outline some typical applications of causal effect estimation to industry; 4) we also endeavor to present a detailed categorization and analysis on relevant datasets, source codes and experiments.

1 Introduction

In general, causality refers to the connection between an effect and the cause of it. Causes and effects of this phenomenon are difficult to define, and we are often only aware of them intuitively[1]. causal effect estimation is a process of drawing a conclusion about a causal connection based on the circumstances surrounding the occurrence of the effect and has a variety of applications in real-world scenarios[2]. For example, causal effect estimation of observational data in advertising[3, 4, 5, 6, 7, 8, 9], developing recommender systems that are highly correlated with causal treatment effect estimates[10, 11, 12, 13, 14, 15, 16], learning optimal treatment rules for patients in medicine[17, 18, 19], estimation of ITE in reinforcement learning[20, 9, 21, 22, 23, 24, 25, 26, 27], causal effect estimation tasks in natural language processing[28, 29, 30, 31, 32, 33], emerging computer vision and language interaction tasks[34, 35, 36, 37, 38], education[39], and strategy resolutions[40, 41, 42, 43, 44], etc.

Deep learning largely contributes to the development of artificial intelligence when applied to big data[45, 46, 47, 48]. In comparison with traditional machine learning, deep learning models are more computationally efficient, more accurate, and hold marvelous performance in various fields. However, many deep learning models are black boxes with poor interpretability since they are more interested in correlations than causality as inputs and outputs[49, 50, 51]. In recent years, deep learning models have been widely used for mining data for causality rather than correlation[40, 42]. Thus, deep causal models have become a core application for the issue of causal effect estimation in observation data with partial samples [19, 43, 44, 52]. At present, many works in the field of causal effect estimation utilize deep causal models to select reasonable treatment options[53, 54, 55, 56].

With the emergence of big data, all trend variables are tended to be correlated[57], so discovering the latent causal relationships is becoming one open problem[58, 59, 60]. In terms of statistical theory, it is the most effective way to

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conduct **randomized controlled trials(RCT)** to infer causality. In other words, the sample is randomly assigned to a treatment or control group. Despite this, real-world RCT data are sparse and have several serious deficiencies. The studies involving RCTs require a large number of samples with little variation in characteristics, which is difficult to interpret and will further involve some ethical issues inevitably. As a matter of fact, it is also not wise to select subjects to try a drug or vaccine in the process of drug development. Therefore, causal effect estimation issues are commonly measured directly using observational data. A central question for obtaining counterfactual results is how to deal with observational data[61]. When observational data are analyzed, treatments are not randomly assigned and the performance of the post-treatment samples are significantly different from that of the general samples[40, 42]. Unfortunately, we can't observe alternative outcomes in theory since the counterfactual results[62] are not available to be observed. For a long time, most of researchers have mainly explored the use of the Potential Outcome Framework as a means of addressing the problem of causal effect estimation from observational data[63]. The Potential Outcome Framework is also known as the Rubin Causal Model[64]. In essence, causal effect estimation is closely connected to deep learning since it is conceptualized using Rubin Causal Model. In order to enhance the accuracy and unbiasedness of estimates, there are some works that attempt to combine deep networks and causal models. Just to illustrate several of them, e.g., the methods considering representations of distribution balance[40, 42, 43], exploiting the effects of covariates confounding learning[52, 65, 66, 67], the methods based on generative adversarial networks[44, 68, 69, 70], and so forth[56, 33, 71].Due to the simultaneous development of deep learning and causal effect estimation, the problem of deep causal models has become more open and diverse[72, 73, 74].

In recent years, various perspectives have been discussed regarding causal effect estimation[75, 1, 76, 77, 78, 79, 80, 81, 2, 82, 83]. In Table 1, we list some of existing representative surveys and their highlight points. An in-depth analysis of the origin and development of causal effect estimation was provided in review[75], as well as the implications of causal learning for the development of causal effect estimation. Immediately after that, due to the rapid development of the field of machine learning, a detailed discussion of the relevance of graphical causal effect estimation to machine learning was presented in survey[76]. In addition, an overview of traditional and cutting-edge causal effect estimation methods and the comparison between machine learning and causal learning can be found in survey[1]. As one of hot focuses in recent years, the studies on the interpretability of machine learning have received great attention. An analytical summary of relevant interpretable artificial intelligence algorithms was elaborated in [77] by combining causal effect estimation with machine learning. Moreover, with the flourishing of causal representation learning, review[78] takes this new perspective to uncover high-level causal variables from low-level observations, strengthening the link between machine learning and causal effect estimation. In survey[82], the structural causal model of counterfactual intervention was comprehensively explained and summarized, and five classes of problems under causal machine learning are systematically analyzed and compared. Furthermore, in review[79], the authors discussed the way in which the latest progress of machine learning is applied to causal effect estimation, and provides a in-depth interpretation of how causal machine learning can contribute to the advancement in healthcare and precision medicine. As argued in review[80], causal discovery methods can be improved and sorted out based on deep learning, and can also be considered and explored from a variable paradigm perspective. Causal effect estimation in recommender systems is the focus of survey[81], which explains how causal effect estimation can be used to extract causal relationships to enhance recommender systems. The Potential Outcome Framework in statistics has long served as a bridge between causal effect estimation and deep learning. As a starting point, review[2] analyzes and compares different classes of traditional statistical algorithms and machine learning that satisfy these assumptions. Recently, survey[83] provides an overview of causal learning from Structural causal model, Potential outcome model, Deep Neural Network and Deep causal discovery, explores the internal mechanism of deep learning to improve the significance of causal learning. Different from the above review, we penetrate into the territory of biased sample observation of causal effect estimation, provide a systematic and comprehensive overview of the deep model in representation learning, debias estimation, counterfactual inference and other perspectives. It is worth mentioning that we provide a comprehensive conclusion of the latest applications of deep causal models in industry. This survey makes four core contributions: 1) we summarize the popularly adopted relevant metrics under multiple treatment, continuous-dose treatment and time series treatment; 2) we cast insight on a comprehensive overview of deep causal models from both temporal development and method classification perspectives; 3) we outline some typical applications of causal effect estimation to industry; 4) we also try to present a detailed categorization and analysis on relevant datasets, source codes and experiments.

The rest of the paper is outlined as follows. In Section 2, some relevant definitions and assumptions of causal effect estimation are inducted. Then, in Section 3, we present the classical examples and metrics, including binary treatment, multiple treatment, and continuous dose treatment. The deep causal model is comprehensively elaborated in Section 4. Next, we categorize the deep causal modeling methods into five groups in Section 5, including representation learning for distribution balancing, covariates confounding learning, methods based on generative adversarial networks, time series causal estimation problem, and methods based on multi-treatment and continuous-dose treatment models. Additionally, in Section 6, We summarize the industrial applications of causal reference. After that, the relevant experimental guidelines are listed in Section 7. Finally, we conclude this article in Section 8.

Table 1: **Highlights of existing surveys on causal learning**

Surveys	Highlights
The Development of Causal Reasoning[75]	The origin and development of causal reasoning
Causality for Machine Learning[76]	The connection between graphical causal effect estimation and machine learning
Causal Inference[77]	Machine Learning Interpretability for Counterfactual Causal Reasoning
Toward Causal Representation Learning[78]	Exploring causal variables in data through causal representation learning
Causal Machine Learning for Healthcare and Precision Medicine[79]	Causal Machine Learning in clinical decision support systems
A Survey of Learning Causality with Data: Problems and Methods[1]	The relationship between causal learning and machine learning in big data
A Survey on Causal Inference[2]	Causal effect estimation of observational data in the potential outcomes framework
A Review and Roadmap of Deep Learning Causal Discovery in Different Variable Paradigms[80]	An improved exploration of causal discovery from the perspective of deep learning and variable paradigms
Causal Machine Learning: A Survey and Open Problems[82]	Five types of causal machine learning problems under structural causal models
Causal Inference in Recommender Systems: A Survey and Future Directions[81]	Optimize the recommendation system by extracting causality through causal reasoning
Deep Causal Learning: Representation, Discovery and Inference[83]	The significance of deep learning for solving traditional causal learning problems
A Survey of Deep Causal Models and Their Industrial Applications	Causal effect estimation in deep models with biased sample observations and challenges in current industrial applications

2 Preliminaries

In this section, we present the background knowledge of causal effect estimation, including task descriptions, mathematical concepts, and pertinent assumptions.

Basically, the aim of causal effect estimation is to estimate the change in outcome that will occur if a different treatment are implemented[2]. Imagine that there are several treatment plans A, B, C, and so on, all of which have different cure rates, and the change in the cure rate is the result of the treatment scheme. Realistically, we can't apply different treatment regimens to the same group at the same time. As opposed to RCT, the main problem to be solved in observational research is the lack of counterfactual data. In other words, the challenge we face is how to find the most effective treatment plan based on past experimental diagnosis and medical history of the patient.

2.1 Definitions

Here, to make the survey self-consistent, we first give some basic definitions under the Potential Outcome Framework[64]. In particular, causality is defined as the result of a treatment scheme applied to a sample, which can either be a specific behavior, a specific method, or some specific treatment scheme. The followings are the concepts related to causal effect estimation, which are benchmarked against the relevant basic definitions in the survey[2].

Definition 1 *Treatment Effect Estimation(Causal Effect Estimation):* Estimate the change in outcome of a given sample receiving an intervention.

Treatment effect estimation and lift modelling have the same goal, the difference is: the lift model is on random experimental data, treatment effect estimates usually need to adhere to the necessary assumptions, applied to experimental and observational data.

Definition 2 *Treatment:* A treatment refers to a scheme or action applied to a sample.

As a medical term, a drug scheme is a treatment. For binary treatments, $T = 1$ is the *treated group*, and $T = 0$ is the *control group*. Multiple treatment can be indicated by the $T \in \{0, 1, 2, \dots, T_N\}$, where $N + 1$ denotes the total number of treatments.

Definition 3 *Observed outcome:* An observational outcome, also known as a *factual outcome*, is a measure of how the sample's outcomes applied to the treatment.

In the case of a specific treatment, observed outcomes can be displayed in Y^F , where $Y^F = Y(T = T_i)$. A donation of in the amount of Y_{T_i} is made as *potential outcome*[84].

Definition 4 *Counterfactual outcome:* A counterfactual outcome is the outcome that differ from a factual outcome.

With binary treatments, counterfactual outcome is denoted as Y^{CF} , and $Y^{CF} = Y(T = 1 - T_i)$. Assuming multiple treatment, let $Y^{CF}(T = T'_i)$ donate the counterfactual result of treatment T'_i .

Definition 5 *Dose:* A dose refers to the amount of drug taken continuously during a particular course of treatment.

In general, there are many medical treatments involving continuous dose parameters, such as vasopressors. A set of continuous dose regimens can be donated as D_T , the treatment-specific factual doses can be donated as can be donated as D^F , and $D^F = D(T = T_i)$. Simultaneously, counterfactual dose can be donated as $D^{CF}(T = T'_i)$.

Definition 6 *Dose-response curve:* A dose-response curve indicates the effect of the samples' response over time after receiving different doses of the intervention.

A better fit to the dose-response curve could make the model more robust and expressive in continuous dose treatments. The actual and counterfactual outcomes on the dose-response curves can be expressed as the sets $Y^F(D^F, T_i)$ and $Y^{CF}(D^{CF}, T_i')$.

Definition 7 *Covariates:* Covariates are variables that are unaffected by treatment choice.

Generally, covariates in the healthcare setting refer to the patient's demographic, medical history, experimental data, etc., usually denoted by X . Covariates can be separated into confounding and non-confounding variables, specifically divided into three categories[65]: instrumental factor I , which only affects treatment T ; confounding factor C , which contributes to both treatment T and outcome Y ; and adjustment factor A , which only determines outcome Y .

2.2 Assumptions

After understanding the basic definition of causal effect estimation, the following three assumptions, which are derived from papers[2, 77], are commonly required to achieve the estimation of causal treatment effects.

Assumption 1 *Stable Sample Treatment Value (SSTV):* One sample's response to treatment is independent of the assignment of other samples.

Based on this assumption, there is no interaction between samples, and there is only one version of each treatment option. SSTV assumption can be expressed as $P(Y_i|T_i, T_i', X_i) = P(Y_i|T_i, X_i)$.

Assumption 2 *Ignorability:* Given the covariate X , the treatment distribution T is independent of the potential outcomes.

On the assumption of ignorability, there should be no unobserved confounding factors. In other words, $T \perp Y(T = T_i), Y(T = T_i')|X$ needs to be satisfied.

Assumption 3 *Overlap:* When given the observed variables, each sample has nonzero probability to receive either treatment status.

In order to estimate the counterfactual treatment effect, it must be assumed that each sample can implement any treatment option, otherwise the overlap assumption will not be valid. That is, $0 < P(T = T_i|X = x) < 1$ and $0 < P(T = T_i'|X = x) < 1$

3 Treatments and Metrics

This section provides an analysis and description of the different performance metrics adopted for different classical application scenarios. In addition to the basic metrics in survey[2], we expand the evaluation from binary to multiple and continuous dose cases.

3.1 Binary treatment

Treatment refers to the action of a sample or a subject. In the medical field, a medication regimen for a patient is a treatment. When there are only two treatment options, the group of units applied with treatment $T = 1$ is the treated group, and the group of units with $T = 0$ is the control group, which can be referred to as the binary treatment[2].

In the binary treatment situation, the most basic and common performance metric is **Average Treatment Effect(ATE)**, which is defined as[85]:

$$ATE = \mathbb{E}[Y(T = 1) - Y(T = 0)], \quad (1)$$

where $Y(T = 1)$ and $Y(T = 0)$ indicate the results of the treatment and control groups in the population. This metric is commonly used to estimate causal effects for the well-known dataset IHDP[86]. Differentiate between treatment and control groups by whether intensive high-quality child care is applied, and the outcome is a score on the cognitive test for the infant.

