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THEORETICAL NOTE

Causal Inference Methods for Intergenerational Research Using Observational Data

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Identifying early causal factors leading to the development of poor mental health and behavioral outcomes is essential to design efficient preventive interventions. The substantial associations observed between parental risk factors (e.g., maternal stress in pregnancy, parental education, parental psychopathology, parent–child relationship) and child outcomes point toward the importance of parents in shaping child outcomes. However, such associations may also reflect confounding, including genetic transmission—that is, the child inherits genetic risk common to the parental risk factor and the child outcome. This can generate associations in the absence of a causal effect. As randomized trials and experiments are often not feasible or ethical, observational studies can help to infer causality under specific assumptions. This review aims to provide a comprehensive summary of current causal inference methods using observational data in intergenerational settings. We present the rich causal inference toolbox currently available to researchers, including genetically informed and analytical methods, and discuss their application to child mental health and related outcomes. We outline promising research areas and discuss how existing approaches can be combined or extended to probe the causal nature of intergenerational effects.

Keywords: causal inference, genetically informed methods, intergenerational effects, quasi-experimental studies, within-family designs

Understanding the role of parents in shaping their children’s outcomes has been a major focus of influential theories across psychology, psychiatry, and behavioral genetics. Delineating the nature of intergenerational effects is however challenging. Randomized controlled trials (RCTs) and experiments are often seen as the gold standard of causal inference but have limited application to intergenerational research due to feasibility and ethical concerns, for example, it would be unethical to design randomized cross-fostering experiments to examine whether environmental or genetic factors explain the intergenerational transmission of risk for schizophrenia. However, observational studies are prone to a variety of biases, including reverse causation, selection biases, or confounding (see Delgado-Rodríguez & Llorca, 2004, for definitions). For example, the outdated theory of the “schizophrenogenic mother” claimed that schizophrenia was caused by an “overprotective but subtly rejecting mother” (Seeman, 2016), which led to social stigma, with maternal behavior being blamed as the cause of their offspring’s psychopathology (Harrington, 2012). Research on the “schizophrenogenic” mother was however, distorted by selection and information biases, reverse causation (e.g., using case-only studies and neglecting offspring effects on maternal behavior) and did not account for genetic factors (Parker, 1982). Today, schizophrenia is known to be a

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highly heritable disorder with hundreds of common and rare genetic variants identified and studies point toward neurobiological underpinnings linked to synaptic processes and brain maturation starting in fetal life (Birnbaum & Weinberger, 2017; Singh et al., 2022; Trubetskoy et al., 2022). Taken together, those findings suggest that exposure to parental behavior is unlikely to cause development of schizophrenia in the offspring.

Despite the presence of confounding and other biases in observational research like that on “schizoprenogenic mothers,” these biases can be addressed by applying causal inference methods. Resolving the questions of causal effects in turn has fundamental implications not only for preventing adverse and promoting favorable child outcomes, but also for the way we conceive the role of parents in shaping their offspring’s development. For example, mistakenly attributing a specific (parental) exposure as the cause of child mental health difficulties would likely result in ineffective interventions due to targeting noncausal factors, thus wasting resources and delaying the development and implementation of more suitable interventions. Furthermore, stigmatizing and blaming either parents or children for being responsible for existing mental health problems could increase mental health problems and negatively impact the parent–child relation.

This review provides a summary of causal inference methods for intergenerational settings that are needed to disentangle putative causal effects from confounded associations, including recent designs using measured genetic variants. For our review, we define exposures as specific parental traits or behaviors with potential effects on a child phenotype. These exposures can either occur prenatally (e.g., maternal stress or drug use during pregnancy), perinatally (e.g., birth complications), or postnatally (e.g., parental education, parent–child relationship quality, parental psychopathology). We draw on empirical examples from the fields of child development, child and adolescent psychology and psychiatry, and allied disciplines, highlighting the multidisciplinary nature of intergenerational research and the broad applicability of such methods beyond those disciplines. We do not offer an extensive background to theories of causal inference and mathematical notation of specific estimands but provide key references. A brief conceptualization of causation is followed by a presentation of existing methods applied to intergenerational settings, where we outline the advantages and limitations of each method and provide references for further readings. Finally, we make suggestions for combining and extending existing approaches to strengthen causal inference.

Causal Inference

Research on the etiology of mental health and behaviors often aims to identify causal effects. An exposure can be considered as a cause for an outcome if changing this exposure results in a change in the outcome of interest, while holding everything else constant. Confounding is a major challenge for causal inference. In the simpler setting of a point-exposure $X$ (i.e., time-fixed exposure), this occurs when a third variable ($Z$) has a causal effect on both the exposure ($X$) and the outcome ($Y$), leading to a spurious association between $X$ and $Y$ (Figure 1a). For example, the effect of smoking during pregnancy on offspring behavioral problems may be entirely confounded by genetic factors predisposing to both smoking behavior and psychopathology (Figure 1b).

**Figure 1**

**Confounding**

(a) $X \rightarrow Y$  
(b) $G \rightarrow SDP \rightarrow ADHD$

Note. (a) Visualisation of confounding, where $X$ has no causal effect on $Y$. A confounder $Z$ has a causal effect on both $X$ and $Y$. Not adjusting for $Z$ would result in a spurious association between $X$ and $Y$. Note that there could also be a real effect ($X \rightarrow Y$), which is only partly confounded by $Z$. (b) Example of the association between smoking during pregnancy (SDP) and attention deficit hyperactivity disorder (ADHD) which is confounded by shared genetic effects ($G$), influencing both maternal smoking and child ADHD. For simplification, we assume continuous variables.

