Estimating the effects of second-line therapy for type 2 diabetes mellitus: retrospective cohort study

Assaf Gottlieb, Chen Yanover, Amos Cahan, Yaara Goldschmidt

ABSTRACT

Objective Metformin is the recommended initial drug treatment in type 2 diabetes mellitus, but there is no clearly preferred choice for an additional drug when indicated. We compare the counterfactual drug effectiveness in lowering glycated hemoglobin (HbA1c) levels and effect on body mass index (BMI) of four diabetes second-line drug classes using electronic health records.


Participants and exposures Our cohort consisted of more than 40 000 patients with type 2 diabetes, prescribed metformin along with a drug out of four second-line drug classes—sulfonylureas, thiazolidinediones, dipeptidyl peptidase 4 (DPP-4) inhibitors and glucagon-like peptide-1 agonists—during the years 2000–2015. Roughly, 17 000 of these patients were followed for 12 months after being prescribed a second-line drug.

Main outcome measures HbA1c and BMI of these patients after 6 and 12 months following treatment.

Results We demonstrate that all four drug classes reduce HbA1c levels, but the effect of sulfonylureas after 6 and 12 months of treatment is less pronounced compared with other classes. We also estimate that DPP-4 inhibitors decrease body weight significantly more than sulfonylureas and thiazolidinediones.

Conclusion Our results are in line with current knowledge on second-line drug effectiveness and effect on BMI. They demonstrate that causal inference from electronic health records is an effective way for conducting multtreatment causal inference studies.

INTRODUCTION

Type 2 diabetes mellitus (T2DM) affects more than 29 million people in the USA and is the seventh leading cause of death.1 2 The American Diabetes Association Standards of Medical Care,3 supported by several studies,4 5 recommends dietary changes and physical exercise as the initial treatment, followed by administration of metformin if lifestyle changes fail to reach glycemic control. According to the Standards of Medical Care, if metformin does not achieve glycemic target within 3 months, one of the following six second-line medications should be added: sulfonylureas (SU), thiazolidinediones (TZD), inhibitors of dipeptidyl peptidase 4 (DPP-4), glucagon-like peptide-1 receptor agonists (GLP-1), sodium-glucose cotransporter 2 (SGLT2) inhibitors or insulin. Currently, the guidelines do not prefer one class over the others. The effectiveness, costs and risk of complication of those drug classes were compared in clinical trials6 and meta-analyses of their results.7–9

Significance of this study

What is already known about this subject?

► The effects of type 2 diabetes second-line drugs on glycosylated hemoglobin levels and on body mass index (BMI) have been evaluated in clinical studies. However, the clinical implication of these studies is limited by small number of participating individuals and the homogeneity of the study populations.

► Meta-analysis studies have increased sample size but potentially suffer from similar homogeneity biases.

What are the new findings?

► This study performs, for the first time, a large-scale analysis of the therapeutic and adverse effects of type 2 diabetes second-line drugs in real-world population using electronic health records.

► We confirm current knowledge for glycosylated hemoglobin levels, while showing better effects on decreasing BMI for inhibitors of dipeptidyl peptidase 4 (DPP-4).

How might these results change the focus of research or clinical practice?

► Our results show that while sulfonylureas are the most commonly prescribed second-line drugs, their estimated reduction in glycosylated hemoglobin levels is significantly smaller than the estimated effects of thiazolidinediones, glucagon-like peptide-1 agonists or DPP-4 inhibitors. DPP-4 inhibitors also show significant reduction of BMI compared with sulfonylureas and thiazolidinediones.

► We demonstrated that causal inference methods can confirm and expand current knowledge in a cost-effective way and should gain increased focus when addressing epidemiological questions.
These comparisons found no significant difference in drug class effect on the percentage of blood glycated hemoglobin (HbA1c); thus, no specific recommendation about the choice of a second drug could be made. Notably, clinical trials are laborious and costly. Trials often include small samples with limited representativeness of the target population (e.g., between 2005 and 2012, the Food and Drug Administration approved drugs based on a median number of 2 clinical trials and the median number of patients enrolled was 760). Meta-analyses of clinical trials may have higher power and be more generalizable, but are also vulnerable to publication bias, small-study effects and limited degree of heterogeneity.