In a sample set, the treatment effect is called the **Conditional Average Treatment Effect (CATE)** that is given as follows[2]:

$$\text{CATE} = \mathbb{E}[Y(T = 1)|X = n] - \mathbb{E}[Y(T = 0)|X = n], \quad (2)$$

where $Y(T = 1)|X = n$ and $Y(T = 0)|X = n$ represent the results under the sample set for the treatment and control groups with $X = n$, respectively. Since different treatments have different effects on different example sets, CATE is also known as heterogeneous treatment effect. The treatment effect at the individual level is called **Individual Treatment Effect (ITE)**, which is defined as[2]:

$$\text{ITE}_n = Y_n(T = 1) - Y_n(T = 0), \quad (3)$$

where $Y_n(T = 1)$ and $Y_n(T = 0)$ represent the results of the sample in the treatment and control groups.

It is also helpful to keep in mind another evaluation metric, which is called **Precision in Estimation of Heterogeneous (PEHE)**. Regardless of fact or counterfactual outcomes, PEHE, as defined below[44], can perform unbiased estimates:

$$\text{PEHE} = \sqrt{\frac{1}{N} \sum_{n=1}^N (Y_1^F(n) - Y_0^F(n) - (Y_1^{CF}(n) - Y_0^{CF}(n)))^2} \quad (4)$$

where $Y_1^F(n)$, $Y_0^F(n)$ and $Y_1^{CF}(n)$, $Y_0^{CF}(n)$ respectively indicate unbiased estimates of the facts and the counterfactuals for the treatment and control groups. The Twins[52] dataset is typical of the application of this metric, and the result is a one-year mortality rate for children.

For the treated group, the treatment effect is referred to as the **Average Treatment effect on the Treated group (ATT)**, which is defined as[2]:

$$\text{ATT} = \mathbb{E}[Y(T = 1)|T = 1] - \mathbb{E}[Y(T = 0)|T = 1], \quad (5)$$

where $Y(T = 1)|T = 1$ and $Y(T = 0)|T = 1$ correspond to potential treated and control outcome of the treated group respectively. This metric is available for the jobs dataset[87], treatment group take part in vocational training, while the control group are not and the outcome is employment status.

Since only factual data is available for Jobs dataset, the testing set is from RCT. Therefore, heterogeneity effect indicators only can be described by **Policy Risk** ($\mathcal{R}_{\text{pol}}(\pi)$), which is defined as[42]:

$$\mathcal{R}_{\text{pol}}(\pi) = \frac{1}{N} \sum_{n=1}^N \left[1 - \left(\sum_{i=1}^K \left[\frac{1}{|\Pi_i \cap T_i \cap E|} \sum_{X(n) \in \Pi_i \cap T_i \cap E} Y_i^F(n) \times \frac{|\Pi_i \cap E|}{|E|} \right] \right) \right] \quad (6)$$

where $\Pi_i = \{X(n) : i = \arg \max Y^{\hat{CF}}\}$, $T_i = \{X(n) : t_i(n) = 1\}$, and E is the subset of RCT.

3.2 Multiple treatment

Unlike the binary treatment problem, the multiple treatment problem can be described by the following stereotype[41]: For each sample, the potential outcomes are represented as a vector Y with N entries Y_j^F where each entry corresponds to the outcome when applying one treatment T_j out of the set of N available treatments $T = \{0, 1, 2, \dots, T_N\}$, with $j \in \{0, 1, 2, \dots, N - 1\}$. The set of available treatments can contain more than two treatments. As training data, samples k and their observed factual outcomes Y^F are received when applying one treatment T_i , the other outcomes can not be observed. The purpose is to train a predictive model \hat{f} that is able to estimate the entire potential outcomes vector \hat{Y} with k entries Y_j .

With the exception of ATE and CATE, an accurate estimate of treatment effect can be determined by using **Root Mean Square Error (RMSE)** for all subgroups, which is defined as[88]:

$$\text{RMSE} = \sqrt{\frac{1}{N} \sum_{n=1}^N \frac{1}{|T|} \sum_{j \in T} (Y^F(n, j) - Y^{CF}(n, j))^2} \quad (7)$$

where T is the set of all possible combinations of treatments, N is the number of observations. $Y^F(n, j)$ displays the true outcome when the j^{th} subset is applied to compute the n^{th} observation, whereas $Y^{CF}(n, j)$ displays the predicted outcome when the j^{th} subset is applied to compute the n^{th} observation. The **Absolute Error** can be calculated in the same way. This metric is commonly used for the News[41] dataset, which the treatments are the choice of viewing tools, such as smartphones, tablets, desktops, and TVs.

To measure the average of the RMSE between the actual and estimated difference between ITE for each treatment with no treatment ITE, We can incorporate multiple treatment into one calculation criterion, which is called **Average PEHE**[88]:

$$\text{AveragePEHE}_j = \frac{1}{|T|} \sqrt{\frac{1}{N} \sum_{n=1}^N ((Y^F(n, j) - Y^F(n, 0)) - (Y^{CF}(n, j) - Y^{CF}(n, 0)))^2}, j \in (T - T_0) \quad (8)$$

where T_0 represents the sample set with no treatment applied. It is worth mentioning that the latest research on estimating the effect of multi-causal treatment COVID-19[89, 90]. Average PEHE typically serve as performance metrics in dataset CHESS.

3.3 Continuous dose treatment

Vary from binary and multivariate treatment issues, the continuous-dose treatment issue can be formulated as follows[69]: Consider to receive observations of the form $(\mathbf{x}^i, T_f^i, Y_f^i)$ for $i = 1, \dots, N$, where, for each i , these are independent realizations of the random variables (\mathbf{X}, T_f, Y_f) , refer to \mathbf{X} as the feature vector lying in some feature space \mathcal{X} , containing pre-treatment covariates. The treatment random variable, T_f , is in fact a pair of values $T_f = (W_f, D_f)$ where $W_f \in \mathcal{W}$ corresponds to the *type* of treatment being administered which lies in the discrete space of N treatments, $\mathcal{W} = \{w_1, \dots, w_k\}$, and D_f corresponds to the *dosage* of the treatment, which, for a given treatment w lies in the corresponding treatment's dosage space, \mathcal{D}_w (e.g. the interval $[0, 1]$). The set of all treatment-dosage pairs can be defined as $\mathcal{T} = \{(w, d) : w \in \mathcal{W}, d \in \mathcal{D}_w\}$.

In continuous dose treatment situation, the sample dose-response curve is adopted as the metric. Furthermore, the metric on the test set is different[55]. Therefore, we can use **Mean Integral Squared Error (MISE)** to measure the accuracy of the estimation of the patient treatment effect on the dose space, which is defined as[55].

$$\text{MISE} = \sqrt{\frac{1}{K} \frac{1}{N} \sum_{T_n \in \mathcal{T}} \sum_{n=1}^N \int_{\mathcal{D}_{T_n}} (Y_n(T_n, u) - \hat{Y}_n(T_n, u))^2 du} \quad (9)$$

where T is the set of treatments in the space, K is the number of samples, and u is the treatment dose taken. For a given treatment, T_n lies within the dose space \mathcal{D}_{T_n} . Moreover, $Y_n(T_n, u)$ and $\hat{Y}_n(T_n, u)$ denote the results predicted by the model and the results at the optimal treatment dose, respectively.

In addition, the **Mean Dose Policy Error (DPE)** is another marvelous metric of a model's ability to predict the optimal dose point for each individual treatment, which can be defined by [55]:

$$\text{DPE} = \sqrt{\frac{1}{K} \frac{1}{N} \sum_{T_n \in \mathcal{T}} \sum_{n=1}^N (Y_n(T_n, D_{T_n}^*) - Y_n(T_n, \hat{D}_{T_n}^*))^2} \quad (10)$$

where, $D_{T_n}^*$ and $\hat{D}_{T_n}^*$ represent the true optimal dose and the model-determined optimal dose under a treatment, respectively. With **Sequential Least Squares Estimation**[91], the optimal dose point for the model can be determined.

In order to compare the optimal treatment dose pair selected by the model with the true optimal treatment dose pair, the **mean policy error (PE)** is given as[55]:

$$\text{PE} = \sqrt{\frac{1}{N} \sum_{n=1}^N (Y_n(T_n^*, D_{T_n}^*) - Y_n(\hat{T}_n^*, \hat{D}_{T_n}^*))^2} \quad (11)$$

where T_n^* and \hat{T}_n^* represent the optimal treatment and the optimal treatment determined by the model, respectively. By calculating the optimal dose for each treatment and then selecting the treatment that yields that optimal dose, the optimal dose pair for the model can be selected. The above metrics are typically used in the TCGA[92] dataset. Among them, drug therapy, chemotherapy, and surgery are the treatment options, and the outcome is a risk of cancer recurrence after treatment.

3.4 Time series treatment

Different from the continuous-dose treatment, the time series treatment issues adopts historical data for causal effect estimation under the time step. Let \mathbf{x}_t be the time-dependent covariates of the observational data at time stamp t such

that $\mathbf{x}_t = \{x_t^{(1)}, x_t^{(2)}, \dots, x_t^{(n)}\}$, where $x_t^{(i)}$ denotes the covariates for i -th individual, n denotes the number of samples. The static features, represented as $c^{(i)}$ (e.g., demographic information), do not change overtime are also considered as observed covariates. At each time stamp t , the treatment assignments are denoted as $\mathbf{A}_t = \{a_t^{(1)}, a_t^{(2)}, \dots, a_t^{(n)}\}$. $a_t^{(i)}$ denotes the treatments assigned to i -th individual at time stamp t . In the case of the binary treatment setting, i.e., $a_t^{(i)} \in \{0, 1\}$, where 1 is considered as treated while 0 as control, time series treatment effect estimation is interested in estimating the effect of the treatment assigned until time stamp T on the outcomes $Y_{T+\tau}$, observed at time stamp $T + \tau$, where τ is the prediction window. We should note that, in observational data, an individual can only belong to one group (i.e., either treated or control group), thus the outcome from the other group is always missing and referred to counterfactual. Combining all covariates and treatments, the observational data for the i -th individual can be represented as $D^{(i)} = \{x_t^{(i)}, a_t^{(i)}\}_{t=1}^T \cup \{c^{(i)}\} \cup \{y_t^{(i)}\}_{t=T+1}^{T+\tau}$. Then the individual treatment effect on the time series observational data is defined as[93]:

$$E_t^{(i)} = \mathbb{E}[y_{1,t}^{(i)} | x_t^{(i)}, a_t^{(i)}, D^{(i)}] - \mathbb{E}[y_{0,t}^{(i)} | x_t^{(i)}, a_t^{(i)}, D^{(i)}], t \in [T + 1, T + \tau] \quad (12)$$

Similar to binary and multiple treatment, **PEHE** or **Average PEHE** can be used as evaluation metrics depending on the type of treatment for ITE. While **RMSE** or **MAE** can be used for ATE and CATE. MIMIC III [94] dataset is commonly used for times series treatment issues, which is a database of electronic health records from ICU patients.

Additionally, many researchers have flexibly applied simulation datasets to differ scenarios. In order to provide more ablation experiments and prove the robustness of the model. A detailed description of the relevant datasets can be found in Section 7.

4 Development of Deep Causal Models

In recent years, there is growing dissatisfaction within industry with current machine learning algorithms, 87% of machine learning projects never make it beyond an experimental phase into production, according to Forbes. However, in the context of big data, with the development of deep learning, the decision-making ability of causal effect estimation has been significantly improved. A recent global survey found that 83% of Ctos and senior data scientists agree that causal deep learning can lead to a more robust business environment model, and 60% of senior data scientists intend to make a significant investment in deep causal models.

With a solid understanding of the basic definitions and model measures of causal effect estimation, this section moves to the core of the paper. We provide an overview of deep causal models during the last several years and a detailed classification of them.

4.1 A timeline of development

In the past few years, research on deep causal models has advanced considerably, and they have become more accurate and efficient in estimating causal effects. In Figure1, we present the development timeline about 40 classical deep causal models from June 2016 to February 2022. It also displays categories of models, the relationship and citation volumes between essays.

Deep causal models have emerged since 2016. For the first time, Johansson et al. publish **Learning Representations for Counterfactual Inference**[40], and propose the algorithm framework BNN and BLR[40], in which deep learning is combined with the causal effect estimation, and the causal effect estimation problem is transformed into a domain adaptation problem. Since then, a number of models, including DCN-PD[95], TARNet and CFRNet[42], have been proposed since then. It is important to note that the CEVAE[52] model proposed by Louizos et al. in December 2017, which is based on classical structural Variational Autoencoders, focuses on the impact of confounding factors on the estimation of causal effects.

