Can a Mendelian Randomization Study Predict the Results of a Clinical Trial? Yes and No

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Abstract: Randomized controlled trials are considered at the top of the evidence hierarchy. However, in several cases randomized trials cannot be conducted or have not yet been completed. In such settings observational studies may provide important inference, yet traditional statistical adjustment methods fall short of controlling for all potential confounders, as unknown confounders cannot be taken care of by even the most sophisticated statistical tools. The mendelian randomization study is a type of research design which simultaneously exploits random transmission of genes and genetic linkage to obtain inferential estimates from the association between specific genetic variants known to modulate given risk factors and the corresponding outcomes of interests. Despite several developments in this field, there remain several areas of further research, and discrepancies between mendelian randomization studies and the corresponding randomized trials have already been recognized. Nonetheless, it is likely that this novel type of study will be used more commonly in the future, and a working knowledge of its pros, cons, and range of validity is crucial for conscientious interpretation and application. We thus aimed to concisely yet poignantly introduce the scholarly reader to this novel type of research design, notwithstanding that complementarity prevails in most cases over overlap between mendelian randomization studies and randomized trials.

Keywords: Adjustment, Confounding, Inference, Mendelian randomization study, Observational study, Prediction, Randomized controlled trial.

INTRODUCTION

Clinical decision-making should best be based on formal scientific experiments. It is widely accepted that a large randomized trial represents the best single piece of clinical evidence, whereas a homogeneous pairwise meta-analysis is considered the best comprehensive source of scholarly information [1-2]. Yet, novel developments in clinical research methods, which are posed to challenge the role of randomized controlled trials and pairwise meta-analyses, are being steadfastly developed, and include mendelian randomization studies, network meta-analyses, and umbrella reviews [3-5]. Whilst network meta-analyses and umbrella reviews typically incorporate one or more randomized trials, thus borrowing their strengths [6], the mendelian randomization approach has been proposed in as early as 1991 to minimize confounding in observational studies [7].

The preamble is of course that it remains impossible to conduct in a timely and ethical fashion a randomized trial for any given topic of interest [8-9]. Accordingly, observational studies (either explicitly designed or retrospectively analyzed) remain an important source of evidence, albeit of lesser strength. Indeed, observational studies are always fraught with a risk of residual confounding by unknown or inappropriately adjusted effect modifiers [10]. From stratification to logistic regression and propensity score matching, traditional adjustment methods have become more and more refined, and in many cases carefully conducted and analyzed observational studies appear to agree with the corresponding megatrials [11]. Yet, several examples of observational inference later disproved by a randomized trial are available.

THE MENDELIAN RANDOMIZATION STUDY

In order to overcome the limitations of traditional adjustment methods, and exploit the unique features of genetics, Gray and Wheatley proposed the mendelian randomization study (Figure 1) [7,12]. This type of research aims to adjust for known and unknown confounders when aiming at observational inference, exploiting Mendel second law of inheritance, which specifies that separate genes are inherited independently from each other, as long as they reside...
Figure 1: Interactions between genetics, confounders, interventions, risk factors, and outcomes. Schematic representation of the logical flow from specific alleles to risk factor modulation and outcomes in a mendelian randomization study (thick arrows). An observational study relies on traditional statistical adjustment, which takes into account the impact on risk factors of known confounders and moderators but cannot adjust for the impact on outcomes of unknown confounders (thin and dashed lines, respectively). A randomized controlled trial focuses on a specific intervention of interest and formally tests its impact on a given risk factor or, more importantly, on an outcome of interest (thin and dashed lines, respectively), and asymptotically balances the distribution of known and unknown confounders.

on separate chromosomes. It basically relies on one or more alleles known to be associated with a risk factor of interest. Once such genetic association is proved beyond random variability, then the association between such genetic variants and the outcomes of interest is explored. As genetic variants are assumed to be randomly assigned at conception, the association between such genetic variants and a given condition can be considered independent of confounders. This is quite different from observational studies and randomized trials. Specifically, an observational study relies on traditional statistical adjustment, which takes into account the impact on risk factors of known confounders and moderators but cannot adjust for the impact on outcomes of unknown confounders. A randomized trial focuses on a specific intervention of interest and formally tests its impact on a given risk factor or, more importantly, on an outcome of interest, and asymptotically balances the distribution of known and unknown confounders.