Electronic health records (EHRs) hold promise as an alternative approach to conduct causal inference experiments that can address some of these limitations. Specifically, secondary use of EHRs requires lower costs, can scale to a large number of patients and better represents the heterogeneity in the population. There are trade-offs to using the EHR approach, and such analyses may suffer from three major limitations: First, patients may get treatment outside the institutions included in the EHR, resulting in missing and fragmented data. Second, confounders play a crucial role in effect size estimation and their identification is challenging. Third, differences in protocols or adoption rate for new medications across institutions may obscure true effect size and might not be generalizable beyond the database from which they are derived.

Here, our aim is to compare the effects of T2DM second-line drugs using a real-world evidence approach. We emulate a multiarm clinical trial of four classes of drugs for diabetes, commonly used as second-line treatment (SU, TZD, inhibitors of DPP-4 and GLP-1). We compare the counterfactual (i.e., potential) effectiveness (in terms of HbA1c levels) and body mass index (BMI) outcomes of 17082 patients over the course of 12 months, adjusting for confounders and censoring (additional 23789 patients). For reference, a recent meta-analysis of antidiabetes drugs was based on data of about 18000 patients. We describe the measures we have taken to address the aforementioned limitations of causal inference from EHR data. Our results are in line with current knowledge, thus demonstrating that causal inference from EHRs is an effective way for conducting multitreatment causal inference studies.

**RESEARCH DESIGN AND METHODS**

**Study design**

**Data source**

We used the Explorys database (IBM), which includes EHR records of approximately 50 million patients, pooled from multiple different healthcare systems in the USA. Data consist of a combination of clinical EHRs, healthcare system outgoing bills and adjudicated payor claims, are standardized and normalized using common ontologies, including SNOMED and The National Drug File - Reference Terminology, and are searchable through a Health Insurance Portability and Accountability Act of 1996-enabled, de-identified database tools. The EHR data include patient demographics, diagnoses, procedures, prescribed drugs, vitals and laboratory values.

**Cohort definition**

We defined a cohort of patients with T2DM based on the Northwestern University diabetes phenotyping algorithm, comprising 40871 patients, using the following criteria:

**Inclusion criteria**

Our analysis considered T2DM patients, identified by having at least two types of evidence for T2DM, out of T2DM diagnosis, T2DM-specific drugs, and indicative lab values (fasting and random glucose or HbA1c levels). We included only patients who were first prescribed metformin and subsequently prescribed, during the years 2000–2015, a second-line drug belonging to any of four classes: SU, TZD, inhibitors of DPP-4 and GLP-1 agonists (online supplementary table 1 lists drugs for each drug class).

The first prescription with order status marked as completed (i.e., that it was not canceled or erroneous) of the second-line drug was considered the ‘index-date’ (emulating the date of intervention allocation and initiation in clinical trials).

We required the patients to have at least 12 months of documented pretreatment observation period prior to the index date.

**Exclusion criteria**

Patients with type 1 diabetes mellitus, identified by either a type 1 diabetes diagnosis code or prescription of pramlintide (approved also for patients with T2DM who use insulin), as well as patients prescribed more than one second-line drug classes on the index date were excluded from the analysis. Our analysis did not include the following three second-line medications: SGLT2 inhibitors, meglitinides and α-glucosidase inhibitors due to the low number of patients receiving it within our data (185, 429 and 153 patients with available HbA1c measurements, respectively). We did not compare insulin because it is not commonly considered as the first choice for second-line therapy in clinical practice; it is also administered to patients with more advanced or severe disease than oral agents; and patient acceptance of insulin often involves unique psychological and social factors that are not part of our cohort data, nor usually recorded in the EHR. These factors are likely to be important confounders and no analysis that uses this type of data could adjust for them.

