When deep learning meets causal inference: a computational framework for drug repurposing from real-world data

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ABSTRACT

Drug repurposing is an effective strategy to identify new uses for existing drugs, providing the quickest possible transition from bench to bedside. Existing methods for drug repurposing that mainly focus on pre-clinical information may exist translational issues when applied to human beings. Real world data (RWD), such as electronic health records and insurance claims, provide information on large cohorts of users for many drugs. Here we present an efficient and easily-customized framework for generating and testing multiple candidates for drug repurposing using a retrospective analysis of RWDs. Building upon well-established causal inference and deep learning methods, our framework emulates randomized clinical trials for drugs present in a large-scale medical claims database. We demonstrate our framework in a case study of coronary artery disease (CAD) by evaluating the effect of 55 repurposing drug candidates on various disease outcomes. We achieve 6 drug candidates that significantly improve the CAD outcomes but not have been indicated for treating CAD, paving the way for drug repurposing.

Introduction

Drug repurposing (a.k.a., drug repositioning) is a strategy to accelerate the drug discovery process by identifying novel uses for existing approved drugs [1]. The primary advantage of drug repurposing over traditional drug development is that it starts from compounds with well-characterized pharmacology and safety profiles and can significantly reduce the risk of adverse effects and attrition in clinical phases [2].

While many successful repurposed drugs (e.g., Viagra for erectile dysfunction) have been discovered serendipitously [3], computation-based repurposing methods are developed recently by leveraging structural features of compounds or proteins [4, 5], genome-wide association study (GWAS) [6], transcriptional responses [7], and gene expression [8]. These methods focus primarily on using pre-clinical information. Unfortunately, the clinical therapeutic effects in humans are not always consistent with pre-clinical outcomes [9].

In healthcare, real world data (RWD) [10] refers to longitudinal observational data derived from sources that are associated with outcomes in a heterogeneous patient population in real-world settings, such as patient surveys, electronic health records (EHRs), and claims and billing activities. Since RWDs are direct observations from human bodies, they become a promising source for drug repurposing. Few researchers have already validated a small number of repurposing drug candidates on RWD [11, 12]. However, there are some limitations with these approaches. First, most studies are complementary (i.e., the original hypotheses usually come from other studies). Second, their studied number of repurposing candidates is limited and unable to proactively generate de novo repurposing drug candidates.

In this study, we follow protocols of randomized clinical trial (RCT) design [13], and computationally screen repurposing candidates for beneficial effect by explicitly emulating the corresponding clinical trials using RWDs. Considering the inherent characteristics of RWD (i.e., temporal sequence data and existing confounding variables [14]), we apply deep learning and causal inference methodologies to control the confounders in RWD, and systematically estimate the drug effects on various disease outcomes. Specifically, the estimated drug effects are obtained by long short-term memory (LSTM) [15] and inverse probability of treatment weighting (IPTW) [16], on MarketScan claims data [17].

As a test case, we apply the proposed drug repurposing framework to coronary artery disease (CAD) cohorts of millions of patients and emulate RCTs for multiple drug candidates, estimating their effects on CAD progression outcomes.

In general, our contributions are three folds:

• We develop a framework for high-throughput screening of on-marked drugs by emulating, for each drug, an RCT that evaluates its beneficial effect. The repurposed drug candidates can be proactively generated on existing large-scale RWDs.
• We present an innovative study design for the estimation of the drug’s effect from longitudinal observational data. The study CAD cohorts are automatically derived under our framework, which accelerates the process of computational drug repurposing.
• We propose a deep learning based propensity score estimation model to correct for confounding and selection
biases. Experimental comparisons to the logistic regression based propensity score estimation model show that our proposed deep learning model effectively estimate drug effects from RWDs, paving the way for drug repurposing.

Overall framework

We develop a high throughput computational drug repurposing pipeline (Fig. 1) that, given a disease cohort (i.e., CAD patients) extracts a list of potential repurposing drug ingredients and, for each, identifies the corresponding user and non-user sub-cohorts. It then computes, for all patients in both sub-cohorts, a large number of features (confounding factors), as well as the disease progression outcomes. The treatment effects are estimated after correcting for confounding and selection biases using the deep learning framework (Fig. 2). Here, the proposed framework is equipped with attention mechanism that provides the interpretability of the model. The drug ingredients with beneficial effect and statistical significance will be considered as repurposed drug candidates and suggested to be used for treating CAD. Algorithm 1 overviews the steps of estimating the effect of assigned treatment on the outcome from observational data.