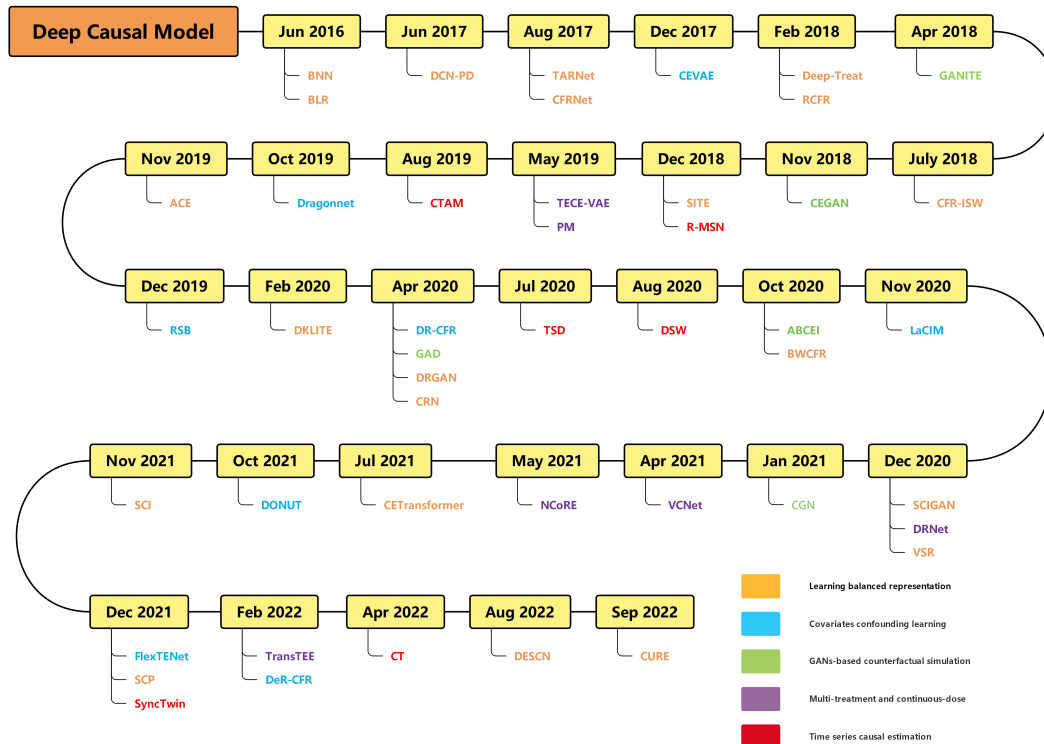


Figure 1: Overview of development timeline of classical deep causal models

In 2018, and going forward into 2019, there emerged an increasing interest in causal representation learning with representative works including Deep-Treat[19] and RCFR[96] models. After the launch of the GANITE[44] model, the use of generative adversarial model architecture for counterfactual estimation becomes mainstream in the field of causal effect estimation. In accordance with the previous works, some novel optimization ideas were proposed in CFR-ISW[97], CEGAN[68], SITE[43].

By applying recurrent neural networks, the R-MSN[71] model aims to solve the problem of continuous dose of multi-treatment time series, which opened up a new theory of deep causal models. Furthermore, PM[41] and TECE-VAE[88], proposed in 2019, attempt to address the issue of estimating causal effects associated with multiple discrete treatment. As a follow-up, the CTAM[33] begins to focus on estimating causal effects for textual data; the Dragonnet[66] adhibits regularizations and propensity score networks into causal models for the first time; the ACE[53] attempts to extract fine-grained similarity information from representation space. After that, RSB[98] model adopts deep representation learning network and PCC regularization for covariate decomposition, uses instrumental variables to control selection bias, and uses confounding factors and moderating factors to predict results.

In 2020, deep causal models got a rapid boom. For DKLITE[54] model, it effectively combines the deep kernel trick and posterior variance regularization. Then, DR-CFR[99] was proposed to decouple selection bias for covariates; GAD[100] focuses on the causal effect of continuous dose treatment; DRGAN[101] defines an innovative generative adversarial network for fitting sample dose effect curves; and CRN[102] estimates time-varying treatment effects by combining counterfactual recurrent neural networks. Following estimating time series causal effects under multi-cause confounding, TSD[103] turns to the estimation of time series causal effects. In the aspect of learning the latent representation space, ABCEI[104] achieves a marvelous balance between the covariate distribution of treatment and the control groups by using GAN. As two variants of previous works[42, 52], both BWCFR[105] and LaCIM[106] are further optimized for the model structure. Additionally, in 2020, SCIGAN[69] and DRNet[55] extended the task of continuous dose treatment to arbitrary quantity, while VSR[107] aggregated the deep latent variables in a reweighted manner.

Up to now since 2021, deep causal models have become more innovative, open, and flexible. The VCNet[56] model implemented an estimator of continuous mean dose-response curves. Moreover, CGN[70] proposed more robust and interpretable classifiers that explicitly expose the causal structure of tasks. As for NCoRE[108], it used cross-treatment interaction models to explain the process of multi-treatment combination at the potential causal level. After that,

inspired by the success application of Transformer in many fields, CETransformer[109] was proposed to characterize the covariates by focusing the attention on the correlation between the covariates. In addition, DONUT[110] and DeR-CFR[65] are optimized for the two previous works[66, 99] respectively. In [72], the subspace methods was further explored for causal representation learning, which formulate the SCI model. From the perspective of multi-task learning e, a multi-task adaptive learning architecture was proposed by FlexTENet[111], in which CATE estimation architecture adaptively learns what to share between the PO functions. Aside from that, SCP[112] attempted to estimate the multifactorial treatment effects using a two-step procedure. To construct synthetic twin matching representation, SyncTwin[113] utilized the temporal structure in the potential outcomes.

In 2022, TransTEE[73] extended the distribution balance approach for spatial representations to continuous, structured, and dose-dependent treatments, applying the latest theory of deep learning to expand the depth and breadth of causal effect estimation. Furthermore, DESCN[114] obtained comprehensive information on treatment propensity, response and hidden treatment effects through a crossover network by means of a multi-task learning. Beside that, CT[115] combined Transformer with the LSTM network to capture complex long-term dependencies between time-varying confounders. Until recently, CURE[116] pre-trained large-scale unlabeled patient data to learn representative background patient representations, and then fine-tunes the labeled patient data for treatment effect estimation.

4.2 Model classification

Through sorting out the existing models, we can find that the current deep causal models are mainly studied from the following aspects that include : 1) Learning balanced representations; 2) Covariate confounding learning; 3) Time series causal learning; 4) GANs based Counterfactual simulation; and 5) Multi-treatment and continuous-dose treatment. In Figure 2, we present a detailed classification of the current deep causal model.

- **Learning balanced representations:** This type of approach has long been a popular research. The core ideology is to use the encoder to map the covariates X to the representation space Φ , combine the processing T , adopt the network h to predict the output outcome Y , and minimize the distribution distance $disc_{\mathcal{H}}$ between the factual and counterfactual outcomes. Classical architectures include BNN[40], CFRNet[42], SITE[43], ACE[53], DKLITE[54], SCI[72], etc.
- **Covariate confounding learning:** This type of approach aims to decomposition of covariant relation in theory. Its main application schemes are unbiased estimation of covariates and removal of confounding factors using decoupling, reweighting, codec reconstruction, etc. The typical structures include CEVAE[52], Dragonnet[66], DeR-CFR[65], LaCIM[106], DONUT[110], FlexTENet[111], and so on.
- **GANs-based counterfactual simulation:** With the great success of GANs in data synthesis in recent years, it is also widely adopted to solve causal effect estimation problems. Two schemes are usually involved in Using GAN networks for counterfactual simulation, i.e., generating the counterfactual output outcomes or balanced representation space distributions. Such classical frameworks include GANITE[44], CEGAN[68], ABCEI[104], CETrnaformer[109], etc.
- **Time series causal estimation:** Temporal causal estimation has been widely concerned. Using RNNs to track contextual covariate information and handle time-varying confounding bias is a long-standing solution adopted by many models. Some typical architectures are R-MSE[71], CTAM[33], CRN[102], TSD[103], and so on.
- **Multi-treatment and continuous-dose treatment:** The issues of Multiple treatment and continuous-dose treatment are one of recent research hotspots in deep causal learning. In general, such issues can be further simplified and structured using schemes such as matching, variational autoencoders, hierarchical discriminators, and multi-headed attention mechanisms. The classical models include PM[41], TECE-VAE[88], DRNet[55], SCIGAN[69], VCNet[56], TransTEE[73], etc.

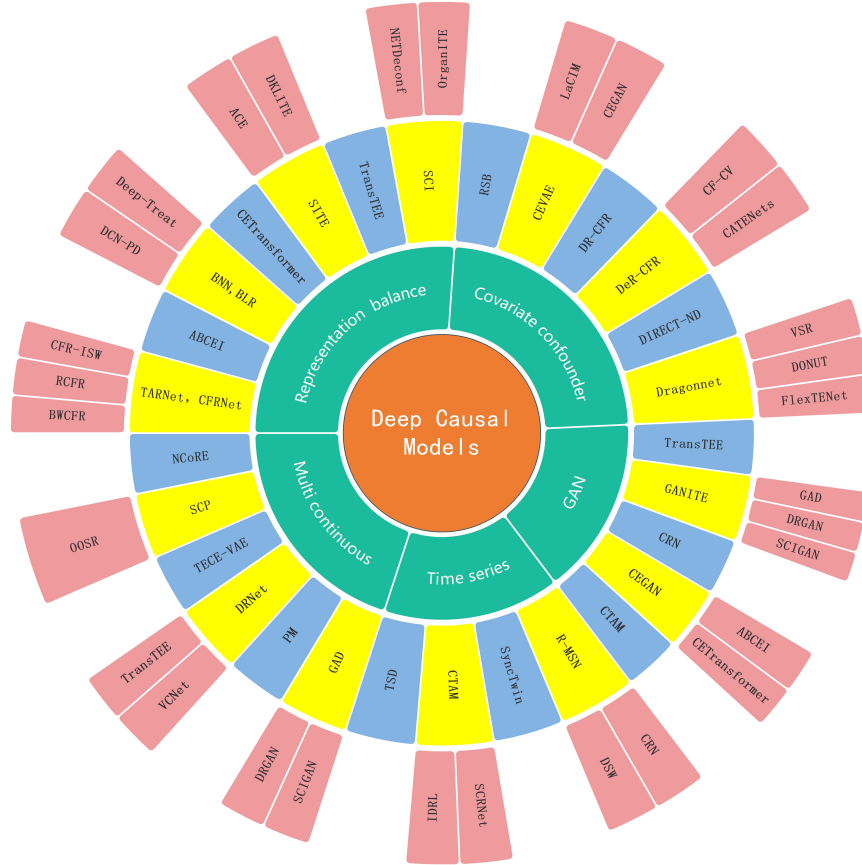


Figure 2: **Taxonomy of deep causal models**

It is worth pointing out, as different application scenarios arise, new research focuses and methods in the aspect of deep causality model will also emerge continuously. In the following section, we will give a detailed introductions to the typical models according to the taxonomy of deep causal models motioned above.

5 Typical Deep Causal Models

As more and more data accumulate in the fields of healthcare, education, economy, etc., deep learning is increasingly being used to infer causal relationships from counterfactual data. As opposed to existing deep causal models, which typically map covariates to a representation space, unbiased estimation of counterfactual data can be achieved by the objective function. Unlike the brief overview of the various classical models that are categorized from the different research perspectives for deep causal models, Table 2 summarizes the classical network architecture applied by those typical deep causal models. In addition, in the following detailed descriptions on the typical deep learning-based causal models, the issues and challenges that these models face are also discussed.

5.1 Learning balanced representation

Most statistical learning theories posit that the test data and training data have independent and identical distributions, but in reality, the distributions of test data and training data are often related, but not identical. Solving this problem requires deep learning models that learn causality rather than correlation in the field of causal effect estimation. There is no standard treatment assignment strategy for observational data, unlike RCTs. As we know, fact and counterfactual distributions are often different because of selection bias caused by known and unknown covariates. Hence, causal effect estimation needs to be transformed into a domain adaptation problem to predict counterfactual outcomes by learning from factual data.

Table 2: **Highlights of deep frameworks on classical causal models**

Models	GAN	AE	RNN	Transformer
DCN-PD[95]				
BNN[40]		✓		
CFRNet[42]		✓		
CEVAE[52]		✓		
Deep-Treat[19]		✓		
GANITE[44]	✓			
CEGAN[68]	✓			
SITE[43]		✓		
R-MSN[71]			✓	
PM[41]		✓		
TECE-VAE[88]		✓		
CTAM[33]	✓	✓		
Dragonnet[66]		✓		
ACE[53]		✓		
RSB[98]		✓		
DKLITE[54]		✓		
GAD[100]	✓			
CRN[102]	✓		✓	
TSD[103]			✓	
ABCEI[104]	✓	✓		
BWCFR[105]		✓		
LaCIM[106]		✓		
SCIGAN[69]	✓	✓		
DRNet[55]		✓		
VSR[107]		✓		
VCNet[56]		✓		
NCoRE[108]		✓		
CETransformer[109]	✓	✓		✓
DeR-CFR[65]		✓		
SCI[72]		✓		
WUNT[117]				✓
FlexTENet[111]		✓		
SCP[112]		✓		
CGN[70]	✓	✓		
SyncTwin[113]			✓	
TransTEE[73]	✓	✓		✓
DESCN[114]		✓		
CURE[116]		✓		✓
DSW[93]		✓	✓	
CT[115]	✓	✓	✓	✓

For counterfactual results to be predicted, effective feature representations are necessary, especially the balanced distributions. According to Johansson et al, BNN[40] is an algorithmic framework as shown in Figure 3 for counterfactual reasoning that transforms the causal effect estimation problem into a representation distribution balance problem.

Upon mapping the covariates to the representation space, the encoder makes use of a two-layer fully connected neural network, balances the distribution distance of the representation space, and then derives the counterfactual results using another two-layer fully connected network. The used regression function is as follows:

$$B_{\mathcal{H},\alpha,\gamma}(\Phi, h) = \frac{1}{n} \sum_{i=1}^n |h(\Phi(x_i), t_i) - y_i^F| + \alpha \text{disc}_{\mathcal{H}}(\hat{P}_{\Phi}^F, \hat{P}_{\Phi}^{CF}) + \frac{\gamma}{n} \sum_{i=1}^n |h(\Phi(x_i), 1 - t_i) - y_{j(i)}^F| \quad (13)$$

where the encoder network is represented by Φ , a predictor network by h , and the metric function is $\text{disc}_{\mathcal{H}}$ that represents the distance between the two distributions. Aside that, Eq.(13) minimizes the error of the training set facts.