**Outcomes and follow-up time**

As outcomes, we used HbA1c and BMI at two follow-up periods, 6 and 12 months after the index date. We averaged the HbA1c and BMI over ±3-month windows for each period (3–9 months for the first follow-up period...
Figure 1: Illustration of the causal inference scheme. (A) Index date is the first prescription of diabetes second-line drug after use of metformin. (B) Potential censoring events include switching to another second-line drug, missing glycated hemoglobin (HbA1c) or body mass index (BMI) measurement, or undergoing bariatric surgery. (C) Outcomes (HbA1c and BMI) are checked after 6 and 12 months from index date. (D) Follow-up ends after 15 months.

and 9–15 months for the second; figure 1). We chose a 12-month pretreatment observation period to ensure that the second-line drug is prescribed for the first time as, for example, 99% of patients receiving SU have prescription period <12 months. It also balances the need for a complete and stable baseline (ie, longer period) with the need to include more patients and avoid bias due to exclusion of patients with relatively limited histories in our data (ie, a shorter period). We chose follow-up periods of 6 and 12 months (averaging over ±3 month windows, resulting in 15 months in total) since they provide a good estimate of the short to intermediate effects of the drugs and correspond to the majority of random clinical trial follow-up time interval.5,8 We required each patient to have at least one HbA1c measurement during the pretreatment observation period.

Analysis methods
We considered two potential biases: (1) selection bias due to censoring; and (2) confounders, affecting both treatment choice and measured outcome (HbA1c levels or BMI).

In order to handle these two biases, we extracted patient characteristics within the pretreatment observation window using the feature engineering framework of Ozery-Flato et al.19 The comprehensive set of features included demographic information (age, sex, ethnicity), insurance type, patient-aggregated diagnoses using Clinical Classifications Software categories, categories of Charlson20 and Elixhauser comorbidity indexes21 prescribed drugs (active ingredients), and laboratory results values over the baseline period. Diagnosis codes and drugs are binary features (measuring the existence of a diagnosis or a drug prescription for that patient). Categorical features, such as insurance type or ethnicity, were split into binary features. For lab values, we included the number of times a lab value was measured within the baseline period, and the maximal, minimal and average values within that period. For the HbA1c outcome, we also included the last measurement before treatment and the time from first diabetes diagnosis (based on the Northwestern University diabetes phenotyping algorithm). As a preliminary step, we filtered features that were dominated (>95% of patients) by a single value or were spurious (>80% with missing values), resulting in 632 features. Missing lab values were imputed using the median value of the test across patients.

Censoring analysis
For both HbA1c and BMI inference, we considered patients as censored if they received a second-line treatment, but during the follow-up period (1) had no 6-month or 12-month HbA1c or BMI measurements; (2) switched or added another antidiabetic drug (including the following drug classes, which were not directly evaluated in this work: insulin, SGLT2 inhibitors, meglitinides and α-glucosidase inhibitors); or (3) underwent bariatric surgery. We corrected for censoring by reweighing the uncensored population using inverse probability of censoring weighting (IPCW)22.

Confounder analysis
We defined the set of confounders in two ways: (1) domain expert confounder set, manually selected by an internist, aided by literature search; and (2) a comprehensive confounder set, treating all the 632 extracted features as confounders. In total, we selected 34 domain expert confounders for HbA1c inference (online supplementary table 2).

Table 1 Descriptive statistics of patients on T2DM second-line drug classes for the HbA1c outcome

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Patients (n)</th>
<th>Treatment change*</th>
<th>Missing outcome*</th>
<th>Average age†</th>
<th>% Female*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfonylurea</td>
<td>26684</td>
<td>4336 (16%, 3e^{153})</td>
<td>12269 (46%, −)</td>
<td>61.2 (2e^{88})</td>
<td>47.7% (2e^{15})</td>
</tr>
<tr>
<td>Thiazolidinedione</td>
<td>4794</td>
<td>1145 (24%, 2e^{12})</td>
<td>2235 (47%, −)</td>
<td>59.6 (0.001)</td>
<td>48.2% (−)</td>
</tr>
<tr>
<td>Glucagon-like peptide-1 receptor agonists</td>
<td>1532</td>
<td>398 (26%, 4e^{9})</td>
<td>735 (48%, −)</td>
<td>52.8 (5e^{−113})</td>
<td>66.6% (3e^{−44})</td>
</tr>
<tr>
<td>Dipeptidyl peptidase 4</td>
<td>7861</td>
<td>2314 (29%, 3e^{118})</td>
<td>3405 (43%, 5e^{−6})</td>
<td>58.9 (2e^{−32})</td>
<td>51.1% (8e^{−5})</td>
</tr>
</tbody>
</table>