![Figure 1. Flowchart of overall drug repurposing framework.](image)

**Figure 2.** Illustration of the deep learning model for predicting treatment probability (a.k.a., propensity score) that used for correcting confounding from temporal time sequence data (including diagnoses $d_t$, prescriptions $p_t$, and demographics $b_t$). It consists of three main components: embedding module, recurrent neural network and prediction module.

**Figure 2.** Illustration of the deep learning model for predicting treatment probability (a.k.a., propensity score) that used for correcting confounding from temporal time sequence data (including diagnoses $d_t$, prescriptions $p_t$, and demographics $b_t$). It consists of three main components: embedding module, recurrent neural network and prediction module.

**Results**

In this section, we first introduce the dataset we use for this study. Then we demonstrate the performance of our model in CAD drug repurposing experiments. We identify more than 90 million patients in MarketScan [17] data from 2012 to 2017, which contain individual-level, de-identified healthcare claims information from employers, health plans, hospitals, Medicare, and Medicaid programs. MarketScan claims data is primarily used for evaluating health utilization and services. The overall patients’ distribution of the recording period is shown in Fig. 3(a). We take both inpatient and outpatient claims into consideration. CAD cohort criteria are defined using International Classification of Diseases (ICD) codes [18] (Supplementary Table 1 for definitions). In total, there are 1,178,997 CAD patients. We refer to the first date when patients were determined to have CAD as CAD initiation date. Figure 3(b) shows the patients distribution of time before/after CAD initiation date.

**Dataset**

We identify three categories of study variables: demographic characteristics, diagnosis codes and prescription medication. Demographic characteristics in MarketScan CAD data include information on age and gender for each patient. Figure 3(d) shows the age and gender statistics and distribution of our dataset. Because a majority of data come from commercial claims, race and ethnicity information is incomplete and is not included in the analysis. Diagnosis codes in MarketScan CAD data are defined using the ICD codes for billing purposes. There are 57,089 ICD-9/10 codes considered in the dataset. Prescription medications in MarketScan CAD data also contain all prescription drug claims which contain prescription drug name (generic and brand), national drug code (NDC), and the number of days of supply approved. By matching NDC to observational medical outcomes partnership (OMOP) ingredient concept ID [19], we get 1,353 unique drugs in the dataset for drug repositioning screening. For drugs with multiple ingredients, we consider each active ingredient separately in the mapping processes.
**Algorithm 1** The algorithmic framework to estimate effect of assigned treatment on the outcomes

**Input:** patient data: assigned treatment, outcomes, values for potential confounders

**Output:** repurposed drug candidates, and their estimated effect, unbalanced feature ratio and significance

1. Generate user and non-user sub-cohorts for the treatment
2. Compute balancing weights for all patients in both sub-cohorts via LSTM based IPTW
3. Estimate the effect over multiple outcomes after correcting for the biases in the confounders (Eq. (1))
4. Compute the unbalanced feature ratio for the treatment after re-weighting using standardized difference (Eq. (2))
5. Estimate the significance of effect and compute adjusted p-value using bootstrapping
6. if estimated effect < 0 and adjusted p-value < 0.05 and unbalanced feature ratio < 2% then
7. return the estimated effect, unbalanced feature ratio and computed p-value
8. end if

To evaluate the drug effect, we define a set of clinically-relevant events linked the CAD as the disease outcomes (e.g., heart failure onset and stroke onset) after consulting domain experts. The definition is based on the ICD codes and can be found in Table 2 and Table 3 of Supplementary Materials. Since CAD is the major risk factor for both heart failure [20] and stroke [21], we hypothesize that an effective drug will lower the risks of CAD patients develop those diseases. Figure 3(c) demonstrates the time to develop outcomes from the CAD initiation date. The confounding variables affect both treatment assignment of patients and an outcome used in the trial. We consult domain experts to compile a list of hypothesized confounders for the CAD case study with respect to the study variables illustrate above: demographics, co-morbidities (diagnosis codes) and co-prescribed drugs.