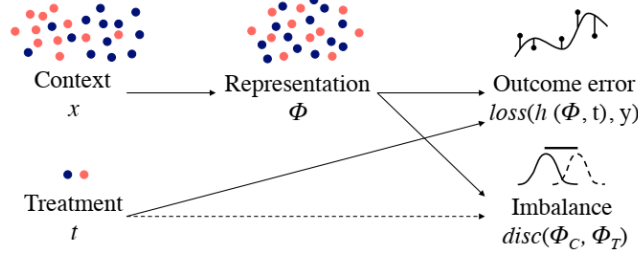


Figure 3: **Balancing Neural Network (BNN)** [40]: Counterfactual inference networks based on representation distribution balance

As an innovative method for measuring the spatial distribution distance between treatment groups and control groups, the literature[42] proposed a CFRNet network structure based on BNN[40] and adopted MMD and WASS for spatial distribution distance representations. When the network is trained, the imbalance penalty is calculated based on the explicit boundary of the distance, and the loss is calculated separately for the treatment group and the control group. As well as adding multiple neural network layers between results prediction layer, DCN-PD[95] combined multi-task deep neural networks with propensity score dropout[118].

On the basis of the CFRNet[42] model, both RCFR[96] and CFR-ISW[97] used the Propensity score to re-weight the representative spatial feature region and the sampling objective function; Atan et al. proposed an unbiased autoencoder network Deep-Treat[19] framework, applying a feedforward neural network to learn the optimal treatment strategy. While it reduces the loss of representation reconstruction as well as the information loss in space, the selection bias is also narrowed.

To maintain local similarity and balance of data representing the treatment and control groups and to improve individual treatment outcomes. Yao et al. proposed the SITE[43] method, which combines position-dependent depth metric PDDM with midpoint distance minimization MPDM into the representation space, and predicts the potential outcomes using a binary result network. In this case, the following loss function is used:

$$\mathcal{L} = \mathcal{L}_{FL} + \beta \mathcal{L}_{PDDM} + \gamma \mathcal{L}_{MPDM} + \lambda \|W\|_2 \quad (14)$$

where \mathcal{L}_{FL} is the loss between the predicted and observed factual outcomes, \mathcal{L}_{PDDM} and \mathcal{L}_{MPDM} are the loss functions of the PDDM and MPDM, respectively, and the last term is the L_2 regularization of the model parameter M .

According to SITE[43], ACE[53] proposed a balanced and adaptive similarity regularization structure to extract spatially fine-grained similarity information; in [54] DKLITE was proposed by applying the deep kernel regression and a posterior regularization to learn the spatial domain overlap information; and BWCFR[105] re-weighted the spatial feature distribution for the domain overlap region.

Unlike the above models, many other works attempt to combine representation distribution balancing with other deep network models. By combining GANs with a mutual information estimator regularization structure, ABCEI[104] tries to balance the covariate distributions of the treatment and control groups in the representation space; In [109], CETransformer was proposed to apply the attention mechanism to focus on the relationships between covariates and then learn a balanced representation distribution; As TransTEE[73] extends the balanced representation distribution method to Continuous, Structured, and Dose-Related treatments, it makes causal effect estimation a more open-ended problem; CURE[116] designed a new sequence encoding for longitudinal (or structured) patient data and merge structure and time into patient embeddings; DESCN[114] learned the processing and response functions jointly across the sample space to avoid processing bias, and used an intermediate pseudo-processing effect prediction network to mitigate sample imbalance; In [72], SCI inducted the concept of a subspace as shown in Figure 4, integrating the covariates into a common subspace, a treatment subspace, and a control subspace simultaneously, thereby obtaining a common representation and two specific representations. Afterwards, the common representation is connected to the specific representation of the treatment group and of the control group, and two potential results are obtained from the reconstruction and prediction network. Based on SCI, NETDECONF[119] used network structure information to infer hidden confounding in the observational data. A personalized treatment effect model that allocates treatments according to scarcity and estimates potential outcomes is proposed by OrganITE[120].

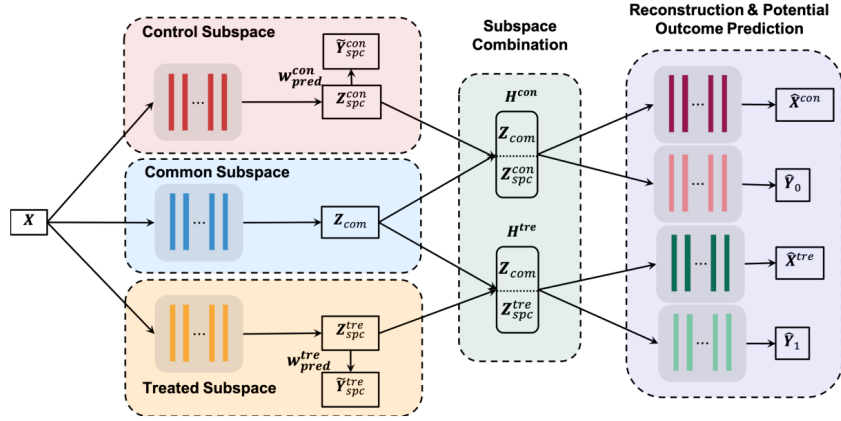


Figure 4: **Subspace Learning Based Counterfactual Inference network (SCI)** [72]: Estimating individual causal effects while preserving information about specific and common subspaces

Due to the improvement in the feasibility of estimating causal effect, the representation distribution balance is becoming the mainstream way, but it is still limited to estimating individual treatment effects, and thus is hard to expand to a broader range of applications like multi-treatments and continuous-dose treatments.

5.2 Covariates confounding learning

The main issue in causal effect estimation is estimating the treatment effect when given a covariate, a treatment, and a predicted outcome. By identifying and correcting for confounders, it is possible to estimate causal effects with greater accuracy from observational data. Nevertheless, in practical cases, there are potential confounders of noise and uncertainty, as well as some non-confounders. For this reason, mining potential confounders and decoupling the associated covariates are important methods to learn unbiased representations of counterfactuals from observed data.

A CEVAE[52] model structure was first proposed by Louizos et al. to capture hidden confounding factors with VAEs in the presence of noise and uncertain confounding factors, and perform treatment and prediction. On the basis of TARNet[42]’s causal relationship diagram structure, Do-calculus[121] derivations are performed for y and t in the inference network, respectively, and z and t in the model network to fit the interaction between potential confounding variables and treatment effects. Overall, the following prediction function is involved in the causal variational autoencoder:

$$\mathcal{F}_{\text{CEVAE}} = \mathcal{L} + \sum_{i=1}^N (\log q(t_i = t_i^* | \mathbf{x}_i^*) + \log q(y_i = y_i^* | \mathbf{x}_i^*, t_i^*)) \quad (15)$$

where, in the training set, the input, treatment, and outcome random variables are observed at points \mathbf{x}_i^* , t_i^* , and y_i^* , respectively.

On the ground of CEVAE[52], Sun et al. proposed the LaCIM[106] latent causal model to avoid false associations and improve the generalization ability of the model; CEGAN[68] used GAN networks to identify the potential confounders unbiasedly.

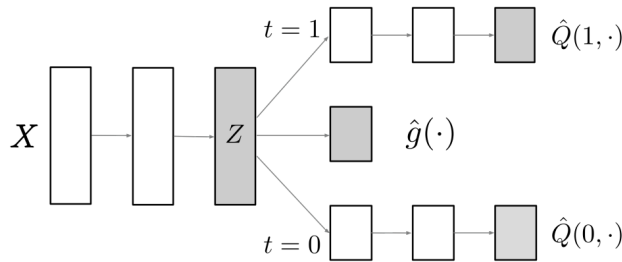


Figure 5: **Dragonnet**[66]: Propensity score adaptive neural network

Proposed by Shi et al., first-ever Dragonnet[66] adhibited the regularization objective functions into nonparametric estimation theory and the Propensity score prediction networks to CFRNet[42], thus making sure that the covariates are adjusted for treatment-related information in them. As can be seen, Figure 5 shows the network structure of the propensity score adaptive neural network. In accordance with Dragonnet[66], VSR[107] proposed a reweighting model that removes the association processing and confounding factors. It also used a deep neural network to aggregate the density ratios of latent variables across the variational distribution for calculateing the sample weight distribution; As part of the estimation process, DONUT[110] has added an orthogonal constraint to the non-confounding factors in the total loss; In addition, an end-to-end regularization and reparameterization method called FlexTENet[111] was proposed to learn a new architecture using multi-tasking framework, by which the shared function between causal structures is obtained adaptively. In DIRECT-ND[122], the entangled representation is solved through hybrid learning, and the multivariate causal effect estimation problem was studied from a new perspective. Additionally, VAE and GAN networks were added to the model to realize the learning of hybrid representation space.

Aiming at balancing selection bias, Zhang et al. [98] proposed the RSB algorithm using autoencoder networks via PCC regularization and instrumental variables, and also adding the confounding variables and moderators for prediction. As extension of CFRNet[42], both , DR-CFR[99] and DeR-CFR[65] were proposed with the main structure as shown in Figure 6 to remove the covariate correlations.

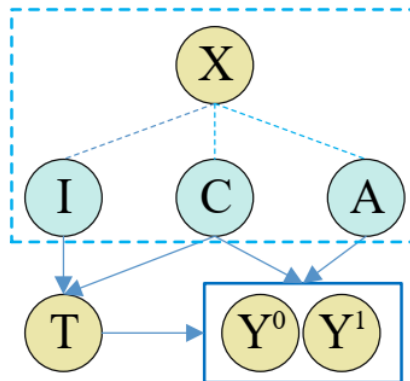


Figure 6: **Decomposed Representation for CounterFactual Regression (DeR-CFR)** [65]: Causal framework with decomposed latent factors

As we can find from Figure 6, there are three possible factors contributing to the observed covariates X in the figure. The instrumental factor I , which only affects the treatment T ; the confounding factor C , which causes the outcome Y along with the treatment T ; and the adjustment factor A , which determines the outcome Y . Learning the decomposed representation for counterfactual reasoning consists of the following steps[65]:

- Three decomposed representation networks for learning latent factors, one for each underlying factor: $I(X)$, $C(X)$, and $A(X)$.
- Three regularizers for confounder identification and balancing are: the first is to decompose A from X by considering $A(X) \perp T$ and $A(X)$ should predict Y as accurately as possible; the second is to decompose I from X by constraining $I(X) \perp Y | T$, and $I(X)$ should be predictive of T , based on Assumption 2; the last is designed for simultaneously balancing confounder $C(X)$ in different treatment arms.
- Two regression networks for potential outcome prediction, one for each treatment arm: $Y^0(C(X), A(X))$ and $Y^1(C(X), A(X))$.

The following is the orthogonal regularizer function used in the decomposition process:

$$\mathcal{L}_O = \bar{W}_I^T \cdot \bar{W}_C + \bar{W}_C^T \cdot \bar{W}_A + \bar{W}_A^T \cdot \bar{W}_I \quad (16)$$

To prevent the representation network from rejecting any input, the sum of \bar{W}_I , \bar{W}_C and \bar{W}_A are constrained to be one. For a hard decomposition, the orthogonal regularizer ensures each variable in X can only flow into one representation network.

Based on DeR-CFR, CATE's prediction performance has been assessed using CF-CV[123], which selects the best model or hyperparameters from potential candidates. In [124], a meta-learning approach was combined together with deep networks, theoretical reasoning, and the optimal counterfactual information.

Causative treatment effect estimation has always been concerned with rationally using confounding variables. The decoupling of covariates to learn related confounding variables can help remove selection bias and generate unbiased output estimates. Despite its theoretical nature, this kind of methods has also some limitations in practical applications, as it requires decomposing covariates into reasonable explanations.

5.3 GANs-based counterfactual simulation

In deep generative models, generative adversarial networks can capture the uncertainty of counterfactual distributions. The generator produces counterfactual results or consistency distribution between control and treatment groups, while the discriminator fits the unbiased estimation of the treatment effects. As well as using factual data, GAN networks also consider the accuracy of counterfactual results when making causal effect estimations. In light of this, generative adversarial models are increasingly used for causal effect estimation.

The first approach suggested by Yoon et al. is for the GANITE[44] network to generate the counterfactual results based on factual data and pass them to the ITE generator. Figure 7 shows the detailed block diagram of GANITE.

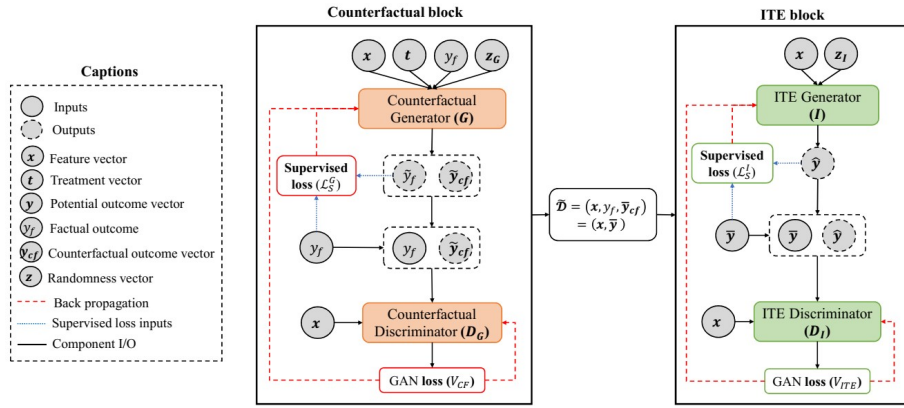


Figure 7: **Generative Adversarial Nets for inference of Individualized Treatment Effects (GANITE) [44]:** Block diagram

For a given feature vector x , GANITE generates the potential output results by first generating factual outputs y_f and then counterfactual samples \hat{y}_{cf} using generator G . After these counterfactual data are combined with the original data, a complete data set \tilde{D} is generated in generator I of the ITE module, which then optimizes \tilde{D} , yielding an unbiased estimate of each treatment’s effect.

