Observational studies of treatment effectiveness: worthwhile or worthless?

Abstract: Observational studies which evaluate effectiveness are often viewed with skepticism owing to the fact that patients are not randomized to treatment, meaning that results are more prone to bias. Therefore, randomized controlled trials remain the gold standard for evaluating treatment effectiveness. However, it is not always possible to conduct randomized trials. This may be due to financial constraints, for example, in identifying funding for a randomized trial for medicines that have already gained market authorization. There can also be challenges with recruitment, for example, of people with rare conditions or in hard-to-reach population subgroups. This is why observational studies are still needed. In this manuscript, we discuss how researchers can mitigate the risk of bias in the most common type of observational study design for evaluation of treatment effectiveness, the cohort study. We outline some key issues that warrant careful consideration at the outset when the question is being developed and the cohort study is being designed. We focus our discussion on the importance of deciding when to start follow-up in a study, choosing a comparator, managing confounding and measuring outcomes. We also illustrate the application of these considerations in a more detailed case study based on an examination of comparative effectiveness of two antidiabetic treatments using data collected during routine clinical practice.

Keywords: epidemiology, therapeutics, diabetes mellitus, public health, effectiveness

Introduction

The randomized controlled trial (RCT) is considered the gold standard design for examining the effectiveness of a treatment.1 This is because randomization increases the likelihood that treatment allocation is undertaken independently of both known and unknown patient characteristics, although there remains a possibility of chance imbalances.2 This chance is inversely proportional to the sample size being studied.3 Such baseline imbalances associated with the outcome under study can confound the findings, leading to biased estimates of effectiveness.

Despite growing interest in the use of observational studies to evaluate effectiveness, their application remains contentious.4 This is because the absence of randomization means that treatment choice is usually influenced by the clinician’s perception of the effectiveness of the treatments being considered.5 Hence, baseline imbalances that can bias estimates of treatment effectiveness are almost always present. A comparison of outcomes across treated and untreated individuals in an observational study may lead one to erroneously conclude that treatment is not effective, when in fact the treatment may have been selectively given to those with the worst prognosis.5
Despite these challenges, observational studies of effectiveness do offer opportunities to examine questions that may not be possible using RCTs. First, they can be used to examine the effectiveness of medication that has already been granted marketing authorization and for which funding for further trials may be limited. Second, they can allow the examination of effectiveness for rare treatment indications. Third, a large observational study can be more representative of a clinical population and less prone to selection bias than a trial. Thus, it can allow investigation of the external validity of trial results in more diverse populations, such as ethnic minority groups and elderly patients, who are often underrepresented in conventional trials.

In this paper, we focus on use of cohort studies to evaluate effectiveness, where individuals are followed up from exposure to a treatment for the development of an outcome of interest. In particular, we will outline important considerations in cohort study design that can help to mitigate the risk of bias and conversely help to identify research questions of clinical effectiveness that are more suited to investigation using such a design. We then illustrate the application of these considerations in depth with reference to a case study based on our own completed work, which examined the comparative effectiveness of two antidiabetic treatments. Alternative study designs, such as self-controlled case series and case-control studies, are well described in the literature and will not be covered in this article.

Important considerations in the design of cohort studies to evaluate treatment effectiveness

In this section, we discuss four important aspects to consider when designing cohort studies to evaluate treatment effectiveness: when to start follow-up, choosing comparators, identification and measurement of confounding, and ascertainment of the outcome.