**Exchangeability**

A key assumption of causal inference methods is exchangeability, that is, the same outcome distribution would be observed if exposed and unexposed individuals were exchanged (Greenland & Robins, 1986). Consider the example of smoking during pregnancy as a dichotomous exposure and offspring attention deficit hyperactivity disorder (ADHD) as a dichotomous outcome, where exposure to smoking in utero is assumed to lead to ADHD in childhood. In the counterfactual framework, an individual can be considered as exposed and unexposed at the same time, resulting in two counterfactual outcomes (also termed potential outcomes), for example, developing ADHD or not. The counterfactual outcomes for an individual can be compared to estimate the causal effect of the exposure, for example, a child developing ADHD when exposed to maternal smoking versus not developing ADHD when not exposed (Little & Rubin, 2000).

However, the assumption of exchangeability is not testable in practice, where each individual cannot be exposed and unexposed at the same time. Hence, causal effects cannot be estimated at the individual level due to missing data (Hernán & Robins, 2020). Causal inference methods therefore aim to approximate the counterfactual scenario to meet the exchangeability assumption at the group level. In our example, exchangeability is fulfilled when the distribution of ADHD diagnoses of individuals exposed to smoking during pregnancy would equal the distribution of ADHD in the unexposed individuals if the latter had been exposed to smoking during pregnancy, and vice versa. Concretely, this means that confounders are balanced in the exposed and unexposed groups, so that the treatment is the only meaningful difference. Note that although we present an example with a dichotomous exposure and outcome, the potential outcomes can also be compared for non-dichotomous exposures and outcomes (Hernán, 2004).

Exchangeability is achieved in RCTs or randomized experiments by random assignment to the exposed and unexposed groups, resulting in balanced confounders at the group level (Cornfield, 1976). Note that exchangeability may not be present even in RCTs, for example if the randomization was not successful (e.g., small sample size) or because of differential attrition between the groups. Exchangeability
is assumed in other causal inference methods which approximate the counterfactual framework to derive an average causal effect for a given population (Hernán & Robins, 2020). Essentially, designs without random assignment assume exchangeability or conditional exchangeability, that is, individuals are exchangeable after conditioning on a sufficient set of confounders (invoking the strong assumption of no unmeasured confounding). For example, offspring of mothers who smoke during pregnancy and of those who do not may be exchangeable after conditioning on the genetic risk for smoking and ADHD, assuming no other common factors of X and Y (Figure 1b).

Natural Experiments

Naturally occurring events like policy changes or natural disasters can be employed by researchers as quasi-experiments that randomly allocate individuals to given circumstances (Rutter, 2007), which are accompanied by differential exposure to the changing factors (e.g., stress or education). For example, exposed and unexposed individuals can be compared before and after the events. Therefore, comparable to RCTs, natural experiments do not require researchers to identify and control for multiple confounders, although further adjustment for confounders by matching or regression has been recommended (Dunning, 2008).

Natural experiments have also been used to study intergenerational effects. For example, a study examining pregnant mothers and their future offspring following a strong earthquake in Chile reported a negative effect of the prenatal exposure on cognitive outcomes of offspring from disadvantaged families (Torche, 2018). The authors used a difference-in-differences design, which allowed them to compare the difference in outcomes of exposed and unexposed mother-child dyads over time (change/difference over time), suggesting a causal effect of stress induced during the prenatal development of the offspring on cognitive outcomes. Another study used a casino opening in the United States as a natural experiment to study the effect of boosted income in an indigenous population. The study found intergenerational effects, in that a reduction in parental poverty improved some mental health outcomes of their children (Costello et al., 2003).

A major strength of natural experiments is that they allow the study of causal effects of exposures that cannot be administered in an experimental setting. However, the application of natural experiments is limited in that only specific naturally occurring exposures can be studied. Generalizability of findings is also limited (O’Connor, 2003)—for example, the effects of a policy change may be specific for a certain country or certain events might be unique, making similar follow-up research difficult. Not least, the choice of adequate control groups is challenging, as compared groups should closely resemble the exposed group and they should only differ because of the “naturally occurring” exposure. For example, the effect of a specific policy change may be difficult to isolate if other changes occurred around that time.

Regression Discontinuity

Originally introduced as an alternative to RCTs, the regression discontinuity design involves individuals that are assigned to different groups based on a predefined cutoff for an assignment variable (Thistlethwaite & Campbell, 1960), for example based on date of

Causal Inference by Using Random Variation in the Exposure

There are numerous quasi-experimental designs which rely on random variation in the exposure to study causal intergenerational effects. As we focus on observational research in this review, we only present such methods that try to achieve exchangeability by naturally occurring means.