Per-confounder statistics appear in online supplementary table 2.
*Proportion test. Missing entries (−) are not significant with FDR <0.05.
†Wilcoxon rank-sum test. Missing entries (−) are not significant with FDR <0.05.
FDR, false discovery rate; HbA1c, glycated hemoglobin; T2DM, type 2 diabetes mellitus.
brevity. We obtained similar counterfactual lab values, into five categories to prevent introduction of stratification of continuous variables, for example, age and the class sizes. Next, similar to Gerhard regularization strength set to one and using balancing of the distribution of the counterfactual outcome (standardization) and a treatment assignment mechanism model (inverse probability of treatment weighting, IPTW). As demonstrated by Bang and Robins,24 DR estimators improve demonstration by Bang and Robins,24 DR estimators improve on either estimators because they are consistent even when only one of the models is correctly specified. This makes DR especially suited for observational data, where one can never be sure that either model is correct. For the outcome model, we used ridge regression with fivefold cross-validation to adjust the regularization coefficient. For the treatment model, we used multiclass logistic regression with fivefold cross-validation. Inference scheme to two negative controls30: patient community.

Based on Groenwold et al,28 we also tested the results of stratification of continuous variables, for example, age and lab values, into five categories to prevent introduction of residual confounding. We obtained similar counterfactual mean values to the non-categorized values but larger confidence intervals (CIs) and thus omitted these results for brevity.

As suggested by Austin,29 the standardized difference, $d$, can be used to quantify covariate imbalance across subject groups. Specifically, for continuous confounders: $$d = \frac{\bar{\chi}_{\text{treatment}} - \bar{\chi}_{\text{control}}}{\sqrt{\frac{S^2_{\text{treatment}} + S^2_{\text{control}}}{2}}}$$ where $\bar{\chi}_{\text{treatment}}$ and $\bar{\chi}_{\text{control}}$ denote the sample mean of the covariate in treated and untreated subjects, respectively, whereas $S^2_{\text{treatment}}$ and $S^2_{\text{control}}$ denote the sample variance of the covariate in treated and untreated subjects, respectively.

And, for dichotomous confounders:

$$d = \frac{\hat{P}_{\text{treatment}} - \hat{P}_{\text{control}}}{\sqrt{\frac{\hat{P}_{\text{treatment}}(1-\hat{P}_{\text{treatment}}) + \hat{P}_{\text{control}}(1-\hat{P}_{\text{control}})}{2}}}$$

where $\hat{P}_{\text{treatment}}$ and $\hat{P}_{\text{control}}$ denote the prevalence or mean of the dichotomous variable in treated and untreated subjects, respectively.

Per-confounder statistics appear in online supplementary table 3.

**Table 2** Descriptive statistics of patients on T2DM second-line drug classes for the BMI outcome

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Patients (n)</th>
<th>Treatment change*</th>
<th>Missing outcome*</th>
<th>Average age†</th>
<th>% Female*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfonylurea</td>
<td>18170</td>
<td>2967 (16%, 2e^{-10})</td>
<td>8611 (47%, 2e^{-16})</td>
<td>60.9 (4e^{-17})</td>
<td>48.4% (0.01)</td>
</tr>
<tr>
<td>Thiazolidinedione</td>
<td>2691</td>
<td>640 (23.8%, 3e^{-5})</td>
<td>1503 (56%, 2e^{-29})</td>
<td>59.2 (7e^{-25})</td>
<td>49.3% (−)</td>
</tr>
<tr>
<td>Glucagon-like peptide-1</td>
<td>1172</td>
<td>293 (25%, 5e^{-5})</td>
<td>441 (38%, 3e^{-5})</td>
<td>52.9 (3e^{-8})</td>
<td>66.6% (7e^{-34})</td>
</tr>
<tr>
<td>receptor agonists</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dipeptidyl peptidase 4</td>
<td>1672</td>
<td>1852 (29%, 3e^{-3})</td>
<td>2352 (37%, 2e^{-49})</td>
<td>58.7 (2e^{-29})</td>
<td>50.9% (0.003)</td>
</tr>
</tbody>
</table>