For the first time, CEGAN[68] has applied the GAN network to balance the distributions between the spatial treatment group and the control group by leveraging the discriminative loss of the GAN network and weighting the Decoder’s construct loss or weight after the Encoder. In order to solve the generator-discriminator min-max problem, the following reward function is used:

$$\min_{(\theta_E, \theta_I, \theta_P)} \max_{\theta_D} \mathbb{E}_{q_E(\mathbf{z}, \mathbf{x}, t, \mathbf{y})} [\log(D(\hat{\mathbf{z}}, \mathbf{x}, t, \mathbf{y}))] + \mathbb{E}_{q_P(\mathbf{z}, \mathbf{x}, t, \mathbf{y})} [\log(1 - D(\mathbf{z}, \mathbf{x}, t, \hat{\mathbf{y}}))] \quad (17)$$

where the encoder-decoder’s joint distribution is represented by $q_E(\mathbf{z}, \mathbf{x}, t, \mathbf{y})$ and $q_P(\mathbf{z}, \mathbf{x}, t, \mathbf{y})$, while their probability estimations are represented by $(D(\hat{\mathbf{z}}, \mathbf{x}, t, \mathbf{y}))$ and $(1 - D(\mathbf{z}, \mathbf{x}, t, \hat{\mathbf{y}}))$, respectively. The discriminator determines which distribution the samples belong to.

In addition to GANITE and CEGAN, there are also many works using GAN networks to estimate the causal effects in other fields. As part of a generative adversarial framework, GAD[100] applied GAN networks to continuous treatment problems to learn a sample-balanced weight matrix, which removes the association between treatment regimens and covariates; to address multiple treatment as well as consecutive doses treatment problems, DRGAN[101] proposed a model architecture consisting of a counterfactual generator, discriminator, and inference block; As a means of better coping with continuous intervention problems, SCIGAN[69] added a hierarchical discriminator based on DRGAN. In addition, CTAM[33] has also applied the generative adversarial ideas to the treatment effect estimation of text sequence information,. It filters out information related to approximate instrumental variables when learning representations, and matches between the learned representations; To eliminate the association between the treatment and patient history,

CRN[102] utilized a counterfactual recurrent neural network to reflect the time-varying treatment effect; In ABCEI[104], the covariate distributions between the control and treatment groups are well balanced with GAN networks, and a regularization function of mutual information estimators is added to reduce bias; To learn the balanced covariate representations, CETransformer[109] combined the attention mechanism with WGAN; In TransTEE[73], Transformer was used for covariate representation, in which the treatment effectiveness is estimated by the Propensity Score Network and the selection bias can be overcome by the GAN network. In particular, the model can also be used for discrete, continuous, structured or dose-related treatments.

In general, it is not hard to extend the ways of individual treatment effect estimation to multiple interventions and continuous dose interventions using the GAN network, and it has a marvelous effect on the balance of representation distribution and the generation of potential results. However, due to lacking a complete theoretical support system, using GAN networks to solve the problem of causal effect estimation requires more impeccable theoretical derivation in the future.

5.4 Time series causal estimation

In treatment effect estimation, most models focus on numerical variables, and it is still difficult to deal with textual information and time series information[125]. Variable decoupling for textual information estimation can reduce estimation bias since there are many covariates in textual information that are unrelated to causal effect estimation. When dealing with time-series information, RNNs have usually been combined to create counterfactual recurrent networks based on historical information.

The R-MSN[71] model is first proposed by Lim et al. in order to address the problems arising with continuous treatment doses and multi-treatments under time series. Figure 8 illustrates the model's frame structure, which uses a recurrent edge network to remove time-dependent confounding, and a standard RNN structure to encode and decode.

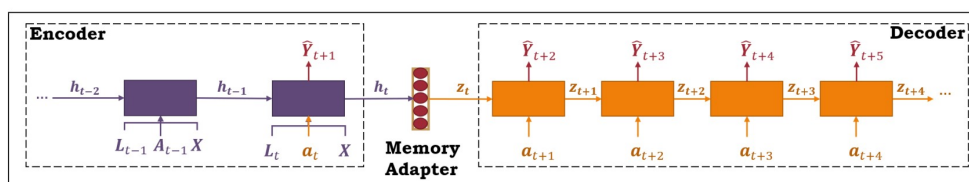


Figure 8: **Recurrent Marginal Structural Network (R-MSN) [71]:** Architecture for multi-step treatment response prediction

To predict the causal effect, R-MSN used the standard LSTM structure, dividing multi-treatment and continuous intervention problems according to the corresponding time interval. As a counterfactual recurrent network, CRN[102] constructs a treatment-invariant representation for each time step based on R-MSN[71], eliminating the patient's medical history association between treatment allocation and treatment allocation and balances time-varying confounding biases; By combining with the current treatment assignment and historical information, the hidden confounders are inferred using recurrent weighted neural networks in DSW[93], and then reweighted using time-varying inverse probabilities. In addition to building a multi-task output RNN factor model, TSD[103] allocates multiple treatment over time, and then estimates the treatment effects with multi-cause hidden confounders by which the latent variables free of treatment can be inferred. Furthermore, it substitutes the unobserved confounders with latent variables, and infers logistic regression results in the absence of treatment; In SyncTwin[113], treatment estimation is performed based on the temporal structure of the prediction results, and synthetic twin samples are constructed and counterfactual predictions are obtained.

Yao et al. proposed a matching treatment-adversarial learning CTAM[33] method that takes into account text sequence information. As shown in Figure 9, when learning representations, it filters out the approximate instrumental variables and then makes matching between the learned representations to estimate the treatment effect.

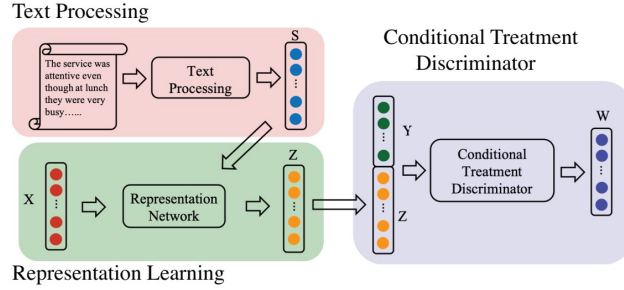


Figure 9: **Conditional Treatment-Adversarial learning based Matching Method (CTAM) [33]: Framework**

Specifically, there are three main components of CTAM[33]: text processing, representation learning, and conditional treatment discrimination. In the first step, the text processing part transforms the original text into a vector representation S , concatenates S with non-text covariates X , and then constructs a unified feature vector that transforms the input into the latent representation Z . As a next step, both Z and Y are fed into the conditioned treatment discriminator, and during the training process, a max-minimum arithmetic unit is calculated between the representation learning network and the conditioned treatment discriminator. In order to filter out information related to instrumental variables, the representation learning network prevents the discriminator from assigning the correlative treatment. As a last step, it implements the match in the representation space Z .

To predict treatment assignment using mutual information between global feature representations and individual feature representations, IDRL[126] proposed to learn Infomax and domain-independent representations. To maximize the ability of capturing the common prediction information among treatment and control groups, the influence of instrumental variables and irrelevant variables were filtered out. In addition, the SCRNet[127] attempted to estimate ITE with different types of variables by dividing the covariates. Recently, CT[115] adopted transformer and lstm to capture complex long-term dependencies between time-varying confounders and proposed a new counterfactual domain confusion loss to address the confusion bias.

Estimation of causal effects of textual time series is often combined with the problems related to multiprocessing and continuous dose processing. Despite the widespread application of this direction, researchers need to develop a standard for measuring intervention effects based on the actual situation, and it is difficult to assess the rationality and reliability of the various working standards used in the industry.

5.5 Multi-treatment and continuous-dose models

Casual estimation for individual treatments focuses on solving binary treatment problems, and extending it to multiple treatment is computationally expensive. However, multiple treatment and continuous-dose treatment models have many applications, such as radiotherapy, chemotherapy and surgery for cancer treatment, as well as prolonged usage of vasopressors for many years. It is therefore beneficial to estimating the effects of ongoing interventions in these various treatment settings in order to make marvelous long-term process decisions.

For the first time, Schwab et al. have extended individual treatment estimation to multi-discrete treatment problems with the PM[41] algorithm. In PM, counterfactual reasoning is utilized in small batches by matching nearest neighbors samples, which makes it easily implemented and is compatible with a wide range of architectures, and there is no need to increase computation complexity or other hyperparameters for treating arbitrary quantity of patients. In order to capture higher-order effects, TECE-VAE[88] modeled the dependence between treatments by using task embedding, extending the problem to arbitrary subsets of multi-treatment situations.

When solving problems involving multi-treatments and continuous-dose treatments, GAN networks are frequently combined. A two-step generative adversarial de-aliasing algorithm proposed by GAD[100] can be used for continuous treatment problems, removing the association between covariates and treatment variables. Specifically, it is along with the following three steps: A) Produce an unbiased distribution with no correlation between the covariates; B) Learn the sample weights and transfer the observed data to the unbiased distribution; C) De-obfuscate the data with generative adversarial networks.

As an improved GAN model, DRGAN[101] takes the form of a generator, a discriminator, and prediction network to generate a complete dose-response curve for each sample by considering both multi-treatment and continuous-dose treatment options; By using a hierarchical discriminator on the basis of DRGAN, SCIGAN[69] was proposed to improve the model's ability of handling the problems of continuous intervention.

Meanwhile, a set of open model benchmark parameters, including MISE, DPE, PE, and model selection criteria, were proposed in DRNet[55], which allows the generation of dose-response curves for an unlimited number of treatments under continuous dose parameters. In VCNet[56] that utilizes a variable coefficient neural network, a continuous ADRF[128, 129, 130] estimator is automatically calculated for the continuous activation function, which is helpful of preventing the processing information from being lost. Moreover, the existing target regularization method was also extended to obtain a double robust ADRF curve estimator.

As part of DRNet[55], continuous treatments are divided into blocks and trained separately into hidden layers, which are then nested into each other to construct a piecewise fit of individual dose-response curves; A continuous prediction head of weighted treatment is created by VCNet[56] by paying closer attention to treatment continuity, and optimizing the individual prediction head into a mapping function of covariates that change with treatment. The structure comparison of DRNet and VCNet models is shown in Figure 10.

TransTEE[73] combines the hierarchical discriminator of SCIGAN[69] with the variable coefficient structure of VCNet[56] and inducts the Transformer multi-headed attention mechanism generic framework to extend the causal effect estimation problem to discrete, continuous, structured, and dose-related treatments.

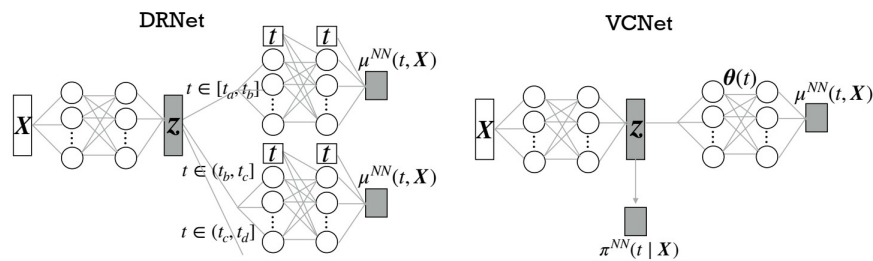


Figure 10: **Dose Response Network (DRNet) [55] and Varying Coefficient Neural Network (VCNet) [56]:** Comparison of network structures

As the first study of the multi-treatment combination problem, NCoRE[108] has used the cross-treatment interaction to infer the causal generative processes underlying multiple treatment combinations, in which the counterfactual representations learned in a treatment setting are combined.

To estimate the multi-cause perturbation treatment effect, Prichard et.al proposed the idea of SCP[112] for the first time. To overcome the confounding bias in two steps, a single-cause CATE estimator is first applied to augment observed data and estimate potential outcomes; As a next step, the augmented data set is adapted for covariates to obtain multi-factor unbiased estimators. In addition to illustrating the relationship between single-factor and multiple-factor problems, SCP shows the equivalence of conditional expectations of single-factor interventions and multiple-factor interventions, which can be verified by the following equation:

$$\mathbb{E}_\alpha (Y (a_k, \mathbf{a}_{-k}) | \mathbf{X}) = \mathbb{E}_k (Y (a_k) | \mathbf{X}, \mathbf{A}_{-k}(a_k) = \mathbf{a}_{-k}) \quad (18)$$

In the first step of augmenting the dataset, outcomes and observations $Y (a_k)$ and $\mathbf{A}_{-k} (a_k)$ are added. As such, by training a supervised learning model on the augmented dataset, it is possible to estimate the expected value on the right side of Eq. 18 as well as the multifactorial intervention effect given by the left side in a way that enhances the generalizability of the estimator. In contrast to SCP, OOSR[131] proposed a prediction model using a reweighting way that puts emphasize on the outcome-oriented treatment.

Recently, more and more researchers have taken interest in the problem of multi-treatment and continuous dose therapy, and have also made significant contributions. Nevertheless, there are still many models in this area that need to be developed. Especially, it is still an urgent issue to solve how to formulate a unified causal effect measurement standard.

6 Current Status of Industrial Applications

Deep learning-based causal effect estimation has been widely applied to data-driven decision making across domains, and plays an important role in incentive allocation, debiased recommendation, precision medicine, and identifying and measuring strategic decision-making. In this section, typical applications in related fields are introduced according to

their corresponding data characteristics, we will focus on industrial applications using deep causal models, involving marketing, E-commerce, finance, medicine, economics, and education, and cast concerns on their social value and application value.