Table 1

<table>
<thead>
<tr>
<th>Classification of Causal Inference Methods</th>
<th>Exchangeability</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Means</strong></td>
<td>(a) Capitalizing on random variation induced in the exposure</td>
</tr>
<tr>
<td>Naturally occurring</td>
<td>Natural experiments, instrumental variable analysis, Mendelian randomization, regression discontinuity</td>
</tr>
<tr>
<td>Artificially occurring</td>
<td>Randomized controlled trials, randomized experiments</td>
</tr>
</tbody>
</table>

Note. Broader definitions of natural experiments often include twin studies and other within-family designs (Rutter, 2007), which are also classified as naturally occurring here but in a separate column. In contrast to the other design-based methods, doubly robust and g-methods are analytical methods that statistically control for confounding.
### Table 2
Causal Inference Methods, Their Rationale to Meet the Exchangeability Assumption, and Important Limitations

<table>
<thead>
<tr>
<th>Type</th>
<th>Design</th>
<th>Rationale</th>
<th>Limitations</th>
</tr>
</thead>
</table>
| Random variation in the exposure | Natural experiments                 | As-if random natural variation in the exposure, for example, following a natural disaster or policy changes. Affect ed individuals are often compared with one or more control groups. Resembles a random experiment and does not require specific control for confounders. | • limited generalizability of findings  
• no available data for many exposures and outcomes  
• choice of adequate control group(s) is a challenge |
| Regression discontinuity | Comparing two groups of individuals that are “on the edge” of belonging to the other group, so that compared individuals are very similar to each other but differ in group status (e.g., in treatment status). | sensitive to misclassification  
• choice of bandwidth around the cutoff is challenging  
• treatment assignment based on more than one indicator can be problematic |
| Instrumental variable    | IV is an unconfounded proxy for the exposure, so that associations of the IV with the outcome are more likely to re flect causal effects of the exposure. | IV estimation with small sample sizes is imprecise, and with large sample sizes, bias is introduced when assumptions are slightly violated  
• less useful for cases of strong confounding  
• adequate instruments may be hard to find for most exposures |
| Intergenerational MR    | Using genetic instruments, taking advantage of random allocation of genetic variants at birth within families. | requires large sample sizes and good IVs (strong and robust associations with exposure)  
• requires genotypes of parents and offspring  
• limited generalizability of findings  
• adoption at birth necessary to exclude any postnatal nurturing effects of biological parents  
• limited generalizability of findings  
• small sample sizes |
| Control-based methods    | Adoption design                      | Effects from genetically unrelated adoptive parents on the offspring are free from genetic confounding. | cannot entirely rule out genetic confounding  
• similarities between parents due to assortative mating or other effects complicate the comparison |
| Parent comparison       | Comparing effects between genetically related parent–child and genetically unrelated parent–child dyads/trios. | risk factor is specific for one parent and the other one is used as a negative control. This might strengthen causal inference (e.g., intraterine effects like smoking, BMI, etc.), because other familial confounding is controlled. |
| Multiple relationships   | Comparing relations in nonnuclear families, including stepparents. Associations between stepparent and child are not genetically confounded, whereas associations between the child and the biological parent with whom the child does not live with mostly reflect genetic effects. | rearing effects can be underestimated, as time lived with the stepparent can substantially differ from time lived with the biological parent  
• nurturing effects of not-lived-with biological parents may not be ruled out (e.g., perinatal effects)  
• confounders that vary between children are not adjusted for  
• carryover effects (e.g., birth order) may be present |
| Sibling comparison      | Comparing an affected child (e.g., mother smoked during pregnancy) with their unaffected sibling, thus adjusting for familial confounding (shared environment and shared genetics). | confounders that vary between children, such as nonshared environment are not controlled  
• direction of causation can only be tested in longitudinal designs  
• parental exposures that do not vary within families (e.g., parental education) cannot be studied |
| Control-based methods    | Twin comparison                      | For DZ twins, this design is equivalent to the sibling comparison, but shared effects are better controlled for as twins are the same age. For MZ twins, all genetic effects are accounted for. | confounders that vary between children, such as nonshared environment are not controlled  
• direction of causation can only be tested in longitudinal designs  
• parental exposures that do not vary within families (e.g., parental education) cannot be studied |
| Children-of-siblings/children-of-twins | For MZ twins, this design compares associations in parent-offspring dyad versus uncle-offspring or aunt-offspring dyad (who share the same amount of genes with their niece/nephew as the parent, but not the family environment). Further, avuncular correlations can be compared between MZ and DZ twin families. | dyadic parental effects (e.g., divorce) cannot be studied  
• age differences between siblings and cousins need to be considered for age-dependent outcomes (e.g., cognitive performance)  
• unobserved time-varying confounding cannot be controlled  
• feedback between time-varying prenatal exposures and outcome examined after birth cannot be accounted |
| G-methods               | Generalized methods to study time-varying exposures and to account for time-varying confounding. | unobserved time-varying confounding cannot be controlled  
• feedback between time-varying prenatal exposures and outcome examined after birth cannot be accounted |
| Doubly robust methods   | Methods that include an exposure model (e.g., propensity score) and an outcome model to adjust for confounding. Doubly robust, in that it would be sufficient if one of the two models is correctly specified. | if both the exposure and the outcome models are mis-specified, results will be biased  
• when the two models are mis-specified, bias is potentially higher compared to other methods |

**Note.** IV = instrumental variable; MR = mendelian randomization; BMI = body mass index; DZ = dizygotic; MZ = monozygotic. In contrast to the other design-based methods, doubly robust and g-methods are analytical methods that statistically control for confounding.
birth as the cutoff for school eligibility, or based on a level of a biomarker as treatment eligibility. By doing this, a selection bias is created, but one that is perfectly controlled. As individuals just below and above the threshold are quasi-randomly allocated to the exposed and unexposed groups due to random variability (e.g., measurement error) in the assignment variable, those individuals are expected to have similar levels regarding all observed and unobserved confounders (Bor et al., 2014). The regression discontinuity design is also commonly used to analyze natural experiments, for example, to test causal effects of altered exposures due to policy changes.