†Wilcoxon rank-sum test. Missing entries (−) are not significant with FDR <0.05.

*Proportion test. Missing entries (−) are not significant with FDR <0.05.

BMI, body mass index; FDR, false discovery rate; T2DM, type 2 diabetes mellitus.

No patients were involved in setting the research question or the outcome measures, nor were they involved in developing plans for design or implementation of the study. No patients were asked to advise on interpretation or writing up of results. There are no plans to disseminate the results of the research to study participants or the relevant patient community.

**RESULTS**

**Study design**

Our cohort included 40,871 patients. Of these, 28,328 also had available BMI before the prescription of second-line drugs and were used for inference of counterfactual BMI (tables 1 and 2, online supplementary figure 2–3). There were significantly more censored patients on TZD, GLP-1 or DPP-4 who switched or added another drug than patients on SU (censored patients, p<2e^{-100}; tables 1 and 2). TZD and SU had significantly higher percentage of patients with missing BMI measurements during the follow-up than GLP-1 and DPP-4 (p<3e^{-8}; table 2).
Finally, the patients on GLP-1 were about 6 years younger on average (p<0) and included significantly higher rate of women (p<3e−44; table 1). The patient age distribution (online supplementary figure 1) is similar to the age distribution published by the Centers for Disease Control and Prevention (CDC) for 2011.31

Analysis methods

We applied causal inference methods to compute the counterfactual HbA1c levels and BMI (for each one of the four drug classes) at each of the two follow-up time points, adjusting for censored patients and confounders (Research design and methods).

Our balancing test (methods) showed that the percentage of balanced confounders, with negligible difference between treatment groups (standardized difference ≤0.1), ranged between 87% and 97% (comprehensive set and domain expert set in BMI outcome, respectively); online supplementary figures 2–5 display scatter plots of the absolute standardized difference before and after the correction. We found no significant differences between patients on different drug classes when using negative controls of patient height, while finding differences of up to 0.08% in HbA1c levels before index date between GLP-1 and SU or TZD (see Discussion).

HbA1c outcome

HbA1c measurements were available for 83% of the patients from up to 90 days prior to initiation of second-line treatment, and for 95% of the patients up to 180 days (see online supplementary figure 6 for complete temporal distribution).

The differences in estimated HbA1c levels using the domain expert and comprehensive sets of confounders were lower than 0.03%. All four drug classes were predicted to reduce HbA1c levels below 7% after 12 months of treatment, with a predicted reduction in HbA1c levels relative to baseline over the entire population of 0.6%–0.61% (SU, domain expert and comprehensive set correction, respectively) to 0.85%–0.83% (GLP-1, domain expert and comprehensive set correction, respectively) (online supplementary table 5, figure 2). Twelve-month HbA1c levels inferred for SU were significantly higher than for TZDs, DPP-4 and GLP-1 by 0.09%–0.24% (Wald test, p<8e−3; online supplementary table 6). Inferred levels for DPP-4 were significantly higher than TZD after 12 months and higher than GLP-1 after 6 months of treatment, but differences became insignificant after 12 months (online supplementary table 6). Notably, both actual and inferred HbA1c levels were lower than those computed using the mixed-treatment comparison (MTC) of clinical trials of McIntosh et al.7 8

BMI outcome

BMI measurements were available for 78% and 83% of the patients as recent as 90 and 180 days prior to treatment date, respectively (see online supplementary figure 7 for complete time distribution).