Start of follow-up

In cohort studies, individuals are ideally followed up from when they are first initiated on a treatment (new-user design). However, this is not always possible, and some studies include individuals who have already been receiving the treatment before the start of follow-up (prevalent user design). There are advantages and disadvantages to both approaches. In the new-user design, by excluding the prevalent users (left truncation) a “prevalent user” bias is eliminated which can be linked to the fact that they have already “survived” a prior period of treatment use without any negative consequences. New-user design allows for the adjustment of confounders at baseline, when the decision was made to initiate treatment, thus helping to eliminate bias. These biases are most relevant when the risk of an outcome of interest is known to be highest in the early stages of treatment. An example of prevalent user bias was seen in studies which demonstrated that hormone replacement therapy (HRT) prevented coronary heart disease, whereas subsequent trials found HRT to be harmful. The observational studies included prevalent users who were taking HRT before study follow-up and had already survived a period of use without any harm. They had a lower likelihood of cardiovascular outcomes at initiation of study follow-up, which led to the bias seen in the risk estimates that suggested a protective effect of HRT.

Restricting inclusion to “new users” only does have limitations. It reduces the sample size of cohort studies and may limit long-term follow-up (right truncation). Provided the risk of biases and potential directionality are carefully considered, the evaluation of prevalent users can still be useful. For example, even though estimates for cardiovascular outcomes were biased in the HRT observational study, effectiveness estimates produced for other outcomes where risks were cumulative over time, such as colon and breast cancer, were unbiased and similar to the trials. In practice, it can sometimes be helpful to split the cohort into “new users” and “prevalent users” and analyze treatment effectiveness separately in each group, so that the limitations of left and right truncation can be acknowledged (Table 1).

Choice of comparators

Although it is common practice in RCTs, estimating the effectiveness of treatments by comparing treated and untreated individuals in a cohort study can lead to bias as treatment may be indicated only for those with a specific prognosis. The results may suggest that treatment is ineffective if an untreated group has a better prognosis or, conversely, may exaggerate effectiveness if the untreated group has a worse prognosis. This type of bias, which often occurs in observational studies, is known as channeling bias or confounding by indication, and arises when the indication for choosing a particular treatment also affects the outcome. Treated and untreated groups commonly differ in terms of disease severity, which can be difficult to measure in a cohort study. For example, using a cohort study design, Freemantle et al investigated the effectiveness of using spironolactone in reducing mortality in patients with severe heart failure treated in clinical practice.
Table 1 Important considerations in the design of cohort studies to evaluate treatment effectiveness and how to mitigate the risk of bias

<table>
<thead>
<tr>
<th>Consideration</th>
<th>Bias</th>
<th>Definition</th>
<th>Problem</th>
<th>Approach that can help</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starting follow-up</td>
<td>Prevalent user bias</td>
<td>Bias that arises from the inclusion of individuals who have already initiated treatment prior to inclusion in study</td>
<td>Prevalent users survived a period of exposure before the study and may be at lower risk of an event</td>
<td>New-user design</td>
</tr>
<tr>
<td>Starting follow-up</td>
<td>Incomplete follow-up bias</td>
<td>Bias that arises owing to too short a period of follow-up for an outcome</td>
<td>If an event is due to a cumulative effect of a treatment, insufficient follow-up may not allow enough time for the event to occur</td>
<td>Prevalent user design</td>
</tr>
<tr>
<td>Comparator choice</td>
<td>Channeling bias</td>
<td>Bias that arises when the clinical indication for choosing a particular treatment also affects the outcome</td>
<td>If there is channeling bias then an effectiveness estimate may be biased, with a directionality based on whether the comparison group has better or worse prognosis</td>
<td>Use of an active comparator where there is equipoise in treatment choice</td>
</tr>
<tr>
<td>Identification and measurement of confounders</td>
<td>Confounding bias</td>
<td>Bias that arises when factors that affect both treatment choice and outcome are not accounted for</td>
<td>The directionality of this bias can be unpredictable and will depend on both accurate identification and measurement of confounders, as well as the extent of unmeasured confounding</td>
<td>Identification, accurate measurement and controlling for confounding</td>
</tr>
<tr>
<td>Outcome ascertainment</td>
<td>Reporting bias</td>
<td>Bias that arises owing to an imbalance in recording of an outcome across treatment groups</td>
<td>This can lead to a treatment appearing more or less effective when, in fact, the outcomes are simply recorded differently across treatment groups</td>
<td>Careful consideration and understanding of how an outcome might be detected and recorded in clinical practice</td>
</tr>
<tr>
<td>Outcome ascertainment</td>
<td>Attrition bias</td>
<td>Bias that arises owing to an imbalance in duration of follow-up across treatment groups</td>
<td>This can lead to lower numbers for an outcome in the group with shorter follow-up time</td>
<td>Ensuring comparison groups have similar follow-up time across which an outcome can realistically be recorded</td>
</tr>
</tbody>
</table>