6.1 Marketing applications

To maximize the return on investment, it is crucial to decide how to allocate the user-specific incentive under budget constraints. Uber and Didi Chuxing have incentivized drivers as well as passengers with commission and coupons to boost ride-sharing business. Mobile payments, such as Apple Pay, Alipay, WeChat Pay, have designed various mobile marketing campaigns to encourage users to pay through their applications. Amazon and JD Digits attempted to give out discounts to enhance attraction to customers. Incentive allocation relies on treatment effect estimation models to estimate users' purchase probability with different incentives. PCAN[132] leveraged the small set of unbiased data to train an unbiased model, and established a biased network to generate a representation close to the unbiased network by the distribution difference between the biased and unbiased data representation. Offline and online experimental results demonstrate that the developed method can alleviate the problem of price-bias and lead to significantly improved performance of the resulting allocation policy in a real-world marketing campaign. Besides, DESCN[114] was developed to capture the relationships among the treatment propensity, the true responses, and the pseudo treatment effect in an integrated manner, which was employed in voucher distribution business in Lazada, a leading South-East Asia E-commerce platform of Alibaba Group. Related results indicate that this method is advantageous in both ITE estimation accuracy and uplift ranking performance. For multi-treatment scenarios, Amazon has presented MEMENTO [133]. It is based on obtaining matching representations of the confounders for the various treatment types, achieved through minimization of an upper bound on the sum of factual and counterfactual losses. Besides incentive allocation, deep causal models are also applied to other tasks in marketing. For example, a user retention model UR-IPW[134] was proposed by AntGroup to account for impression-revisit effect, where users could revisit the APP even if they do not explicitly interact with the recommender system. The model makes full use of both explicit and implicit interactions in the observed data, and estimate revisit rate by using inverse propensity weighting accounting for the selection bias caused by user's self-selection.

Practical Application: In bilateral business relations, effective marketing strategy should also motivate merchants as well as customers. For example, to encourage mobile payment activities, merchants get shared incentives with customers after them scanning QR-codes and paying with Alipay. However, making independent optimization for both sides can be nonoptimal. Therefore, [135] and [136] took the mutual influence into account, and used graph neural networks to represent merchants and customers jointly by modeling the underlying bipartite influences. Extensive experimental results demonstrate the effectiveness of the proposed approaches.

6.2 E-commerce applications

Recommender systems play a crucial role in E-commerce. Traditional recommender systems extract user preference based on learning correlation in data observational, resulting in biases including selection bias, exposure bias, position bias, conformity bias, etc. To address this issue, Amazon, Netflix, Criteo, Alibaba, AntGroup, JD, Kuaishou, etc. have all begun to utilize causal effect estimation to extract causality. Datasets and recent works on recommendation debiasing can refer to [137] and [138]. In [81], debiased information bottleneck was proposed by Huawei, which is applicable to various types of bias. The architecture constrains the model to learn a biased embedding vector with independent biased and unbiased components in the training phase, and uses only the unbiased component in the test phase to deliver more accurate recommendations. As mentioned in [139], Wei et al. explored the popularity bias issue from a cause-effect perspective, which performs multi-task learning to achieve the contribution of each cause, and during testing, performs counterfactual inference to remove the effect of item popularity. Following that, ESCM²[140] leveraged the sequential pattern of user actions to address data sparsity issue and employed a counterfactual risk minimizer as a regularizer to address both issues in inherent estimation bias for CVR estimation and potential independence priority for CTCVR estimation simultaneously. Furthermore, in some scenarios, another set of uniform data can also be used to alleviate the bias problems and improve the performance. In [15], Criteo proposed a multi-task objective that jointly factorizes the matrix of biased data and the matrix of uniform data. In [141], a general knowledge distillation framework that enables uniform data modeling was proposed, which consists of four modules including label-based distillation, feature-based distillation, sample-based distillation and model structure based distillation. Differently, in [142], AutoDebias optimized the debiasing parameters by solving the bi-level optimization problem with a small set of uniform data.

Practical Application: In [140], CausalMTA was proposed by Alibaba, which systemically eliminates the confounding bias from both static and dynamic perspectives by reweighting and learn an unbiased conversion prediction model. Its effectiveness is demonstrated by a real ad impression data, which includes 30 days of ad impression data from mobile

phone shops, where touchpoints are categorized into 40 channels, including interact, feed, display, search, live show, etc.

6.3 Financial applications

In the last few decades, economists have learned to take very seriously the old admonition from undergraduate econometrics that “correlation is not causality”. We have surveyed a number of recent developments in the econometrics toolkit for addressing causality issues in the context of estimating the impact of policies. Hennessy et al. [143] analyzed the meaning and utilization of alternative causal effect measures in the types of settings commonplace in finance and economics, those where individual agents have private information and outcome variables are mediated by the beliefs of other agents in the economy. They suggested the utilization of two distinct causal effect definitions which include partial causal effects and total causal effects.

Tiffin [144] focused mostly on the Causal Forest algorithm, and took the costs of a financial crisis as an illustrative example, the paper has endeavored to show how such techniques can produce plausible results, i.e., the estimated average impact of a crisis. A natural application of personalized treatment effect estimation is to estimate optimal policy functions in observational data. Recently, Athey and Wager [145] brought in insights from semiparametric efficiency theory in econometrics to propose a new estimator for optimal policies and to analyze the properties of this estimator. Policies can be compared in terms of their “risk”, which is defined as the gap between the expected outcomes using the (unknown) optimal policy and the estimated policy. Arpino et al. [146] proposed an explicitly model, which was applied to the evaluation of a policy implemented in Tuscany on small handicraft firms, to address the violation of the Stable Cell Handling Value Assumption due to the interference between cells. Results show that the benefits from the policy are reduced when treated firms are subject to high levels of interference.

Practical Application: In the field of Fin Tech, MYbank of Ant Group adopted causal counterfactual inference debiasing methods [147, 107, 112, 108, 148, 149] to solve the biased problem of loan marketing AB experiment. In order to effectively measure the intervention effect, we use the method of causal counterfactual inference to construct a homogeneous control group from the full experimental population based on observational data, so that horizontal comparisons can be made.

6.4 Medical applications

Since the advent of precision medicine and the availability of large amounts of observational data from electronic health records, the research community has started to explore quantitative individual-level effect of a treatment. The goal of clinical decision-makers is to determine the optimal treatment course for any given patient, using static or time-series observational data. However, observational datasets are prone to treatment assignment bias. To address it, a wide variety of models have been proposed.

Many medical applications involve simultaneous intervention on multiple variables. In order to estimate the multi-cause treatment issue, Single-cause Perturbation[112] was proposed, which started by augmenting the observational dataset with the estimated potential outcomes under single-cause interventions, and then performed covariate adjustment on the augmented dataset to obtain the estimator. Following that, GraphITE[150] was proposed, which obtained the representations of the graph-structured treatments using graph neural networks, and also mitigated the observational biases by using the HSIC regularization to increase the independence of the representations of the targets and the treatments.

Observational data also captures information on complex time-dependent treatment scenarios. To estimate the effectiveness of treatment over time, SCIGAN[69] was proposed, which is flexible and capable of simultaneously estimating counterfactual outcomes for several different continuous interventions. Additionally, TE-CDE[151] was proposed, which allows the potential outcomes to be evaluated at any time point. Adversarial training is used to adjust for time-dependent confounding which is critical in longitudinal settings.

Practical Application: The van der Schaar Lab identified and targeted an extensive range of potential clinical applications, including COVID-19, organ transplantation, etc. To decide “one best model” for each scenario, a first-of-its-kind validation procedure is introduced in [152] for estimating the performance of causal effect estimation methods using influence functions, which utilized a Taylor-like expansion to approximate the loss function of a method on a given dataset in terms of the influence functions of its loss on a “synthesized”, proximal dataset with known causal effects.

6.5 Economic applications

Causal effect estimation has attracted more and more attention of researchers in economic science, especially how to better combine with deep learning to solve practical problems. Learning about cause and effect is arguably the main goal in applied econometrics.

For instance, unobserved confounding factors threaten the internal validity of estimates, data availability is often limited to non-random, selection-biased samples, causal effects need to be learned from surrogate experiments with imperfect compliance, and causal knowledge has to be extrapolated across structurally heterogeneous populations. A powerful causal effect estimation framework is required in order to tackle all of these challenges, which plague essentially any data analysis to varying degrees [153]. It nests the prediction power of deep learning to obtain consistent causal estimators under high dimensional covariates [154, 155, 156]. In [157], Angrist and Steve Pischke coined the term “credibility revolution”. They argued that economics turns to transparent empirical strategies applied to specific causal problems. Problem-driven methodological agendas are mostly based on the propensity score theorem of Rosenbaum and Rubin [158]. This theorem changes applied econometrics to focus our attention on the process of determining treatment assignment rather than models for outcomes. Dehejia and Wahba [85] was the first to demonstrate the value of this approach. More recently, Belloni et al. [159] utilized deep learning to model scores while modeling outcomes. This work can be seen as an extension of Robins’s notion of dual robustness to a broader class of empirical strategies. Angrist [160] introduced a novel framework, the local average treatment effects framework for causal effect estimation, to help make the empirical strategies in economics should be transparent and credible. The LATE theorem can tell us for whom particular instrumental variables and regression discontinuity estimates are valid. Hal R. Varian [161] argued that the powerful techniques used in deep learning may be useful for developing better estimates of the counterfactual, potentially improving causal effect estimation. For generalized neighbor matching to estimate individual and average treatment effects, Vikas [162] proposed the use of deep learning techniques in econometrics, specifically for causal effect estimation and for estimating individual as well as average treatment effects.

6.6 Educational applications

Education plays a central role in modern society, especially in earlier years [163], a wave of new studies on the effects of educational interventions on student performance has emerged. Therefore, the need for evidence-based policy in the field of education is increasingly recognized. Education policy-makers and practitioners want to know which policies and practices can best achieve their goals. However, providing empirical evidence suitable for guiding policy is not an easy task, because it refers to causal effect estimations that require special research methods which are not always easy to communicate due to their technical complexity. From a Bayesian perspective, David [164, 165] introduced a review and synthesis of the problem of causal inference in large scale educational assessments which requires the articulation of framework for causal effect estimation followed by a statistical approach that closely matches the framework and can yield the causal estimate of interest. Zhao et al. [39] proposed the Residual Counterfactual Networks in an Intelligent Tutor System can decide which hint is more suitable for a specific student. However, the effectiveness of an intervention is necessarily multifaceted and complex effects differ between students, as a function of implementation [166], and, potentially, as a function of time and location. Sales et al. [167] explored a different sort of treatment effect heterogeneity differences in effectiveness for different outcomes. Specifically, different posttest items measuring different skills. Carvalho [168] conducted causal effect estimation on students’ online behavior patterns using a different toolkit, GeNIe3, using data from a learning management system. Chen et al. [169] developed a causal discovery framework that utilized TETRAD [170].

7 Guideline for Experiment

After an in-depth description of the deep causal modeling approach, this section will provide a detailed experimental guide, including a comprehensive conclusion and analysis of datasets, source codes as well as experiments.

7.1 Datasets

As counterfactual results can never be observed in real life, finding datasets that satisfy experimental requirements is difficult. Most of the datasets used in the literature are semi-synthetic. Following the survey[2], we further supplemented in Table 3 more relevant datasets that are popularly used for causal analysis. Meanwhile, we also conclude the application scenarios for these datasets. In addition, Table 4 summarizes the web links to these datasets and the classical models that are evaluated on them. Below are the detailed descriptions of the available datasets.

IHDP. the Infant health and development[86] dataset was generated from a randomized controlled trial of preterm infants with low birth weight. Various features of the children and their mothers are measured as pre-treatment

covariates, such as birth weight, head circumference, neonatal health index, prenatal care, mother's age, education, drugs, and alcohol. Intensive high-quality childcare is presented to infants in the treatment group, such as specialist home visits[171]. The outcome is a score on the cognitive test for the infant. Moreover, noisy subsets of treatment groups need to be removed to build unbiased selection models.

Jobs. The employment dataset studied by Jobs in LaLonde (1986)[87] is composed of randomized data based on state-supported work programs and nonrandomized data from observational studies. The pre-treatment covariates include eight variables such as age, education, race, and income in 1974 and 1975. The treatment group has taken part in the vocational training, while the control group has't. The outcome is the employment status.

Twins. The Twins dataset comes from data on twin births in the United States from 1989-1991[172]. An evaluation of 40 covariates pertaining to pregnancy, twin births, and parents is carried out for every pair of twins, including gestational weeks just before birth, quality of care during pregnancy, pregnancy risk factors (anemia, alcohol, smoking, etc.), nursing, residence, and more. The outcome is a one-year mortality rate. A twin dataset is available with results from the treatment (heavier of the twins) and control groups (lighter of the twins). Selection bias is typically simulated by assigning different treatments based on user-defined criteria.

News. The News dataset consists of 5000 randomly sampled news articles from the New York Times corpus. The news dataset contains data on media consumers' perceptions of news items. A sample is a news item consisting of word counts, the results are readers' opinions, and the treatments are a variety of devices that can be used to view the news item, such as smartphones, tablets, computers, and TVs.

ACIC. A causal effect estimation data analysis challenge has been held every year at the Atlantic Causal Inference Conference since 2016, which presents different data sets for a variety of causal effect estimation problems. Below we describe in detail two typical datasets, ACIC2016 and ACIC2018. A summary of the latest conference dataset about ACIC can be found in [173].

The ACIC 2016 consists of 77 datasets with different degrees of nonlinearity, sparsity, correlation between treatment assignment and outcome, and overlap between treatment effects. Covariates are derived from real data from the IHDP[86] dataset, which consists of 58 variables and 4802 samples[174]. The simulation model generates treatment, factual and counterfactual outcomes, while the selection bias is created by removing treated children who are mothers of nonwhites. The ACIC 2018 is a benchmarking dataset for causal effect estimation that is commonly used[175]. It is a semi-synthetic dataset about infant births and deaths[176] and contains 63 datasets with each drawn randomly from a different distribution, which are then generated by a simulation process.