**Model performance**

**Evaluation metrics**

**Treatment effect estimation** In this study, we leverage average treatment effect (ATE) to examine the treatment effect at the population level, which is defined as

\[
\text{ATE} = \mathbb{E}(Y_1) - \mathbb{E}(Y_0)
\]  

(1)

where \(\mathbb{E}(Y_1)\) and \(\mathbb{E}(Y_0)\) are the expected potential treated and control outcome of the whole population respectively. The values of ATE are used to determine whether the given treatment can improve disease outcomes or not.

**Testing feature balance** We evaluate the performance of models by measuring features' balance between the weighted user and non-user sub-cohorts generated by the IPTW. Given patient weights from IPTW, we quantify the balance for each feature using its standardized mean difference (SMD), which is the difference in the variable means between the two treatment groups, divided by the combined standard deviation. To be exact, we use the following definition of the standardized difference,

\[
\text{SMD} = \frac{|\mu_{user} - \mu_{nonuser}|}{\sqrt{(s^2_{user} + s^2_{nonuser})/2}}
\]  

(2)

where \(\mu_{user}\) and \(\mu_{nonuser}\) are the mean in user cohort and nonuser cohort; \(s^2_{user}\) and \(s^2_{nonuser}\) are sample variance of variables in two sub-cohorts. For binary variables, the variance \(s^2\) is calculated by \(\mu(1 - \mu)\). We consider a standardized difference greater than 0.1 as unbalanced [22] and compute the unbalanced feature ratio (i.e., \(#\text{unbalanced features} \big/ \#\text{all features}\) before/after weighting to evaluate the performance of balancing. The user and non-user sub-cohorts are considered as balanced if their unbalanced feature ratio is below 2% after weighting.

**Confidence intervals and significance of effect** We use the bootstrapping method [23] to calculate the confidence intervals of estimators of \(\mathbb{E}(Y_1)\) and \(\mathbb{E}(Y_0)\), and statistical significance of ATE. For each candidate ingredient, we repeatedly generate multiple different control drugs via random sampling with replacement, and the analysis is repeated in each bootstrap sample. The 95% confidence interval is then computed by using the standard normal approximation: \(\pm 1.96 \times \text{standard error}\) estimate of the standard error. The p-value of the effect estimator can be computed by the normal cumulative distribution function of estimators. We further use adjusted p-value [24] as a statistically significant measurement. We consider a repurposing drug candidate as significant if its adjusted p-value is below 0.05.

**Performance over repurposing drug candidates**

We identify 55 ingredients as drug repurposing candidates following the study design (see Methods). Then we estimate the treatment effect on various disease outcomes (i.e., heart failure and stroke) and demonstrate the distribution of ATE in Fig. 4. Here, we only show the drug candidates with the balanced user and non-user sub-cohorts after re-weighting and statistically significant estimates (adjusted p-value). All the drugs are ranked from the left side to the right side according to the increasing order of estimated ATE values. Based on the definition of ATE (i.e., the weighted average of observed outcomes from the user and non-user sub-cohorts), the drug ingredients with ATE values that smaller than 0 are identified to improve the disease outcomes, while the drug ingredients with ATE values larger than 0 are identified to worsen the disease outcomes. For the drugs with beneficial effects, we color those with known CAD indication in red and those without known CAD indication in blue (The drug label information is collected from SIDER [25] database and DrugBank [26]).

From the results, we observe that 9 drugs yield a beneficial effect on disease outcomes among 16 selected significant drug candidates. Specifically, only 3 drugs have been used as CAD indication according to their drug labels information. The remaining 6 drugs which have not been indicated
for treating CAD but can improve the disease outcomes are considered as repurposed drug candidates. We find evidence support for these 6 drug candidates from related literature and web resources as follows: (1) Metoprolol is one of the most commonly used beta-blockers for treating high blood pressure and chest pain. It shows beneficial effects in patients with heart failure associated with CAD [27]; (2) Fenofibrate is mainly used to treat abnormal blood lipid levels and also appears to decrease the risk of CAD in patients with diabetes mellitus [28]; (3) Hydrochlorothiazide which is often used to treat high blood pressure and diabetes insipidus [29], has already completed phase 4 trials for CAD treatment [30]; (4) Pravastatin has also been studied to have a beneficial effect on CAD [31]; (5) For simvastatin, results from randomized clinical trials show that it can reduce the occurrence of heart failure in patients with CAD [32]; (6) Valsartan, a kind of angiotensin receptor blocker, results in improved coronary micro-vascular flow reserve, suggesting direct beneficial in hypertensive patients with stable CAD [33].