randomized aldactone evaluation study (RALES) clinical trial, which found that spironolactone reduced mortality; their observational study found a lower mortality in the untreated group than in those treated with spironolactone. These contrasting results were explained by the difference in disease severity across the two groups, as spironolactone was primarily prescribed to those with more severe heart failure and the worst prognosis.

While it may often be difficult to estimate the effect of treatment against no treatment based on data from clinical practice owing to channeling bias, Smeeth et al successfully replicated the findings from large trials in their cohort study. They showed that statin use was effective in reducing vascular outcomes compared to non-use. In this instance, they were able to match statin users to non-users with similar disease severity and hence mitigate the risk of bias.

Another approach in cohort study design that can often help to yield more accurate estimates involves the inclusion of an active comparator group, if the clinical question allows. For example, consider a cohort study comparing two alternative first-line antihypertensive agents, the angiotensin-converting enzyme inhibitors ramipril and perindopril, for reducing blood pressure. The choice of either ramipril or perindopril is unlikely to be driven by many prognostic factors other than prescriber preferences or local formulary policy, and hence patients are likely to have similar disease severity at baseline. However, such a study is limited to providing estimates of the relative effectiveness of the two treatments only, and not of the effectiveness of treatment compared to no treatment.

Identification and measurement of confounding
In clinical practice, scenarios where there is complete baseline balance in disease severity are rare, and the design and analysis of most observational studies of effectiveness will need to actively remove sources of potential confounding bias. This involves the identification of all factors that cause the outcome and are associated with treatment choice, but are not on the treatment–outcome pathway. This can be achieved with visual maps called direct acyclic graphs.
Once confounders have been identified, several analytical approaches can be applied to remove the influence of confounders on effect estimates, such as propensity-score based methods and standard multivariable regression methods.\textsuperscript{17,20} Patorno et al used propensity score matching in a large cohort study examining the antidiabetic agent canagliflozin. They demonstrated that canagliflozin effectively reduced admission to hospital due to heart failure compared to several other antidiabetics, with estimates consistent with previously completed clinical trials.\textsuperscript{21} Propensity score matching facilitated the removal of baseline imbalances in disease severity across treatment groups, which allowed reliable estimates to be obtained. The removal of baseline imbalance was not possible in the study by Freemantle et al, however, as imbalances in disease severity could not all be captured through simple matching.\textsuperscript{5} Although the analytical approach is important, evaluation of the completeness and validity of the recording of the confounding variables and risk of unmeasured confounding is equally crucial.\textsuperscript{17,22} If the source of data does not include information on confounding variables, as in the study by Freemantle et al,\textsuperscript{5} it creates a problem of unmeasured confounding and will bias analyses.\textsuperscript{17} Methodological approaches involving the use of proxy variables for confounders and sensitivity analysis can be considered to explore the impact of unmeasured confounding on the analysis.\textsuperscript{23} However, despite these approaches, the limitations of such a study must be reconsidered, especially if unmeasured confounding is suspected to be highly influential.

\textbf{Outcome ascertainment}

In any cohort study comparing the effectiveness of different treatments, all groups at baseline must have an equal chance of recording the outcome being investigated. A thorough consideration of whether individuals receiving one treatment may have longer follow-up, or are more likely to be screened for an event, to be intensively managed or to have better data recorded, must be made at the outset.\textsuperscript{7} For example, individuals prescribed the anticoagulant warfarin, which requires regular international normalized ratio blood testing, as opposed to direct oral anticoagulants, which do not, may have more frequent health care contacts and thus greater opportunity to report symptoms that lead to identification of an outcome being considered, eg, minor stroke. This could falsely lead to higher reporting and recording of an outcome in the warfarin group, resulting in estimates suggesting that warfarin is inferior in effectiveness when this is purely due to a reporting bias or an attrition bias (imbalance in the duration of follow-up).