TCGA. As the world's largest and most comprehensive genomic database, the Cancer Genome Atlas (TCGA)[92] contains billions of genomes. A total of 9658 individuals are included in the TCGA[92] dataset, the treatment regimens are drug treatment, chemotherapy, and surgery, and the outcome is the risk of developing cancer after treatment.

PK-PD model of tumor growth. A model of pharmacokinetic-pharmacodynamics (PK-PD)[177] can be used to explore dose-response relationships and suggest optimal treatments. Among its key functions are the combination of chemotherapy and radiotherapy effects, post-treatment cellular regeneration, patient death or recovery, and cancer-based different gaze distributions of tumor size at the diagnostic stage, which make this model an excellent one for treating non-small cell lung cancer patients. The PKPD model enables clinicians to explore hypotheses about dose-response relationships and suggest optimal treatment options[178, 179]. In the most classic example of PK-PD, tumor growth[177] with time-dependent confounding can be predicted by observing the expected response to treatment, chemotherapy, and radiotherapy.

MIMIC III. Medical Information Mart for Intensive Care(MIMIC III)[94] is a database of electronic health records from ICU patients. The benchmark dataset consists of 7413 samples with 25 covariates after filtering for missing values. As far as treatment options go, antibiotics, vasopressors, and mechanical ventilators are the most common options in the ICU to treat patients with sepsis. A number of laboratory use patient vital signs over time as a measure to assess the effects of antibiotics, vasopressors, and mechanical ventilators on the following covariates, including white blood cells, blood pressure, and oxygen saturation. A comprehensive and detailed description of the clinical data can be found in [180].

NICO. There is a bias in sample selection when using the image dataset NICO with context for object classification[106]. Cat or dog classification in the "animals" dataset in NICO is seen as a benchmark distribution for non-i.i.d. The parameters include the time of sampling, whether to sample, the context, and the semantic shape of cats and dogs, as well as the "grass" and "snow" environment.

ADNI. Alzheimer's Disease Neuroimaging Initiative(ADNI)[181] dataset has three latent representation outputs Alzheimer's Disease, Mild Cognitive Impairment and Normal Control. The covariates are age and TAU[182], which determine whether Magnetic resonance imaging should be used as an input for therapy.

Table 3: Overview of some open available datasets for causal analysis and their application scenario

Datasets	Binary	Multiple	Continuous dose	Times series	Structured
IHDP	✓	✓			
Jobs	✓				
Twins	✓				
News	✓	✓	✓		
ACIC		✓	✓		
TCGA			✓		✓
Tumor Growth				✓	
MIMIC III			✓		
NICO	✓				
CMNIST	✓				
ADNI		✓			
COVID-19		✓			
CPRD				✓	
BlogCatalog		✓			✓
Flickr		✓			✓

COVID-19. During the first peak of the pandemic, dataset COVID-19[90, 89] Hospitalization in England Surveillance System (CHES) collected individual-level risk factors, treatments, and outcomes from 3090 ICU patients. There are a number of covariates, including factors such as age and multiple morbidity, as well as treatment parameters, such as ventilation and antiviral drugs. The outcome is the length of stay in the intensive care unit[183].

CPRD. Clinical Practice Research Datalink (CPRD) contains records from NHS general practice clinics in the United Kingdom, covering approximately 6.9 percent of the country’s population[184]. National mortality records and hospital event statistics indicate that CPRD is associated with secondary care admissions. Low-density lipoprotein is measured after CPRD is initiated, and treatment is defined as the date of first prescription. As temporal covariates, the following risk factors is measured before treatment initiation: high-density lipoprotein cholesterol, blood pressure, pulse, creatinine, triglycerides, and smoking status. HPS registry participants are selected from 125,784 individuals who meet the eligibility criteria. A total of 17,371 treatment groups and 24,557 control groups are divided into three equally sized subsets for training, validation, and testing.

BlogCatalog. BlogCatalog is an online community where users post blogs. In the dataset, each instance is a blogger[185]. Each edge represents a social relationship between two bloggers. Blog descriptions contain keywords represented as a bag-of-words. Blog reader opinions as input, whether blog-created content gets more comments on mobile or desktop as therapy, parameters for individual treatment effect estimation are derived from the content of readers’ comments on mobile (than desktop). The blogger belongs to the treatment group if people reads more on a mobile device than on a desktop device, and vice versa.

Flickr. Flickr is an online social networking site where users can share photos and videos[186]. The dataset consists of instances representing users, and edges representing social relationships between them. Tags of interest are represented by the features of each user. Generally, Settings and assumptions are the same as for BlogCatalog dataset.

Table 4: The web links to the open available datasets and the related models that have used them for performance evaluation

Datasets	Links	Methods
IHDP	https://www.fredjo.com	[40, 95, 42, 52, 19, 96, 44, 43, 41, 33, 66, 53, 98, 54, 99, 104, 105, 56, 109, 65, 111, 124, 109, 73, 127]
Jobs	https://www.fredjo.com	[42, 52, 44, 43, 53, 54, 104, 109, 110, 72, 127, 126]
Twins	www.nber.org/data/linked-birth-infant-death-data-vital-statistics-data.html	[52, 44, 68, 43, 53, 54, 100, 104, 109, 110]
News	https://archive.ics.uci.edu/ml/datasets/bag+of+words	[40, 41, 33, 101, 69, 72, 55, 56, 73, 126]
ACIC	https://www.synapse.org/ACIC2018Challenge	[66, 111, 104, 187, 188, 189, 190, 191, 192, 193, 114]
TCGA	https://gdc.cancer.gov/	[101, 69, 72, 73, 131, 194, 195, 196, 197, 198]
Tumor Growth	www.nature.com/scientificreports/	[71, 102, 199, 200, 201, 202]
MIMIC III	https://mimic.physionet.org/	[103, 104, 69, 55, 72, 115, 203]
NICO	https://www.dropbox.com/sh/8mouawi5guauyb/AAD4fdy8rA6fn3PgSmhKwFgvadl=0	[106, 204, 205, 206, 207, 208]
ADNI	https://adni.loni.usc.edu/	[106, 209, 210, 211, 212]
COVID-19	https://www.heywhale.com/mw/dataset/5e8ee81fe7ec38002400f9cb	[112, 213, 214, 215]
CPRD	https://academic.oup.com/ije/article/44/3/827/632531	[113, 216, 217, 218, 219, 220, 221]
BlogCatalog	https://www.blogcatalog.com	[119, 222, 223]
Flickr	https://www.flickr.com	[119]

7.2 Source codes

Most existing reviews are limited to an overview of traditional algorithm code. Here we summarize the source code associated with deep learning and the programming frameworks used. In academia and industry, PyTorch and TensorFlow are the mainstream deep learning frameworks, with more than 80 % scholars and programmers using them for model construction. In addition to the popularly used datasets as motioned above, we also list in Table 5 the available source codes for some representative models and the relevant dataset they have used.

Table 5: Overview of some available datasets and web links to them

Methods	Datasets	Framework	Links
DCN-PD[95]	IHDP	Pytorch	https://github.com/Shantanu48114860/Deep-Counterfactual-Networks-with-Propensity-Dropout
BNN[40]/CFRNet[42]	IHDP,Jobs,News	Tensorflow	https://github.com/clinicalml/cfrnet
CEVAE[52]	IHDP,Twins,Jobs	Tensorflow	https://github.com/AMlab-berkeley/CEVAE
GANTTE[44]	IHDP,Twins,Jobs	Tensorflow	https://github.com/jyoon0923/GANTTE
SITE[43]	IHDP,Twins,Jobs	Tensorflow	https://github.com/qsier-Yi/SITE
R-MSN[71]	PK-PD model of tumor growth	Tensorflow	https://github.com/sjblm/rmsn_nips_2018
PM[41]	IHDP,News	Tensorflow	https://github.com/d916b/perfect_match
Dragonnet[66]	IHDP,ACIC	Tensorflow	https://github.com/claudiahi57/dragonnet
DKLITE[54]	IHDP,Twins,Jobs	Tensorflow	https://github.com/vanderschaarlab/mlforhealthlabpub/tree/main/alg/dk-lite
CRN[102]	PK-PD model of tumor growth	Tensorflow	https://github.com/vanderschaarlab/mlforhealthlabpub/tree/main/alg/counterfactual_recurrent_network
TSJ[103]	MIMIC III	Tensorflow	https://github.com/vanderschaarlab/mlforhealthlabpub/tree/main/alg/time_series_deconfounder
ABCE[104]	IHDP,Twins,Jobs,ACIC,MIMIC III	Tensorflow	https://github.com/octeuffer/Adversarial-Balancing-based-representation-learning-for-Causal-Effect-Inference
LaCIM[106]	NICO,CMNIST,ADNI	Pytorch	https://github.com/vubotong/LaCIM
SCIGAN[69]	TCGA,News,MIMIC III	Tensorflow	https://github.com/soanabica/SCIGAN
DRNet[55]	TCGA,News,MIMIC III	Tensorflow	https://github.com/d909b/drnet
VCNet[56]	IHDP,News	Pytorch	https://github.com/lushleaf/varying-coefficient-net-with-functional-tr
DeR-CFR[65]	IHDP	Tensorflow	https://github.com/angwu/DeR-CFR
DONUT[110]	IHDP,Twins,Jobs	Tensorflow	https://github.com/tohhatt/donut
FlexTENet[111],CATE Nets[124]	IHDP,Twins,ACIC	Jax,Pytorch	https://github.com/AliciaCurth/CATE Nets
SCP[112]	COVID-19	Pytorch	https://github.com/vanderschaarlab/Single-Cause-Perturbation-NeurIPS-2021
SyncTwin[113]	CPRD	Pytorch	https://github.com/vanderschaarlab/SyncTwin-NeurIPS-2021
TransTEE[73]	IHDP,News,TCGA	Pytorch	https://github.com/hlzhang109/TransTEE
CF-CV[123]	IHDP	Tensorflow	https://github.com/usaaito/counterfactual_cv
CGN[70]	MNIST,ImageNet	Pytorch	https://github.com/autonomousvision/counterfactual_generative_networks
DESCN[114]	ACIC,Epilpsy	Pytorch	https://github.com/kailiang-zhong/DESCN
CT[115]	MIMIC III	Pytorch	https://github.com/Valentyn1997/CausalTransformer

By combining related methods, datasets, and source codes, we can more easily identify the innovation points in each model. Meanwhile, it will also facilitate more fair comparison in performance evaluation. In addition, undoubtedly, these source codes will also greatly promote the development of research community about causal effect estimation. As an example, the covariate decomposition is applied to the Dragonnet[66] model when combined with the DeR-CFR[65] model to make a further model optimization. By applying the TransTEE[73] attention mechanism to the representation balance part of VCNet[56] or DRNet[55], the continuous dose estimation curve can be fitted more accurately. It also means the latest advances in causal analysis have also benefited from or inspired by some previous representative works.

7.3 Experiment

We also provide an experimental summary of the binary treatment, multiple treatment and continuous-dose treatment scenarios on the above datasets, respectively. it should be pointed out that the reported results are under unified dataset setting for the compared models. Detailed experimental results and performance comparisons can be found in Tables 6, 7, 8, 9 and 10, respectively, in which the **Mean \pm Std** of the estimated treatment effect results (lower better) are presented.

7.3.1 Settings and results for binary treatment

In binary treatment situation, the datasets are the same as[42, 43, 53], which are three public datasets, i.e., IHDP[86], Jobs[87], and Twins[172]. On IHDP and Twins datasets, the average over 10 realizations with 63/27/10 ratio of train/validation/test splits, and on Jobs dataset, the average over 10 train/validation/test splits with ratios 56/24/20. Relevant experimental results can be found in Tables 6, 7 and 8.

IHDP: The treatment group is made unbalanced by removing a subset of the biased treatment population prior to the experiment. The simulated results adopt the setting "A" of the NPCI package. As with[171], the real effect is calculated using noise-free results. Moreover, by constructing a biased subsample of the IHDP dataset the impact of the growing imbalance between the original treatment groups can be studied.

Twins: This version focuses on same-sex twin couples weighing less than 2,000 grams. After eliminating the records containing missing features, the final dataset contains 5409 records. The procedure $T_i \mid \mathbf{x}_i \sim \text{Bern}(\text{Sigmoid}(\mathbf{w}^T \mathbf{x} + n))$, where $\mathbf{w}^T \sim \mathcal{U}((-0.1, 0.1)^{40 \times 1})$ and $n \sim \mathcal{N}(0, 0.1)$ is used in the experiments to selectively choose one of the twins as an observation and hide the other one, generating a selection bias. The average over 10 realizations with 63/27/10 ratio of train/validation/test splits.

Jobs: Jobs is a binary classification task, in which the goal is to use the feature set of [224] to predict unemployment. The LaLonde trial sample [225] consists of 297 treatments and 425 controls, and the PSID comparison groups contains 2490 controls. At the end of study, a total of 482 (15%) subjects are unemployed.