We further list the sub-cohort size, feature balancing and estimated ATE values for each drug candidate in Table 1. The results of all 55 drugs can be found in Table 4 of Supplementary Materials. The first column lists the drug names corresponding to drugs in Fig. 4. The second and third columns denote the number of patients in user and non-user
Table 1. Sub-cohorts size, feature balancing and estimated effects for CAD over balanced and statistically significant drug ingredients. Bold in the table denotes the ingredients without known CAD indication (repurposed drug candidates).

| # drug_name     | # user | # non-user | pre.unbalanced covariates | post.unbalanced covariates | # covariates | post.unbalanced.ratio % | pre.ATE     | post.ATE    |
|-----------------|--------|------------|---------------------------|-----------------------------|--------------|--------------------------|-------------|-------------|
| metoprolol      | 9730   | 29190      | 38.308                    | 23.231                      | 1270         | 1.8                      | -0.023      | -0.043      |
| fenofibrate     | 1352   | 4056       | 39.340                    | 13.200                      | 1038         | 1.3                      | -0.051      | -0.038      |
| rosuvastatin    | 2420   | 7260       | 24.020                    | 9.620                       | 1097         | 0.9                      | -0.063      | -0.030      |
| hydrochlorothiazide | 2001   | 6003       | 32.500                    | 15.320                      | 1180         | 0.7                      | -0.050      | -0.026      |
| amlodipine      | 4613   | 13839      | 21.340                    | 8.300                       | 1154         | 0.7                      | -0.050      | -0.026      |
| pravastatin     | 2007   | 6021       | 11.260                    | 9.640                       | 1085         | 0.9                      | -0.016      | -0.022      |
| simvastatin     | 1605   | 4815       | 10.060                    | 13.240                      | 1044         | 1.3                      | -0.032      | -0.020      |
| valsartan       | 1316   | 3948       | 24.940                    | 13.740                      | 1026         | 1.3                      | 0.010       | -0.015      |
| amlodipine      | 1044   | 3132       | 28.360                    | 13.080                      | 1007         | 1.3                      | -0.010      | -0.013      |
| isosorbide      | 1482   | 4446       | 33.320                    | 9.560                       | 1039         | 0.9                      | 0.045       | 0.034       |
| prasugrel       | 1316   | 3948       | 41.500                    | 18.340                      | 1019         | 1.8                      | -0.043      | 0.036       |
| ramipril        | 887    | 2661       | 25.340                    | 14.840                      | 973          | 1.5                      | 0.020       | 0.043       |
| potassium chloride | 1110  | 3330       | 43.460                    | 20.240                      | 1016         | 2.0                      | 0.169       | 0.090       |
| carvedilol      | 3959   | 11877      | 38.280                    | 8.140                       | 1154         | 0.7                      | 0.198       | 0.124       |
| furosemide      | 1545   | 4635       | 50.880                    | 17.080                      | 1064         | 1.6                      | 0.301       | 0.179       |
| spironolactone  | 1292   | 3876       | 70.620                    | 12.920                      | 1034         | 1.3                      | 0.393       | 0.190       |

sub-cohorts, respectively. The next two columns denote the average number of unbalanced covariates before and after re-weighting. The "post unbalanced ratio" column represents the percentage of unbalanced covariates after re-weighting (i.e., the number of unbalanced covariates divided by the total number of covariates). And the last two columns are the estimated ATE before and after re-weighting. We rank the drugs by increasing of re-weighted ATE values. We see that our proposed method successfully corrects for most biases in the original data which results in a decrease in the number of unbalanced covariates.

**Case studies: attention visualization**

Having presented that our model successfully identified repurposed drug candidates for CAD treatment, we further demonstrate the interpretability of our framework achieves via attention mechanism. To exemplify this, we select two case drug candidates: diltiazem and fenofibrate. According to Table. 1, diltiazem and fenofibrate both have beneficial effect on CAD disease outcomes. Diltiazem has already been used for treating CAD [34], while fenofibrate does not have CAD indication on its drug label.