Regression discontinuity designs using predefined cutoffs for kindergarten eligibility showed an effect of being relatively young for their grade on elevated risk of children’s ADHD diagnoses (Elden, 2010; Evans et al., 2010). To our knowledge, regression discontinuity designs have not been applied to intergenerational effects on psychological outcomes. However, Ali and Elsayed (2018) used the regression discontinuity design to examine the effects of parental education on child health. The authors compared families including parents that started school just before and after an education reform that resulted in a reduction of compulsory schooling by 1 year. They found little evidence for a causal effect of higher paternal education on offspring’s nutritional status and no evidence for an effect on child mortality or other measures (Ali & Elsayed, 2018).

The regression discontinuity design assumes exchangeability or continuity—that is, continuous variation of the assignment variable around the cutoff (Oldenburg et al., 2016). This design allows the study of exposures that are closer to “real life” than by using RCTs, but require larger sample sizes than RCTs (Bloom, 2012). Other limitations, which include the potential bias arising from misspecification of the functional form around the cutoff (e.g., assuming a linear form whereas the “true” functional form might be nonlinear), are addressed in more detail elsewhere together with arguments for the generalizability of the effects (Bloom, 2012).

**Instrumental Variables Analysis**

Instrumental variable (IV) analyses use an IV associated with an exposure of interest as a proxy to study the exposure’s effect on an outcome, and account for both observed and unobserved confounding in the exposure-outcome association (Figure 2a).

For example, educational attainment of parents was shown to influence their offspring’s educational attainment using cohorts’ finishing school during a period of unrest in France in 1968 as the instrumental variable (Maurin & McNally, 2008). The conflicts between students and universities led to a simplification of the national exams, which subsequently allowed more students to attend higher education. The authors first found increased educational attainment of this generation compared to cohorts’ finishing school before and after 1968. Strikingly, these effects were transmitted to the next generation, resulting in higher educational attainment of the students’ offspring, and providing evidence for a causal effect (Maurin & McNally, 2008).

Notably, IV analysis requires strong assumptions in addition to exchangeability (exchangeability here would mean that the IV is not associated with any confounders of the exposure-outcome association). The exclusion restriction assumption requires that there should not be any direct association between the IV and the outcome, and that the observed effect should result from the pathway through the exposure (i.e., no other pathways from the exposure to the outcome via other potential mediators). Another assumption is the relevance of the IV to the exposure, which can be tested by examining the magnitude of the association between the IV and the exposure. IV analysis is generally robust against unobserved confounding of the exposure-outcome association, but larger degrees of unobserved confounding reduce the strength of the IV and thus likely violate the assumptions of IV analysis, that is, the relevance assumption (Martens et al., 2006). Inferences from IV analysis can be biased even when assumptions are only slightly violated, or when using small sample sizes (Bound et al., 1995; Martens et al., 2006). Furthermore, it may be challenging to find appropriate instruments for most exposures, as the IV needs to be strongly associated with the exposure and exclusively associated with the outcome through the exposure of interest.

**Intergenerational Mendelian Randomization**

Mendelian randomization (MR) is similar to IV analysis and make use of measured genetic variants as a proxy for an exposure of interest (Didelez & Sheehan, 2007; Smith & Ebrahim, 2003; see Figure 2b). Individuals inherit one of two copies of each genetic variant randomly from both parents, thus enabling the use of genetic variants with established associations with an exposure as instruments for such exposure. Mendelian randomization has been widely used for causal inference and was recently extended to intergenerational settings (Davies et al., 2019; Zhang et al., 2015).

MR was applied using a genetic variant strongly associated with heaviness of smoking as proxy for maternal smoking during pregnancy, to examine the effect of maternal genotype on autism spectrum disorder and related traits of the offspring (Caramaschi et al., 2018). Consistent with other causal inference methods, the MR analysis did not support a causal effect of smoking during pregnancy on the examined outcomes (Caramaschi et al., 2018).

An advantage of MR is that the association between the genetic instrument and the outcome is largely free from environmental confounding but can be influenced by population effects including population stratification or assortative mating. Population stratification describes the fact that certain effect alleles vary in frequency between (sub-)populations, whereas assortative mating indicates that individual mating choices are not random but influenced by phenotypic similarities, for example, tall individuals choosing tall mates (Freedman et al., 2004; Yengo et al., 2018). These biases can be better accounted for in within-family MR analyses (Hwang et al., 2020). For valid inference from intergenerational MR analyses, one needs to include parental genotypes as instruments for the parental exposure and further control for the offspring’s genotype (thus requiring data from trios) to block the pathway of genetic transmission (Figure 2b; Davies et al., 2019). Intergenerational MR requires large samples, which are still rare for intergenerational settings. Further limitations include selection bias (Hwang et al., 2020) as well as general limitations of MR and IV analysis such as weak instrument bias (Burgess & Thompson, 2011).