Table 6: Overview of performance comparisons on IHDP dataset by some representative models in the case of binary treatment

Methods	IHDP(ϵ_{ATE})		IHDP(ϵ_{PEHE})	
	In-sample	Out-sample	In-sample	Out-sample
BNN[40, 42]	0.37 ± 0.03	0.42 ± 0.03	2.2 ± 0.1	2.1 ± 0.1
TARNet[42]	0.26 ± 0.01	0.28 ± 0.01	0.88 ± 0.02	0.95 ± 0.02
CFR _{MMD} [42]	0.30 ± 0.01	0.31 ± 0.01	0.73 ± 0.01	0.78 ± 0.02
CFR _{WASS} [42]	0.25 ± 0.01	0.27 ± 0.01	0.71 ± 0.02	0.76 ± 0.02
CEVAE[52]	0.34 ± 0.01	0.46 ± 0.02	2.7 ± 0.1	2.6 ± 0.1
GANITE[44]	0.43 ± 0.05	0.49 ± 0.05	1.9 ± 0.4	2.4 ± 0.4
SITE[43]	—	—	0.604 ± 0.093	0.656 ± 0.108
ACE[53]	—	—	0.489 ± 0.046	0.541 ± 0.061
DKLITE[54]	—	—	0.52 ± 0.02	0.65 ± 0.03
DR-CFR[99]	0.240 ± 0.032	0.261 ± 0.036	0.657 ± 0.028	0.789 ± 0.091
CETransformer[109]	—	—	0.46 ± 0.02	0.51 ± 0.03
DeR-CFR[65]	0.130 ± 0.020	0.147 ± 0.022	0.444 ± 0.020	0.529 ± 0.068
DONUT[110]	0.13 ± 0.01	0.19 ± 0.02	—	—

Table 7: Overview of performance comparisons on Twins dataset by some representative models in the case of binary treatment

Methods	Twins($\hat{\epsilon}_{ATE}$)		Twins(ϵ_{PEHE})	
	In-sample	Out-sample	In-sample	Out-sample
BNN[40, 42]	0.0056 ± 0.0032	0.0203 ± 0.0071	0.307 ± 0.001	0.309 ± 0.004
TARNet[42]	0.0108 ± 0.0017	0.0151 ± 0.0018	0.314 ± 0.001	0.313 ± 0.002
CFR _{MMD} [42]	—	—	0.312 ± 0.001	0.316 ± 0.003
CFR _{WASS} [42]	0.0112 ± 0.0016	0.0284 ± 0.0032	0.308 ± 0.001	0.309 ± 0.003
GANITE[44]	0.0058 ± 0.0017	0.0089 ± 0.0075	—	—
SITE[43]	—	—	0.309 ± 0.002	0.311 ± 0.004
ACE[53]	—	—	0.306 ± 0.000	0.301 ± 0.002
DKLITE[54]	—	—	0.288 ± 0.001	0.293 ± 0.003
CETransformer[109]	—	—	0.287 ± 0.001	0.289 ± 0.002
DONUT[110]	0.0025 ± 0.0016	0.0033 ± 0.0026	—	—

Table 8: Overview of performance comparisons on Jobs dataset by some representative models in the case of binary treatment

Methods	Jobs(ϵ_{ATT})		Jobs($\mathcal{R}_{poi}(\pi_f)$)	
	In-sample	Out-sample	In-sample	Out-sample
BNN[40, 42]	0.04 ± 0.01	0.09 ± 0.04	0.20 ± 0.01	0.24 ± 0.02
TARNet[42]	0.05 ± 0.02	0.11 ± 0.04	0.17 ± 0.01	0.21 ± 0.01
CFR _{MMD} [42]	0.04 ± 0.01	0.08 ± 0.03	0.18 ± 0.00	0.21 ± 0.01
CFR _{WASS} [42]	0.04 ± 0.01	0.09 ± 0.03	0.17 ± 0.01	0.21 ± 0.01
CEVAE[52]	0.02 ± 0.01	0.03 ± 0.01	0.15 ± 0.00	0.26 ± 0.00
GANITE[44]	0.01 ± 0.01	0.06 ± 0.03	0.13 ± 0.01	0.14 ± 0.01
SITE[43]	—	—	0.224 ± 0.004	0.219 ± 0.009
ACE[53]	—	—	0.216 ± 0.005	0.215 ± 0.009
DKLITE[54]	—	—	0.13 ± 0.01	0.14 ± 0.01
CETransformer[109]	—	—	0.12 ± 0.01	0.13 ± 0.00
DONUT[110]	0.01 ± 0.00	0.06 ± 0.05	—	—
SCI[72]	—	—	0.204 ± 0.008	0.225 ± 0.014

7.3.2 Settings and results for multiple and continue-dose treatment

In multiple and continuous dose treatment situations, there are three public datasets including TCGA[92], News[40] and MIMIC III[94] that are derived from[55, 69]. The datasets are split into 64/16/20% for training, validation, and testing respectively. The experimental results of TARNet[42], TransTEE[73] and VCNet[56] on the TCGA dataset in Table 9 are derived from the re-implementation of the source code.

Three treatments are used in the experiment[101, 69], each with a dose. In addition, each treatment, w , is associated with a set of parameters, $\mathbf{v}_1^w, \mathbf{v}_2^w, \mathbf{v}_3^w$. These parameters are sampled randomly by sampling a vector from $\mathcal{N}(\mathbf{0}, \mathbf{1})$ and then normalized. The experiments add $\epsilon \sim \mathcal{N}(0, 0.2)$ noise to the results. Interventions are assigned by drawing a dose d_w from the beta distribution for each treatment, $d_w | \mathbf{x} \sim \text{Beta}(\alpha, \beta_w)$. After that, the experiment assigns a treatment based on $w_f | \mathbf{x} \sim \text{Categorical}(\text{softmax}(\kappa f(\mathbf{x}, d_w)))$ where increasing κ increases the selection bias.

TCGA: This version uses the measurements of the 4000 most variable genes. Each feature of the gene expression data is scaled in the $[0, 1]$ interval. The experiment gives meaning to the treatment and dose by considering the treatment as chemotherapy/radiotherapy/immunotherapy and its corresponding dose[55].

News: The experiments intercepted 10,000 news articles, each with 2,858 features. Treatment and dose are given meaning, with treatment referring to the viewing device used to read the article (e.g., phone, tablet, etc.) and dose referring to the amount of time spent reading the article[42, 55].

MIMIC: The experiments intercepted 3000 patients that receive antibiotics treatment, using 9 clinical covariates, namely age, temperature, heart rate, systolic and diastolic blood pressure, SpO2, FiO2, glucose, and white blood cell count, measured at start of ICU stay. Likewise, the features is scaled in the $[0, 1]$ interval. Different antibiotics and their corresponding doses are considered as treatment[69].

Table 9: Overview of performance comparisons on TCGA, News, and MIMIC datasets by some representative models in the case of multiple and continue-dose treatments

Methods	TCGA			News			MIMIC		
	MISE	DPE	PE	MISE	DPE	PE	MISE	DPE	PE
TARNet[42]	5.76 ± 0.15	0.53 ± 0.06	0.65 ± 0.07	—	—	—	—	—	—
DRN-W[55]	3.71 ± 0.12	0.50 ± 0.05	0.63 ± 0.05	5.07 ± 0.12	4.21 ± 0.11	4.56 ± 0.12	4.47 ± 0.12	0.53 ± 0.05	1.37 ± 0.05
DRNet[55]	3.64 ± 0.12	0.51 ± 0.05	0.67 ± 0.05	4.98 ± 0.12	4.39 ± 0.11	4.17 ± 0.11	4.45 ± 0.12	0.52 ± 0.05	1.44 ± 0.05
DRGAN-SCIGAN[69]	1.89 ± 0.05	0.31 ± 0.05	0.25 ± 0.05	3.71 ± 0.05	4.14 ± 0.11	3.90 ± 0.05	2.09 ± 0.12	0.51 ± 0.05	0.32 ± 0.05
VCNet[56]	6.36 ± 0.11	0.22 ± 0.04	0.32 ± 0.02	—	—	—	—	—	—
TransTEE[73]	6.40 ± 0.14	0.17 ± 0.03	0.78 ± 0.05	—	—	—	—	—	—
TransTEE+TR[73]	6.26 ± 0.13	0.08 ± 0.02	0.96 ± 0.06	—	—	—	—	—	—
TransTEE+PTR[73]	6.36 ± 0.14	0.18 ± 0.03	0.81 ± 0.05	—	—	—	—	—	—

7.3.3 Settings and results for time series treatment

For time series treatment, there are two public datasets derived from [115], one semi-synthetic data generated base on MIMIC-III[94] and one real-world data of MIMIC-III[94]. For estimating counterfactual outcomes over time, five methods are evaluated, including MSMs[226], RMSNs[71], CRN[102], G-Net[227] and Causal Transformer[115]. For comparability, train/validation/test subsets are splited with ratios 60/20/20. The hyperparameter tuning for these methods are all via random grid search with respect to the factual RMSE of the validation set. The means and standard deviations are over five runs.

MIMIC-III semi-synthetic: In this semi-synthetic dataset, trajectories are generated with outcomes under endogeneous and exogeneous dependencies while considering treatment effects. Specifically, 25 time-varying covariates and 3 static covariates are extracted. The ground-truth counterfactuals for evaluation are accessible.

MIMIC-III real-world: For real-world dataset, the same 25 time-varying covariates and 3 static covariates are used. Vasopressors and mechanical ventilation are considered as two treatments, and (diastolic) blood pressure is considered as outcome. The application of vasopressors is highly confounded by previous and current levels of blood pressure, as they aim to raise low blood pressure.

Table 10: Overview of performance comparisons on semi-synthetic and real-world MIMIC-III datasets by some representative models in the case of time series treatments for τ -step-ahead prediction

Methods	MIMIC-III semi-synthetic (RMSE)					MIMIC-III real-world (RMSE)				
	$\tau=1$	$\tau=2$	$\tau=3$	$\tau=4$	$\tau=5$	$\tau=1$	$\tau=2$	$\tau=3$	$\tau=4$	$\tau=5$
MSM[226]	0.37 ± 0.01	0.57 ± 0.03	0.74 ± 0.06	0.88 ± 0.03	1.14 ± 0.10	6.37 ± 0.26	9.06 ± 0.41	11.89 ± 1.28	13.12 ± 1.25	14.4 ± 1.12
RMSN[71]	0.24 ± 0.01	0.47 ± 0.01	0.60 ± 0.01	0.70 ± 0.02	0.78 ± 0.04	5.20 ± 0.15	9.79 ± 0.31	10.52 ± 0.39	11.09 ± 0.49	11.64 ± 0.62
CRN[102]	0.30 ± 0.01	0.48 ± 0.02	0.59 ± 0.02	0.65 ± 0.02	0.68 ± 0.02	4.84 ± 0.08	9.15 ± 0.16	9.81 ± 0.17	10.15 ± 0.19	10.40 ± 0.21
G-Net[227]	0.34 ± 0.01	0.67 ± 0.03	0.83 ± 0.04	0.94 ± 0.04	1.03 ± 0.05	5.31 ± 0.05	11.88 ± 0.20	12.91 ± 0.26	13.57 ± 0.30	14.08 ± 0.31
Causal Transformer[115]	0.20 ± 0.01	0.38 ± 0.01	0.45 ± 0.01	0.49 ± 0.01	0.52 ± 0.02	4.58 ± 0.09	8.99 ± 0.21	9.59 ± 0.22	9.91 ± 0.26	10.14 ± 0.29

Based on the experimental analysis, we found that with the development and evolution of the deep model, it has significant implications on solving the core issues and challenges such as representation learning, de-biasing, and counterfactual inference for causal effect estimation. Whether it is in the decision making of therapeutic interventions, the fitting of dose curves or the capture of time-varying confounding, deep causal models give us a new perspective to explore a broader range of directions.

8 Conclusions and Future Prospect in Industrial Application

8.1 Conclusions

Deep causal models have become increasingly popular as a research topic because of the development of causal effect estimation and deep learning. It is possible to improve causal effect estimation accuracy and unbiasedness by applying deep network models to causal effect estimation. Moreover, the deep network can be optimized and improved by the profound theory used in causal effect estimation. This survey presents the development of deep causal models and the evolution of various methods. Firstly, the basic knowledge related to the field of causal effect estimation is adhibited. Then, we present the classical treatments and metrics. Additionally, we provide a comprehensive analysis of the deep causal model from temporal development. Next, we divide the deep causal modeling methods into five groups with an overview and analysis. Furthermore, We furnish a comprehensive conclusion of the application of causal effect estimation in industry. Finally, we summarize the relevant benchmark datasets, open source codes, and performance results as experimental guidelines.

Since 2016, causal effect estimation has been combined with deep learning models for the first time in the binary treatment case for estimation of counterfactual outcomes. So far, deep causal models have been used for time-series, multivariate treatment, and continuous-dose treatment situations. It is inseparable from the proposal of deep network models such as AE, GAN, RNN, and Transformer by researchers in the field of deep learning, the generation and simulation of datasets such as IHDP, Twins, Jobs, News, and TCGA by researchers in the field of statistics, and the exploration of ATE, PEHE, MISE, DPE by researchers in industry guided with the theory of Potential Outcome Framework. We believe that with the joint efforts of everyone in the community of causal learning, deep causal models will flourish for the benefit of society and humanity.

8.2 Future prospect in industrial application

For marketing applications, in addition to incentive efficacy evaluation scores, the interpretability of deep learning is also necessary to understand why we predict the output and use it as a basis for business innovation. Furthermore, marketing decisions may involve ethical and legal issues, such as loan applications. Therefore, it makes sense to use causal effect estimation as a guarantee of fairness.

For e-commerce applications, existing debias methods in recommendation systems are usually designed to address only one or two specific biases. There is an urgent need for a universal debias framework to deal with all kinds of prejudice. In addition, how to evaluate a recommendation system fairly and impartially is also an important issue. Existing methods either require accurate propensity scores or rely on unbiased data. Therefore, deep causal models are urgently needed to provide theoretical assurance.

For financial and economic applications, it is desirable to integrate both micro- and macro-level data to study causal effects for stability. In addition, uncertainty quantification is also beneficial to decision making processes, which can be realized by Bayesian approximation and ensemble learning techniques with deep learning.

For medical and educational applications, deep causal models can be adopted in a variety of fields including processing high-dimensional data, enriching randomized trials with real-world data, evaluating spillover causal effects, and moving from studies conducted on specific populations to other populations of interest.

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