We want to identify the covariates that significantly biased between the user and non-user cohorts in original data but balanced after re-weighting. The learned attention weights enable visualization of each covariate and its SMD values before/after balancing between user and non-user cohorts. We select the top 20 well-balanced (i.e., large deviations of SMD during balancing) covariates and plot the distribution of SMD values for two case drugs in Fig. 5. The original unweighted data are denoted in blue dots and LSTM weighted data are in orange dots. The covariates are ordered from bottom to top according to the increase of differences between SMD values of unweighted data and LSTM weighted data. According to the figure, we see that for both two drugs, the SMD values in original data are greater than 0.1 (i.e., the threshold of balancing), which indicates that the original observational data is highly biased and exists much confounding variables. The maximum SMD value is about 0.6 for diltiazem and 0.35 for fenofibrate. While the SMD values estimated in the LSTM weighted data are smaller than 0.1, which means that no major biases between user and non-user cohort in terms of selected covariates. The selected covariates include demographics (e.g., age), co-prescribed drugs (metformin, metoprolol, etc.) and co-morbidities (e.g., acute myocardial infarction, cardiac dysrhythmias, etc.). After correcting for these confounding variables, we can have a more accurate estimation of the treatment effect on the diseases.

**Discussion**

In this study, we present a computational drug repurposing framework for high-throughput screening of on-marked drugs by emulating a corresponding RCT for each drug and evaluating its treatment effect on various disease outcomes. We propose a deep learning based propensity score model for correcting selection biases and confounding in longitudinal observational data. We demonstrate our framework in a case study of CAD and evaluate 55 different repurposing drug candidates on two disease outcomes. According to the results, we obtain 6 drug candidates (i.e., metoprolol, fenofibrate, hydrochlorothiazide, pravastatin, simvastatin, valsartan) that improve the CAD outcomes with statistical significance but have not been indicated for treating CAD.
Figure 5. The SMD values of top 20 well balanced covariates. Fig. 5(a) shows results of diltiazem. Fig. 5(b) shows results of fenofibrate.

We also develop a base version of our model that uses logistic regression (LR) for computing propensity score and treatment effect estimation. We conduct comparison experiments on the base model (LR-IPTW) and our model (LSTM-IPTW) on the above two case drugs and show the results for diltiazem in Fig. 6 (The results for fenofibrate can be found in Supplementary Materials).

As the feature balancing is one of the most important evaluation metrics, we first plot the distribution of absolute SMD values computed by LSTM-IPTW and LR-IPTW in Fig. 6(a) and Fig. 6(d). In both LSTM weighted data and LR weighted data, many features exhibit large absolute SMD values (greater than 0.1) in the original data, while most features exhibit low absolute SMD (below 0.1) after re-weighting. Specifically, less features exhibit absolute SMD values above 0.1 threshold after weighted by LSTM model than weighted by LR model. This indicates that the data is well-balanced by LSTM-IPTW and the estimated ATE from LSTM-IPTW should be more accurate than LR-IPTW. Figure 6(b) and Figure 6(e) show the propensity distribution plot over user and non-user cohorts using LSTM-IPTW and LR-IPTW models. We observe that the propensity distribution of LSTM-IPTW is more smooth (i.e., the propensities are normally distributed) than the distribution of LR-IPTW. Under LR-IPTW model, many of the patients in non-user cohorts are predicted to have a propensity of 0. We also evaluated our models using conventional metrics. The ROC curve is a standard metric widely used to estimate the performance of prediction models. The area under ROC curve (AUC) characterize the accuracy of the prediction results. Figure 6(c) and Figure 6(f) show the ROC curves for LSTM-IPTW model and LR-IPTW model. The "Propensity" curves in the figures are the standard ROC curves of LSTM.
model and LR model. By comparing the AUC values of two models, we see that LSTM model yields more accurate prediction results than the LR model. With the accurate treatment predictions, the model would generate better weights for balancing and treatment effect estimates in the following tasks. Besides the standard ROC curve, we also show another two curves: weighted propensity curve and expected curve. The weighted propensity curve is obtained by re-weighting the standard ROC curve using the weights drawn from the propensity model (the same weights applied in covariates balancing and effect estimates). This curve should be very close to the curve that would arise by a random assignment (i.e., with an AUC close to 0.5) which indicates our assumption that the weighting can emulate an RCT. From the plots, we can find that LSTM-IPTW performs better than LR-IPTW in terms of more close value to 0.5. Compared to the standard propensity ROC curve, "Expected" ROC curve duplicates the population and assign weights to each individual based on the propensity.