**Causal Inference by Using Naturally Occurring Adjustment Methods**

As many exposures are not randomly distributed (e.g., whether a child’s parents are highly educated or not is usually not random), there is a need for causal inference methods to study potential effects...
of these exposures using observational data. This can be achieved by confounder control, including methods that control by matching or blocking. Within-family designs are of specific interest when investigating both genetic and environmental effects. Examples of different familial constellations and genetic relatedness in different study designs are shown in Figure 3 and described below.

**Adoption Studies**

Associations between biological parents’ traits or behaviors and the outcomes of their biological offspring might be biased by genetic factors, as parents provide both their genes and the rearing environment for their children. If both the parental exposure and the child outcome are genetically influenced, the association between them can be genetically confounded (e.g., Figure 1b). Adoption designs allow researchers to explore parent–child associations in the absence of potentially confounding effects arising from genetic relatedness. Associations between adoptive parents’ traits and the phenotype of the genetically unrelated offspring cannot result from genetic transmission and are thus more likely to reflect environmental transmission (Figure 3a).

A prospective adoption study found no effect of maternal anxiety in the adoptive mother on the anxiety symptoms of the adopted child (Ahmadzadeh et al., 2019), suggesting that previous findings of observational associations between anxiety of biological mothers and anxiety in their biological children (e.g., McClure et al., 2001) may not reflect a causal environmental effect. In contrast, adoption studies support the role of maternal depression as environmental risk factor for offspring internalizing and externalizing problems, as well as for their neurobiological development (reviewed by Natsuaki et al., 2014).

Similar to other designs, the possibility of reverse causation—that is, effects from the child on parental behavior—is a major limitation of this design unless a longitudinal approach is adopted. A specific limitation of adoption studies is that samples may not be representative of the general population, resulting in limited generalizability of findings. Furthermore, although genetic confounding can be ruled out, the design does not control for shared or nonshared environmental confounders (Thapar & Rice, 2020). Previous environmental effects from the biological parents on the offspring’s outcomes cannot be ruled out (e.g., during pregnancy or prior to adoption), which also restricts the variety of exposures that can be studied (i.e., only postnatal). However, if additional information on birth mothers and fathers prior to adoption is available (e.g., prenatal environment), maternal effects can also be examined and distinguished from the postnatal rearing environment provided by the adoptive parents. If data on both birth parents are available, prenatal environmental effects can be further distinguished from genetic effects (Loehlin, 2016), which we discuss in the section on parent comparisons.

**Assisted Conception**

Assisted conception or in vitro fertilization (IVF) can be carried out in multiple ways, so that either both parents are biologically unrelated to the child, only one parent is biologically related to the child or both parents are (Figure 3b). Similar to adoption designs, genetic confounding can be ruled out in biologically unrelated parent–child dyads or trios. If the child is carried during pregnancy by the future rearing mother, prenatal effects can be examined in addition to perinatal and postnatal effects, which expands the applicability of this design in comparison to the adoption design (e.g., information on the birth mothers may not be available in adoption studies).

Studies using the IVF design showed that maternal smoking during pregnancy is only associated with child ADHD and antisocial behavior in genetically related but not in unrelated families, suggesting genetic confounding of the association between smoking during pregnancy and child psychopathology (Rice et al., 2009; Thapar et al., 2009).
When parental risk factors are highly correlated with cannabis use during pregnancy, it could not be examined independently (El Marroun et al., 2019). Pregnancies that also had a spouse consuming cannabis, thus maternal pregnancy also had a spouse consuming cannabis, thus maternal twin pregnancies and twin comparisons indicate a blocked pathway. Genetic relatedness for the offspring generation can be inferred by path tracing, for example, for Figure 3e, the relatedness between the child and the sibling is equal to the paths via the mother (\( G_M \rightarrow G_F \rightarrow G_C \)) and father (\( G_F \rightarrow G_F \rightarrow G_C \)), resulting in a genetic relatedness coefficient of 0.5 \times 0.5 + 0.5 \times 0.5 = 0.5. Similarly, in Figure 3f the correlation between \( G_C \) and \( G_C \) is 0.5 \times 1 \times 0.5 = 0.25 when the parents are MZ twins, similar to half-siblings, and 0.5 \times 0.5 \times 0.5 = 0.125 when the parents are DZ twins or siblings; similar to typical cousins. \( C = \) child; \( F = \) father; \( G = \) genotype; \( M = \) mother; \( P_1/P_2 = \) Partner 1/Partner 2; \( S = \) sibling; \( SC = \) stepchild; \( SP = \) stepparent; \( IVF = \) in vitro fertilization; \( DZ = \) dizygotic; \( MZ = \) monozygotic.

Limitations of the assisted conception design are similar to the adoption design, mainly the extent of generalizability, as families using IVF might differ from families conceiving naturally, for example, showing higher rates of perinatal complications or lower degrees of prenatal risk exposure (Thapar & Rice, 2020). However, comparing both biologically related and unrelated families using IVF may control for confounding that might be specific for families using assisted conception. Like adoption studies, present IVF studies are limited by small sample sizes.

**Multiple Relationship Designs**

Multiple relationship designs investigate not only nuclear families but focus on families including related individuals not living together, and both related and unrelated individuals living together. Therefore, these designs allow researchers to disentangle genetic and environmental effects and can either be parent-focused—that is, multiple parenting relationships of one (step)parent—or child-focused, that is, multiple relationships between a child and their (step)parents (Figure 3d).