Several authors have welcomed this innovative tool which builds on the expanding knowledge base on genetics as a means to test novel hypotheses or retest apparently established ones, given the assumed superiority to traditional statistical adjustment tools [7,12-13]. There remain some theoretical caveats, though. First, a prerequisite of any mendelian randomization study is a reliable association between genotype and exposure and a clear understanding of the pathophysiology of the genetic variants of interest. This can be taken for granted in many cases, but more robust proof may remain elusive in some cases. Second, confounding may still occur due to linkage disequilibrium. Third, genetic variants with multiple effects may confound the results. Fourth, canalization, which can be defined as the modulation of the effects of genetic variations during development, may also significantly impact on the association between genotype and outcomes. Accordingly, the real virtue of mendelian randomization studies for causal inference rather than association appraisal remains debated, and it is not unexpected that mendelian randomization studies still represent only a paucity of observational studies. Indeed, a dedicated PubMed search (updated on July 28, 2015) for ‘(mendelian AND random*)’, and restricted to works published within the prior 5 years, highlighted 541 citations, which pale in comparison to the 89,911 for ‘(logistic AND regression)’, the 11,639 for ‘cox AND hazard AND analysis’, and the 6,912 for ‘propensity AND (score* OR match*)’, representing, respectively, citations of studies reporting on logistic regression, Cox proportional hazard analysis, and propensity score adjustment/matching.

CASE STUDIES

Under the premise of aiming at a simplified take at mendelian randomization studies, we may elaborate that two typical examples of such studies can be found in the scholarly literature. The first is a study exploiting
specific genetic variants and which does not relate to any given clinical intervention yet. Accordingly, it cannot be directly compared to prior, current or future randomized controlled trials. For instance, the mendelian randomization study with the largest number of scholarly citations in Google Scholar is an observational study using both traditional statistical adjustment techniques and mendelian randomization to appraise the association between C-reactive protein and blood pressure/hypertension [14]. As no pharmacologic intervention specifically and selectively targeting C-reactive protein levels or activity is yet available, the findings from this study cannot be directly confirmed or disproved, nor formally applied.

The second type of mendelian randomization analysis is instead a study which focuses on an association which can also be translated into a randomized controlled trial. Perusal of a set of key applications of mendelian randomization studies in this specific fashion may inform the scholarly reader on the pros and cons of this novel analytical approach (Table 1). For instance, the impact of alcohol consumption on blood pressure has been analyzed within the context of mendelian randomization by Chen et al., with results similar to those provided in a comprehensive meta-analysis of randomized controlled trials on alcohol cessation [15-16]. Specifically, a common polymorphism in aldehyde dehydrogenase 2, which is an established surrogate for measuring alcohol consumption, was associated with increased blood pressure and increased risk of hypertension, similarly to the established impact of alcohol intake on blood pressure and hypertension risk.

The above and other favorable cases of agreement between mendelian randomization studies and randomized controlled trials are reassuring, yet it is not easy to identify cases in which there was a substantial agreement or discrepancy between a mendelian randomization study and the corresponding randomized controlled trial. Specifically, an informal scholarly search enabled us to recognize substantial

<table>
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<tr>
<th>Association</th>
<th>Mendelian randomization study (MRS)</th>
<th>Randomized controlled trial (RCT)</th>
<th>Comparative analysis</th>
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<tr>
<td>Alcohol – blood pressure</td>
<td>Chen (2008): MRS with meta-analysis of 8 studies of a common polymorphism in aldehyde dehydrogenase 2 (ALDH2) as a surrogate for measuring alcohol consumption and blood pressure/hypertension, suggesting that alcohol intake is significantly associated with blood pressure and hypertension</td>
<td>Xin (2001): meta-analysis of 15 RCTs comparing the effect alcohol cessation versus control on blood pressure, suggesting that alcohol cessation is associated with significant reductions in blood pressure</td>
<td>Agreement between MRS and RCT, with both supporting prior evidence of an association from traditional observational studies</td>
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<td>B-type natriuretic peptide (BNP) – diabetes</td>
<td>Pfister (2011): MRS with meta-analysis of 11 case-control studies of the variant rs198389 within the BNP locus and the incidence of type 2 diabetes, suggesting a significant association between BNP and diabetes</td>
<td>Yancy (2004): RCT in 210 patients with heart failure receiving nesiritide (recombinant form of the 32 amino acid human BNP) versus placebo, suggesting no significant impact of nesiritide on glucose tolerance</td>
<td>Discrepancy between MRS and RCT, with the latter not confirming the MRS results</td>
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<td>Vitamin D – blood lipid profile</td>
<td>Skaaby (2013): MRS combining 3 studies and including 12,911 subjects of 3 filaggrin gene mutations (R501X, 2282del4, and R2447X) and blood lipid profile, suggesting a significant association between the mutations and an adverse lipid profile</td>
<td>Pilz (2015): RCT in 200 subjects of vitamin D3 versus placebo, suggesting no significant impact of vitamin D3 on blood lipid profile</td>
<td>Discrepancy between MRS and RCT, with latter not confirming the MRS results</td>
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<td>Vitamin D – mortality</td>
<td>Trummer (2013): MRS including 3316 subjects of 3 common single-nucleotide polymorphisms (SNPs) associated with 25-OH-vitamin D concentrations and mortality, suggesting no significant association between them</td>
<td>Chowdhury (2014): meta-analysis of 22 RCTs on vitamin D2 and D3 supplementation, suggesting that only vitamin D3 is associated with a significantly reduced risk of mortality</td>
<td>Discrepancy between MRS and RCT, with latter suggesting a mortality benefit apparently disproved by the MRS results</td>
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<td>Testosterone – Cardiovascular risk</td>
<td>Haring (2013): MRS in 1882 men of polymorphisms at the SHBG gene (rs12150660) and X chromosome (rs5934505) and cardiovascular risk</td>
<td>Corona (2014): meta-analysis of 75 RCTs on testosterone-boosting medications, suggesting no significant impact of these drugs on cardiovascular risk</td>
<td>Agreement between MRS and RCT, with both not supporting prior evidence of an association from traditional observational studies</td>
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discrepancies only in pairs of studies appraising the impact on B-type natriuretic peptide on diabetes, and of vitamin D on blood lipid profile and mortality [17-22].