In this setting, each patient contributes their propensity to the true positives and\(1 - \text{propensity}\) to the false positives. The standard propensity ROC curve should be close to the expected propensity ROC. We observe that LSTM-IPTW’s "Propensity" curve is much close to its "Expected" curve when compared to LR-ITPW.

This study can be extended in multiple directions in the future. For the study, we use the hypothesized confounders including demographics, co-morbidities and co-prescribed drugs. Some other potential confounders such as time elapsed from the first disease diagnosis to index date and outcome value calculated over the baseline period could be considered to build the model in the future work. Also, we can consider drug combinations and estimate the effect of entire combinations on the disease outcomes.

In summary, we demonstrate that the proposed computational drug repurposing framework can successfully identify drug candidates that have a beneficial effect on disease out-
comes but not have been indicated for CAD patients yet. The new LSTM-IPTW model shows better performance for correcting biases and estimating treatment effect than LR-IPTW, and remaining the interpretability for recognizing significant confounding.

**Methods**

In this section, we first introduce the study design, which includes definitions of cohorts and study variables. Then we illustrate our deep learning model with three main components in detail.

**Study design**

Our framework identifies drug repurposing candidates using MarketScan CAD data to emulate a bulk of corresponding RCTs. Below we describe the design of the emulated trials and the key components of our framework for CAD drug repurposing.

**User and non-user cohorts**

Given the drug tested in trial, a patient is assigned to the user cohort if the following inclusion criteria are satisfied: (1) the patient has persistently prescribed the drug (e.g., the interval between two prescriptions is less than 30 days); (2) the patient is eligible for trial at the time of the first prescription for the drug. In the CAD study, this condition is that the first prescription is after CAD initiation data; (3) the patient had a history in the database of at least one year (365 days) prior to the first prescription of the drug.

Estimating the effect of a drug requires comparing the user cohort to a control group assigned with alternative drugs. Once the alternative drugs are determined, the framework further excludes from the non-user cohort any patient prescribed with the trial’s drug. In our study design, alternative drugs are selected randomly from the prescribed ingredients, excluding the trial drug itself. Such a control group directly compares the trial’s drug to drugs of the same therapeutic indication, reducing confounding by indication. We use the term index-date to refer to the date of the first prescription of the assigned drug, that is, the first time the trial’s drug (respectively, the alternative drug) was prescribed for patients in the user (respectively, non-user) cohort.

**Baseline and follow-up periods**

We refer to the time period prior to the index-date for which we have information on the patient as the baseline period. We use the baseline period for characterizing the patients prior to the beginning of the treatment with the assigned drug. The follow-up period starts at the index-date, that is, at the beginning of the treatment with the trial’s drug in the user cohort, and the control-drug in the non-user cohort. The effect of the drug is evaluated during the follow-up period. In the CAD study, the baseline period is at least 365 days, and the follow-up period is 2 years (730 days). Figure 7 demonstrates the definition of user and non-user cohorts.

**Outcomes and hypothesized confounders**

The effect of the drug during the follow-up period is defined with respect to various disease outcomes. In this CAD drug repurposing case study, we consult domain experts to define a set of clinically-relevant events linked with CAD as the outcome, e.g., heart failure onset (Supplemental Table 2) and stroke onset (Supplemental Table 3). The treatment effect is estimated on these outcomes during the follow-up period (i.e., 730 days after index-date). The patient is considered to have the disease outcome if either of them happens in follow-up period.

Confounders are variables affecting both treatment assignment of patients and an outcome used in the trial, thus creating a "backdoor path" that may hinder the true effect of the drug on the outcome. We consult domain experts to compile a list of hypothesized confounders for the CAD study, including demographics (e.g., age at the index date, sex), co-morbidities (e.g., indicator per each ICD-9/10 diagnosis class) and co-prescribed drugs. Since confounders affect treatment assignment, they are computed on the baseline period.