**Case study example**

**Overview**

In this case study, we discuss how we applied the considerations detailed in the previous section to design a cohort study to compare the effectiveness of two antidiabetic treatments, sitagliptin vs sulfonylureas, as add-on to metformin for the treatment of type 2 diabetes mellitus. Both treatments are widely used add-on options for managing type 2 diabetes mellitus when metformin alone has proved inadequate. Guidelines from the UK National Institute of Health and Care Excellence, as well as other international guidelines, do not discriminate between these add-on treatments in terms of effectiveness.\textsuperscript{24} Our study investigated their glycemic effectiveness when used as part of routine clinical care in UK general practice.

We undertook this study in The Health Improvement Network Primary Care Database (version 15), which contains anonymized data from around 670 general practices across the UK. Scientific approval to undertake this study was obtained from the IQVIA World Publications Scientific Review Committee in August 2016 (reference number 16-072).\textsuperscript{25} This retrospective cohort study examined changes in HbA\textsubscript{1c} from baseline after 12 months of treatment between those prescribed sulfonylurea vs sitagliptin as add-on to metformin for type 2 diabetes mellitus. The driver behind this study was to investigate the external validity of several trials which had concluded that both treatments produced a similar glycemic reduction after initiation. Details of how we identified individuals with type 2 diabetes mellitus have been previously described in depth.\textsuperscript{26}

The baseline characteristics of our cohort, and how this cohort study population differed from the corresponding trial populations, are shown in Table 2.

In summary, our cohort study population was older, had worse baseline HbA\textsubscript{1c} control and had higher weight than the populations in the completed trials.

After adjustment for baseline HbA\textsubscript{1c}, sex, age and other identified potential confounders in our analysis, we found that 12 months after treatment initiation the HbA\textsubscript{1c} level was on average 1 mmol/mol (mean difference 0.89 mmol/mol, 95% CI 0.33–1.45) higher for those prescribed sitagliptin compared to sulfonylureas (Table 3). Despite its statistical significance, a difference of up to 1.45 mmol/mol is not considered clinically significant, given than such a small quantitative difference in HbA\textsubscript{1c} would not impact on the short- or long-term prognosis of diabetes.\textsuperscript{28} In fact, clinically relevant differences in HbA\textsubscript{1c} are those that typically exceed 5.5 mmol/mol and ideally 10.9 mmol/mol in magnitude.\textsuperscript{27} Our
cohort study estimate was found to compare favorably with that from the meta-analysis of completed RCT previously undertaken (Figure 1), which also highlighted no significant difference (weighted mean difference 0.54 mmol/mol, 95% CI –0.28 to 1.35).28

Case study in context
We approached the study question having considered each of the design issues detailed in the earlier section (see section “Important considerations in the design of cohort studies to evaluate treatment effectiveness”). Our choice of a new-user design was taken to mitigate the risk of prevalent user bias that would arise by including individuals who had already been exposed to the treatment and hence experienced a glycemic benefit.10 This was achieved by following individuals from their first prescription of sitagliptin or sulfonylurea and ensuring that they had not been prescribed any antidiabetic agents other than metformin in the preceding 12 months.