A study examined triparental families comprising children, their biological mother, stepfather and biological father with whom they did not live (Kendler et al., 2015). In investigating parental transmission of different externalizing behaviors, the authors found that parental effects were largest, followed by effects of not-lived-with biological fathers and then stepfathers, indicating that both environmental and genetic transmission were present (Kendler et al., 2015). Using a similar design, positive effects of higher quality relationships of both stepfather–child and nonresident (biological) father–child relations on child educational attainment were found (King et al., 2020), supporting an environmental effect of parent–child relationships on child education.
These designs are limited—somewhat similar to adoption designs—in that not-lived-with biological parents may in fact live with their offspring in early sensitive stages of their development or often have contact with their child, thus, environmental effects of those parents cannot be ruled out (Kendler et al., 2015, 2019). Furthermore, as stepparents may spend less time with the offspring (e.g., joined the family some years after the child was born), using stepparents as an index for environmental parent-offspring effects can lead to substantial underestimation of rearing effects (Kendler et al., 2015).

### Sibling and Twin Comparisons

The sibling comparison contrasts an affected child with their unaffected sibling (i.e., discordant siblings in case of binary exposure), thus controlling for familial confounding including shared genetic and environmental effects (Figure 3e). For instance, siblings may be differentially exposed to risk or protective factors like parental age, parental education, and also parenting behavior. If the outcomes of both siblings are similar, a causal effect of the specific exposure is not likely. A recent article proposed a counterfactual-based framework for the sibling comparison design and describes how a causal effect can be estimated and interpreted (Petersen & Lange, 2020).

Consistent with findings from the IVF design, studies using sibling comparisons suggest that maternal smoking during pregnancy does not causally affect child internalizing and externalizing problems, but that observed associations reflect familial confounding instead (Meier et al., 2017; Obel et al., 2016). Other studies use twin comparisons to examine the associations of differential parent–child relationships with differences in behavioral outcomes of monozygotic twins, which can rule out all genetic and shared environmental confounding (Asbury et al., 2003; Burt et al., 2006). In twin comparison studies, differential parenting behaviors such as harsh discipline have been associated with differences in externalizing and other maladaptive behaviors of the twins (Asbury et al., 2003).

Limitations of these designs are that unobserved nonshared genetic and environmental confounders (i.e., factors that are specific for each child) are not controlled for. Furthermore, the assumption that the direction of effects in parent–child associations is from the parent to the child is not always reasonable (Bell, 1968). Many studies have shown that children can influence their parents (e.g., Ahmadzadeh et al., 2019; Lifford et al., 2008; McAdams et al., 2015). The possibility of reverse causation is therefore something which should be considered, and which can make it more difficult to accurately estimate the effect of the parent on the child. However, this limitation is not unique to the sibling comparisons, but also applies for example, to the adoption design or other designs. Exploiting a longitudinal approach for example, could help to disentangle associations and provide insights in the direction of effects or potential bidirectional effects. Sibling comparisons can also be biased by carryover effects, that is, the exposure of an individual affects the outcome or the exposure of the sibling, but these can be tested (e.g., birth order effects; Sjölander et al., 2016). Furthermore, sibling comparisons cannot be used to study shared parental exposures, for example, household income or parental education, unless such factors vary over time and siblings are differentially affected at a specific age (Sariaslan et al., 2021). In contrast, twin comparisons control for more shared confounding but are even more limited in terms of the exposures that can be studied, for example, precluding the ability to study effects of the prenatal environment or of parental age at birth (D’Onofrio et al., 2014).

### Children-of-Siblings Designs

These designs use parents who are related to one another (e.g., twins, siblings, half-siblings) and their children to study associations between parental characteristics and child outcomes, using the consisbling as a matched control to enable causal inference (Figure 3f; Latvala et al., 2015; Rodgers et al., 2008). A special case of this design is the children-of-twins design (D’Onofrio et al., 2003), which we outline here in more detail. Because MZ twins are genetically identical, they are related to their cotwin’s child (their niece/nephew) to the same degree as to their own child (sharing 50% of their genes). As a result, children of MZ twins (who are cousins) are biologically half-siblings (they share 25% of their genes on average). In contrast, children of dizygotic (DZ) twins or of siblings share 12.5% of their genes on average. Genetic and environmental effects can be estimated in different ways by comparing associations between twins discordant for a phenotype and their children’s phenotype, and associations can be contrasted for children of MZ and DZ twins (D’Onofrio et al., 2003). Using structural equation modeling or other methods, the association between parental and child phenotype can be estimated, while controlling for shared environmental and genetic confounding (McAdams et al., 2018).

The children-of-twins design has been used to examine effects of parental depression on offspring psychopathology. Studies report effects of exposure to parental depression in childhood and adolescence on offspring’s depressive symptoms that remain after accounting for shared environment and genetics (McAdams et al., 2015; Silberg et al., 2010; Singh et al., 2011). However, the association between prenatal depression and subsequent child psychopathology, which has been reported in previous observational studies (Barker et al., 2008), does not remain in the children-of-twins design and might be attributable to genetic confounding (Hannigan et al., 2018).