**Repurposing drug ingredients**

We regard a drug as a repurposing candidate if it satisfies the following condition: (1) an active ingredient (i.e., the ingredient directly involved in achieving the mediation objectives). (2) persistently prescribed to a large enough number of patients in the disease cohort. Specifically, an ingredient is considered being used by a patient only if it was prescribed in two or more distinct dates, as least one month apart. And a minimum of 500 patients prescribed a certain ingredient was required. For each repurposing candidate, we can compute the user and non-user cohorts according to the above definition of cohorts. After obtaining the corresponding user and non-user cohorts, we can extract outcomes and hypothesized confounders for each individual patient from the database. Every patient in their sub-cohort is represented by a sequence of events, with each event providing the patient information (i.e., co-morbidities, co-prescribed drugs, etc.) that corresponds to each visit. The available data within these visits during the baseline period, combined with demographic characteristics
We define average treatment effect (ATE) of a drug on the outcome \( Y \) with treatment \( \alpha \) as \( \text{ATE} = \text{E}(Y_1) - \text{E}(Y_0) \), where \( \text{E}(Y_\alpha) \) denotes the potential expected prevalence of patients that would have experienced an outcome event during a complete follow-up period if all patients in the trial had been assigned with treatment \( \alpha \). The potential outcomes are referred as counterfactual as only one of these is observed for any given individual. By running RCT, we can measure the outcomes within user and non-user groups into which individuals are randomly assigned: \( \text{E}(Y_1) \) can be directly estimated as \( \text{E}(Y|\alpha = 1) \) and \( \text{E}(Y_0) \) as \( \text{E}(Y|\alpha = 0) \). However, in observational data (e.g., our MarketScan CAD data), treatment assignment is usually far from being random, which may depend on confounders (affecting both treatment assignment and outcome). We need to assign weights to the individuals in each group to avoid the influence of confounders.

In order to control the influence of confounders, we apply inverse probability of treatment weighting (IPTW) to create a pseudo-population from the original one by assigning a weight \( w_i^r \) to an individual \( i \) with treatment \( \alpha \). The weight is defined as the inverse of conditional probability (aka propensity score) that an individual is treated with \( \alpha \) given the confounding values. One common issue with IPTW is that individuals with a propensity score very close to 0 will end up with an extremely high weight, potentially making the weighted estimator unstable. We address this problem by adopting an alternative weighting function called standardized IPTW [22], which uses the marginal probability of treatment instead of 1 in the weight numerator.

For estimating the propensity score, logistic regression is the most popular method in statistics [35, 36]. In longitudinal observational data, those observational covariates are not a set of static feature vectors (one for each patient), but irregularly sampled time series (recording diagnoses, medications, etc. at each timestamp). Thus, logistic regression is not ideal for effectively modeling longitudinal observational data.

Model for propensity score weighting

The schematic view of our model is shown in Figure 2, which consists of three main components: embedding module, recurrent neural network and prediction module. Briefly, the model estimates the propensity score by first transforming the input features using an embedding layer. These embedded features are then fed into LSTM, the output of which at every time point is aggregated through an attention layer for automatically focusing on important time points. The aggregated features are fed into a prediction module that provides the probability of receiving treatment. Each of these is discussed below in detail.

Embedding module

The embedding module is to convert the initial high-dimensional and sparse input features into a lower-dimensional and continuous data representations, which is beneficial to the following prediction task. As shown in Fig. 2, the input features are consist of three components: diagnosis, prescription and demographic information (age and gender). The diagnosis codes for each patient at each timestamp can be denoted as \( \{d_1, d_2, ..., d_t\} \), and prescription can be denoted as \( \{p_1, p_2, ..., p_t\} \). Here, \( d_i \) and \( p_i \) are both one dimensional binary vector with the size of diagnosis code dictionary \( r \) and prescription code dictionary \( s \), respectively. For each element in the vector, the value one in the \( j \)-th column indicates that code \( j \) is documented in \( t \)-th visit. We use two linear embedding modules to represent diagnosis and prescription respectively. That is, we define \( e_i = W_{emb}^d d_i, f_i = W_{emb}^p p_i \), where \( e_i \in \mathbb{R}^r \) denotes the embedding of the input vector \( d_i \in \mathbb{R}^r, m \) the size of the diagnosis embedding dimension, \( W_{emb}^d \in \mathbb{R}^{m \times r} \) the embedding matrix. \( f_i \in \mathbb{R}^s \) denotes the embedding of the input vector \( p_i \in \mathbb{R}^s, n \) the size of the diagnosis embedding dimension, \( W_{emb}^p \in \mathbb{R}^{n \times s} \) the embedding matrix. The age is normalized into range of \([0,1]\) using min-max normalization and the gender is represented as a binary vector. Having the embedded vectors of patients, we input them to LSTM.