We chose to use an active comparator in this study to mitigate the risk of channeling bias that might have arisen if we had compared the treatment group to a non-treated group, as there would have been a substantial difference in disease severity.5 Both sitagliptin and sulfonylurea are commonly used as add-on to metformin in the type 2 diabetes clinical pathway, which would help to balance disease severity at baseline. Nevertheless, there were differences at baseline between the groups in HbA1c and weight, which we believed would influence both treatment choice and outcome and hence could lead to confounding bias. We explored the recording and measurement of these confounders across treatment groups to ensure adequacy before controlling for them using a multivariable regression model.17 We undertook several sensitivity analyses exploring subgroups such as those who persisted with treatment for the study duration to ensure that the findings were robust. Finally, to eliminate the risk of recording bias, we analyzed both treatment arms to ensure that the frequency of HbA1c recording across both groups over time was similar.7

To investigate the robustness of our estimate, we compared our study to the existing literature and highlighted how the comparative effectiveness estimate of this cohort study compared favourably to a meta-analysis of completed RCTs.28 The absolute change in HbA1c observed with both treatments, however, was greater in our study than that observed in the trials. This may have been due to the fact that our baseline population had worse disease (worse glycemic control) at treatment initiation than those in the previously completed trials, and hence had differential scope for improvement.

### Table 2

Comparison of baseline characteristics from three randomized controlled trials and the present case study

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants, n</th>
<th>Age (years), mean (SD)</th>
<th>Male, n (%)</th>
<th>HbA1c % (SD) [mmol/mol, SD]</th>
<th>Weight (kg), mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sita Sulf</td>
<td>Ahrén et al (2014)30</td>
<td>302</td>
<td>54.3 (9.8) Sita 516</td>
<td>307</td>
<td>54.3 (9.8) Sulf 534</td>
</tr>
<tr>
<td>Sita Sulf</td>
<td>Arechavaleta et al (2011)31</td>
<td>516</td>
<td>56.3 (9.7) Sita 534</td>
<td>519</td>
<td>56.3 (9.7) Sulf 556</td>
</tr>
<tr>
<td>Sita Sulf</td>
<td>Seck et al (2010)32</td>
<td>588</td>
<td>56.8 (9.3) Sita 584</td>
<td>556</td>
<td>56.8 (9.3) Sulf 556</td>
</tr>
<tr>
<td>Sita Sulf</td>
<td>Sharma et al (present study)</td>
<td>3,306</td>
<td>58.9 (11.2) Sita 3,306</td>
<td>15,880</td>
<td>58.9 (11.2) Sulf 3,306</td>
</tr>
</tbody>
</table>

Abbreviations: Sita, sitagliptin; Sulf, sulfonylurea.
Comparison to previously completed trials is a common approach used to demonstrate the validity of results in observational studies. As in our example, the studies by Smeeth et al and Patorno et al detailed earlier were able to compare their findings to estimates from previously completed trials for consistency. A comparison trial, however, is not always available, and therefore careful consideration of the issues outlined in this article can help to ensure a more robust approach to designing observational studies of effectiveness and mitigating the risk of bias. Equally, traditional challenges common to all observational study designs, such as handling missing data and the risk of exposure misclassification, remain when undertaking cohort studies and must also be carefully considered and managed.

**Conclusion**

In this manuscript, we describe some key considerations for clinical researchers that can help to mitigate the risk of bias when designing cohort studies evaluating effectiveness. These considerations can also help researchers to identify clinical questions that are more suited to such a cohort study design. Their overall purpose is to ensure that the characteristics of groups across which the treatments are being compared are as similar as possible at baseline in terms of disease severity and, in addition, that the occurrence of the outcome of interest is reported equally in all groups. They must also be assessed in the context of traditional methodological challenges of observational studies, such as the possible existence of missing data and the risk of exposure misclassification. However,
despite these obstacles, these considerations can help clinical researchers and epidemiologists to identify focused clinical questions where observational studies of effectiveness may be most worthwhile and, potentially, even advantageous.

Transparency declaration
I, Manuj Sharma, lead author, confirm that this manuscript is an honest, accurate and transparent account of the studies being reported; that no important aspects of the studies have been omitted; and that any discrepancies from this study as planned from our protocol have been explained.

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Author contributions
MS, IN and IP collectively planned the study. MS performed the analyses and wrote the manuscript. All authors contributed to data analysis, drafting and revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

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References