This design does not allow causal inference when studying dyadic effects that cannot be examined for the twins independently but also involve their partners (e.g., divorce or parental conflicts; Eaves et al., 2005). Further limitations and considerations are discussed elsewhere in more detail (D’Onofrio et al., 2003; McAdams et al., 2018).

### Genetic Nurture Effects

The concept of genetic nurture describes indirect genetic effects of parental genes on child phenotypes that are assumed to work through environmental pathways, such as parental nurturing (Kong et al., 2018). Polygenic scores—summary scores that capture the genetic predisposition for a trait and include weighted effects from many genetic variants associated with that trait—can be used to infer genetic nurture effects from associations between parental genetic scores and the offspring’s outcome (Pingault, Allegrini, et al., 2022). If parental nontransmitted genes are associated with the child phenotype (i.e., ruling out genetic transmission), this effect may occur through the parental environment (Bates et al., 2018; Kong et al., 2018). Genetic nurture effects have been examined recently to investigate associations between parental genotype and offspring educational and mental health outcomes (Jami et al., 2020; Pingault, Barkhausen, et al., 2022; Wang et al., 2021). Figure 4 is an extension of Figure 3 and additionally includes environmental
parent-to-offspring effects. Furthermore, indirect genetic effects from parental genotype to the offspring’s outcome via parental phenotype can be traced in Figure 4 (e.g., GM → PM → GC). Studies designed to test for genetic nurture effects are not prone to reverse causation, as the child cannot influence parental genotypes. However, causal inference from such indirect genetic effects needs to be treated with caution, as one often cannot distinguish between population effects such as population stratification or assortative mating and genetic nurture effects (Young et al., 2019), although new methods can control for some population effects (Balbona et al., 2021; Kim et al., 2021). Furthermore, genetic nurture effects typically index a global effect without measurement of any specific parental traits (usually only a proportion of genetic variants that are associated with a specific trait are measured), research should be conducted to further characterize genetic nurture effects and to identify environmental mediators. In fact, parental phenotypic data can be integrated in novel approaches to test specific environmental pathways (Balbona et al., 2021; Kim et al., 2021), and a study following up genetic nurture effects on child depression showed partial environmental mediation of these effects by maternal anxiety and depression symptoms (Cheesman et al., 2020). Additionally, specific associations may be followed up using intergenerational MR to examine a causal effect.

Causal Inference Using Control-Based Statistical Methods

In contrast to the above-described methods, which control for unmeasured genetic and familial confounding by study design, many statistical methods aim to control for measured confounders. Among existing methods, some causal inference methods are available that have explicit assumptions for interpreting a causal effect. As mentioned, control-based statistical methods usually assume no unobserved confounding, a strong assumption that can be easily violated. Here we present some exemplar control-based causal inference methods using statistical modeling. Other methods exist such as propensity score methods (i.e., combining measured covariates into one score that captures the propensity of being exposed and adjusting for this score, e.g., by matching or weighting, Austin, 2011), which we did not address here, as they can be more biased or less efficient than g-methods or doubly robust methods (Chatton et al., 2020; Gruber & van der Laan, 2010).

G-Methods

Robins’ generalized methods (g-methods) are dedicated to the analysis of longitudinal data with multiple measurements of...
(time-varying) exposures. The g-methods include the g-formula, g-estimation of structural nested models, and inverse probability weight estimation of marginal structural models (Robins & Hernán, 2009). Mathematically, these methods use different formulae to estimate the probability densities of outcomes in the exposed and unexposed individuals, given a set of measured confounders (described in detail elsewhere, Robins & Hernán, 2009). In longitudinal designs, individuals can be compared with themselves over time, thus differences between individuals (including differences in confounders) are better accounted for. Other conventional methods, such as fixed effect models, which control for unobserved time-invariant confounding (Gunasekara et al., 2014) are likely to be insufficient to estimate an unbiased causal effect of a time-varying exposure (Robins & Hernán, 2009) as many (parental) exposures and potential confounders are not stable over time, for example, smoking status or income. G-methods can enable estimation of a causal effect of an exposure that varies over time while controlling for time-varying confounding.

For example, a recent study used g-methods (marginal structural models) to examine the effect of maternal smoking during pregnancy on offspring’s risk of depression in a sibling design and found similar results compared to generalized estimating equations, reporting an association between maternal smoking and offspring’s depression beyond confounding (Shemassa et al., 2023). Each of the g-methods has specific limitations and one or the other may be favored depending on the data structure and model of interest (Daniel et al., 2013). In general, inverse probability weighting and g-estimation are more robust than the g-formula, although it is possible that all methods converge in some scenarios (Robins & Hernán, 2009). A brief example of g-methods can be found elsewhere (Naimi et al., 2017), including references for more detailed background (e.g., Daniel et al., 2013; Robins & Hernán, 2009). G-methods can also be applied to nonvarying (time-fixed) exposures. However, it may be limited when studying time-varying prenatal exposures on child outcomes that are assessed after birth, as the feedback between exposure and outcome at different stages cannot be examined.