Recurrent neural network and Attention mechanism

Long short-term memory (LSTM) [15], which is a kind of recurrent neural network (RNN) equipped with memory cells, can better model temporality of observational data. LSTM and its variations are widely adopted in the scenario that contains sequential and temporal data, such as in language translation [37], speech recognition [38] and image captioning [39]. A common LSTM unit contains a cell, an input gate, an output gate and a forget gate. The cell can remember values over irregular time intervals and the three gates moderate the flow of information into and out of the cell. The inputs to the LSTM are embedded confounding vectors from the embedding module and the output of which is patient’s latent health status at the time of visit. We use two LSTMs, LSTM\(_\alpha\) and LSTM\(_\beta\) to separately model diagnosis and prescription codes of patients.

\[
\begin{align*}
  h_1, h_2, ..., h_t &= \text{LSTM}_\alpha(e_1, e_2, ..., e_t) \\
  g_1, g_2, ..., g_t &= \text{LSTM}_\beta(f_1, f_2, ..., f_t)
\end{align*}
\]  

where \( h_i \in \mathbb{R}^r, g_i \in \mathbb{R}^r \) are hidden state vectors at \( t \)-th visit, and \( u, v \) denote the size of hidden layer of LSTM\(_\alpha\) and LSTM\(_\beta\). Then those patient hidden states are aggregated through two separate attention layers for automatically focusing on impor-
where $y_c \in \mathbb{R}^n$, $b_a \in \mathbb{R}^r$, $W_\beta \in \mathbb{R}^r$ and $b_\beta \in \mathbb{R}^r$ are the parameters to learn. Using the generated attention weights for the aggregated patient states from attention layer $c_\alpha$ and $c_\beta$ with vectorized age and gender to predict the probability of receiving a treatment (propensity score).

### Prediction module

The aggregated patient states from attention layer $c_\alpha$, $c_\beta$, combined with the demographic features $c_{demo}$, are passed through a fully-connected neural network predict the probability of receiving a treatment as follows,

$$
\hat{y} = \text{Sigmoid}(W^T c_i + b)
$$

(5)

where $c_i = \text{ReLU}(W_c[c_\alpha, c_\beta, c_{demo}]) + b_c$, $W_c \in \mathbb{R}^{k \times (n+r+2)}$, $b_c \in \mathbb{R}^k$, $W \in \mathbb{R}^k$, $b \in \mathbb{R}$ are the model parameters. We use cross-entropy to calculate the prediction loss as follows,

$$
\mathcal{L} = -\frac{1}{N} \sum_{i=1}^{N} (y_i \log \hat{y}_i + (1 - y_i) \log (1 - \hat{y}_i))
$$

(6)

where $y_i$ is the ground truth of observed treatment for patient $i$.

### Experiment settings

The model is implemented and trained with Python 3.6 and PyTorch 1.4, on a high-performance computing cluster with four NVIDIA TITAN RTX 6000 GPUs. For each drug candidate, we train a model using the adaptive moment estimation (Adam) algorithm with a batch size of 50 subjects and the learning rate is 0.001. We run each model for 50 iterations for computing p-value and confidence interval. We randomly split the input data into training, validation and test sets with a ratio of 70%, 10%, 20%. The information from a given patient is only present in one set. The training set is to train the proposed models. The validation set is used to improve the models and select the best model hyperparameters.

### Data availability

The access of the MarketScan data analyzed in this manuscript is provided by The Ohio State University. The dataset may be available from IBM at https://www.ibm.com/products/marketscan-research-databases.

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1https://pytorch.org/

### Code availability

The code for this paper is available at https://github.com/ruoqi-liu/DeepIPW

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Author contributions
PZ conceived the project. RL and PZ developed the method. RL conducted the experiments. RL, LW and PZ analyzed the results. RL, LW and PZ wrote the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplemental Table 1. The definition of coronary artery disease (CAD) from observational health data. Supplemental Table 2-3. The definition of heart failure and stroke from observational health data. Supplemental Table 4. Main results for all 55 repurposing drug candidates. Supplemental Figure 1. Performance comparison of LSTM-IPTW and LR-IPTW on case drug: fenofibrate.