The predicted BMI after 12 months was significantly lower for patients on DPP-4 than for patients on SUs or TZDs by 0.47–0.81 kg/m² (Wald test, p<8e−4; figure 3, online supplementary figure 7–8), but not significantly lower than the predicted BMI for patients on GLP-1. On average, patients on GLP-1 had higher BMI (by 4.2–4.4 kg/m²; online supplementary figure 7) before the prescription of second-line treatment compared with patients prescribed one of the other studied drug classes. These GLP-1 patients had lowered their BMI by 0.87 kg/m² on average. Our predicted BMI shows significant advantage for GLP-1 over TZD based on the domain expert confounder set, but not statistically significant based on the comprehensive confounder set. It also shows no significant advantage over SU in a population with lower initial BMI (online supplementary supplementary table 8; see also discussion on BMI and GLP-1).

DISCUSSION

We presented a causal inference analysis of observational EHR data to compare the effect of adding a second-line treatment for T2DM on HbA1c and BMI, in patients already treated with metformin. Our inferred HbA1c levels for up to 12 months of follow-up suggest that the effect of TZD, DPP-4 and GLP-1 inhibitors is comparable,
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under-represent children and young adults, as this popu-
Medicare and Medicaid. While the data potentially
negligible effect on BMI, DPP-4 and GLP-1 reduce BMI
whereas that of SU is smaller. While TZD and SU have a
negligible effect on BMI, DPP-4 and GLP-1 reduce BMI
after 12 months of treatment.

The analyzed data contain privately insured employees,
Medicare and Medicaid. While the data potentially
under-represent children and young adults, as this popula-
ration is rarely confronted with T2DM, there was no
major bias introduced, as can be seen by the comparison
of the age distribution in online supplementary figure 1
to the distribution published by the CDC for 2011.31

We addressed three challenges associated with causal
inference from EHRs, including fragmented data, identifi-
cation of the true set of confounders, and differences in
protocols or adoption rate across institutions. To address
fragmented data, we corrected for potential selection
bias using patients with incomplete data as censored. In
order to reduce the probability of incorrectly specifying
confounders or correcting for them, we took the following
three measures: For the first measure, we compared
confounder sets based on domain expertise with a compre-
hsive set of confounders based on available clinical and
demographic information of the patient to find minor
differences in predicted outcomes. We showed the DR
estimator improves balancing of confounders for both
confounder sets and especially for the domain expert
sets, but as noted by Austin29 for propensity score models,
in many settings it is likely that one can safely include
all measured baseline characteristics in the models. For
the second measure, we tested whether we could reduce
residual confounding by stratifying continuous values,
our study were treated with GLP-1 after 2005 and 38% of them treated during or after 2013, suggesting physicians may have considered this evidence when prescribing GLP-1. Also, patients on GLP-1 are typically younger than patients on other drug classes, in line with an observation made by others. Finally, patients with higher BMI tend to be prescribed GLP-1, and this is likely due to its known positive effect on weight. In our analysis, though, DPP-4 inhibitors are estimated to lead to BMI reduction comparable to GLP-1 agonists.

TZD is the only class predicted to maintain HbA1c at a stable level in 6 and 12 months, whereas HbA1c levels are predicted to increase over time in the other studied classes. A gradual weaning of the effect of SU on HbA1c levels had been previously described.

The estimates of HbA1c reported in the meta-analysis (MTC) we used for reference were higher than our EHR-based inference. We note that we predicted HbA1c in exact periods, while the MTC method combined heterogeneous time point measurements across the different clinical trials, some listed as having up to 5 years of follow-up. This may suggest that the meta-analysis captured later stages in the progression of T2DM, characterized by higher HbA1c levels.

As demonstrated by our analysis, as well as by others, EHR data can support causal inference and allow replication of clinical trial results. The advantages of this approach in terms of the labor and costs required to expand evidence-based medicine are clear. As the availability of EHR data increases and the many theoretical and technical challenges associated with detecting and correcting for confounders are addressed, we expect causal inference based on observational data to become more widely used.

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