Another recent and interesting case study in mendelian randomization analysis is the one focusing on interleukin-1 receptor antagonist (IL-1Ra), and its association with the risk of adverse blood lipid profile, coronary heart disease, stroke, and aortic aneurysm [23]. Notwithstanding the general limitations of mendelian randomization [24], in this specific study the use of the term ‘cardiovascular risk’ was potentially misleading, as risk implies the likelihood of an event occurring over time, and therefore one would need to explore the incidence of an event, rather than the prevalence. In addition, the value of this type analysis inevitably depends on the quality of the original studies, and therefore it cannot be free of enrollment or survivor biases deriving from the original studies. In addition, its findings are not clearly consistent with prior genetic association data exploring the impact of IL-1, IL-1Ra and the effects of proven cardioprotective agents such as HMG-CoA inhibitors, such as those reported Waehre et al. [25].

More specifically, let us consider a hypothetical scenario by which gene X modulates the ‘risk’ of dying from disease A, by which carriers of X are more likely to have a more severe form of A, by which when they become ill, they develop more complications, and die sooner. If we were to perform a cross-sectional study of all subjects with the disease A, we could paradoxically find that less subjects carrying X with the disease A will be represented because X-carriers have a shorter duration of the disease A before severe complications or death occur. In this specific case, the authors found that carriers of hyper-producing IL-1Ra gene alleles have mildly higher levels of IL-1Ra and mildly lower levels of C-reactive protein, yet mildly higher levels of low-density lipoprotein cholesterol. The authors suggested that IL-1Ra is increasing low-density lipoprotein cholesterol (LDL), which is known to associate with coronary artery disease, and hence have IL-1Ra enhancing the likelihood of coronary artery disease through increased LDL (Figure 2). Yet, what if it were not the case, and, on the other hand, higher IL-1Ra levels were reducing mortality/increasing longevity in patients with coronary artery disease [26], a way by which IL-1Ra would be modulating the negative effects of LDL? This example, while interesting and potentially informative, is however based on several untested and non-evidence-based assumptions.

**Figure 2: Case mendelian randomization study.** In a recent mendelian randomization study exploring the association between interleukin-1 (IL-1) receptor antagonist (IL-1Ra) genotypes with higher IL-1Ra plasma levels and higher prevalence of coronary artery disease, the authors suggested that IL-1Ra causes coronary artery disease, possibly through increased levels of low-density lipoprotein (LDL) cholesterol. While this is biologically possible it is not evidently plausible as there are no biologic data linking IL-1Ra to increased LDL levels nor to increased prevalence of coronary artery disease. An alternative hypothesis would be that higher IL-1Ra levels may affect the prevalence of coronary artery disease by increasing survival so that the actual prevalence of this condition is increased. This hypothesis is possible and also plausible, as there are preclinical and clinical data linking enhanced IL-1 activity and worse outcomes in coronary artery disease.

Accordingly, only time and the cumulative evidence provided by recent and ongoing randomized controlled trials on this topic will provide a definite answer to this research question [27-30], and similar considerations may apply to other mendelian randomization studies.

**CONCLUSIONS**

Despite their inherent limitations, the success and impact of mendelian randomization studies will likely continue to increase over time. Concomitantly, comparative research on their accuracy and precision in comparison to other types of observational research and pertinent randomized controlled trials will accrue. These efforts will further bolster the comprehensiveness of the many different layers of clinical research tools (Figure 3), under the key premise that complementarity will most often prevail over overlap between mendelian randomization studies, randomized trials and other types of research tools [31].
CONFLICTS OF INTEREST

Drs. Abbate and Dinarello hold a patent entitled ‘Compositions and methods for modulating cardiac conditions’ (US 20130195859 A1).

REFERENCES


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