Doubly Robust Methods

Doubly robust methods combine two approaches to control for observed confounding, making them more robust against model misspecification (Funk et al., 2011). Confounders are included in the specification of the outcome-regression model (e.g., as covariates) and of an exposure model, for example, by using propensity scores that describe the probability of being exposed given the observed confounders (Funk et al., 2011). Both the exposure model and the outcome-regression model are prone to misspecification, but combining the two approaches enables a reliable estimation even when one of the two models is mis-specified (Bang & Robins, 2005). A commonly used doubly robust estimator is the targeted maximum likelihood estimator (van der Laan & Rubin, 2006). This method combines maximum likelihood estimation with function-based estimation, where an initial (targeted) density estimator can be updated to maximize the log-likelihood fit (van der Laan & Rubin, 2006).

A study used targeted maximum likelihood estimation to study effects of childhood adversity, providing evidence for an effect of low parental education, neglect and several other forms of adversity on lower fluid intelligence of adolescents (Platt et al., 2018). Recently, a doubly robust method has been introduced to control for confounding by cluster (Zetterqvist et al., 2016), which occurs in family designs due to shared genetics and environment. Applying this method to an existing data confirmed the results from the original study (Kuja-Halkola et al., 2014), providing no evidence of a causal effect of smoking during pregnancy on child cognitive outcomes.

In general, doubly robust methods are limited in that, under correct model specification, they can be less efficient than maximum likelihood estimation and more biased when both models are misspecified (Funk et al., 2011; Kang & Schafer, 2007). Hence expert knowledge and careful model design (including the choice of confounders) is essential.

Sensitivity Analysis

In general, sensitivity analysis is an umbrella term for various post hoc analyses to test the robustness of observed findings. Here we define sensitivity analysis—also termed bias analysis—as a method that specifically tests for potential effects of unmeasured confounding (e.g., Cornfield et al., 1959; Ding & VanderWeele, 2016; Lash et al., 2009). Control-based statistical methods, including the ones described above (g-methods and doubly robust methods), assume no unobserved confounding; however, this assumption cannot be tested and is often unrealistic. The rationale of sensitivity analysis is to test how strong the effect of an unmeasured confounder must be to change the observed finding. Combined with knowledge about effect sizes of existing confounders, one can assess the robustness of the observed association and could draw conclusions about the causal nature of the association.

A study detected an effect of higher prenatal exposure to airborne particles with a diameter <2.5 μm on worsened children’s mental and psychomotor development (Tozzi et al., 2019). A sensitivity analysis showed that findings were robust, as effects of unmeasured confounders had to be unrealistically large to erase the observed effect (Tozzi et al., 2019).

Sensitivity analyses may be limited in that they often have strong and untestable assumptions about the frequency or the distribution of the unobserved confounders, albeit some more flexible approaches exist (Ding & VanderWeele, 2016; Shen et al., 2011). Sensitivity analysis still constitutes a valuable tool and complements the above-described methods to strengthen causal inference.

Discussion

Confounding remains a major issue in observational research, including genetic confounding which might partly or fully explain observational associations in intergenerational research, highlighting the importance of causal inference methods. We have described a broad range of methods for causal inference in intergenerational settings, which aim to achieve exchangeability by either (a) capitalizing on random variation in the exposure or (b) adjusting for confounding resulting from nonrandom variation in the exposure. We elaborated on the advantages and limitations of each design and provided an overview of existing methods with references to potential designs for specific research questions.

As each method described here has its own limitations, we advocate combining more than one approach to infer causality. Where feasible, different approaches should be combined in the same study (Caramaschi et al., 2018; Liu et al., 2021), for example, using designs with random variation in the exposure and control-based methods,
performing sensitivity analyses and including negative controls (e.g., pseudo-outcomes; Imbens, 2015). Triangulating evidence from independent studies, that is, using different methods with distinct assumptions to test the same hypothesis, is pivotal to infer causal effects (Lawlor et al., 2016; Munafò et al., 2021; Munafò & Smith, 2018), and no method should be seen as sufficient on its own. Coming back to an example from the introduction (Figure 1b) highlights how a variety of different methods jointly provide insights into the causal relations between intergenerational risk factors and psychopathology. For example, within-family designs using IVF (Thapar et al., 2009), parent (Gustavson et al., 2017; Langley et al., 2012), or sibling comparisons (Knopik et al., 2016; Kuja-Halkola et al., 2014; Skoglund et al., 2014), as well as studies using genetic instruments (Haan et al., 2021) showed that smoking during pregnancy seems to not causally affect child ADHD, but rather reflects familial confounding. Triangulation can also be obtained via systematic reviews and meta-analyses of various genetically informed methods with different assumptions, which have provided evidence for environmental effects of parental anxiety on child internalizing problems (Ahmadzadeh et al., 2021), of prenatal alcohol exposure on child cognitive outcomes (Mamluk et al., 2020), and of several parental risk factors on child mental health and related outcomes (Jarii et al., 2021) beyond genetic confounding.

**Outlook**

The multidisciplinary nature of intergenerational research holds promise for the development of novel methods to further strengthen causal inference. In general, cooperative research efforts are crucial to maximize insights by increasing sample sizes, and by gathering collective expertise across disciplines. However, smaller but more nuanced studies, which include detailed and harmonized phenotyping and longitudinally examine the samples, are also of importance and constitute a large part of the research cited here. In this review, we only focused on studies and methods that test direct effects of parental exposures on child outcomes and did not address mediating or moderating factors, which is a limitation. Identifying mediating pathways from parental (risk) factors to child outcomes or moderating factors of parental effects is clearly important. For example, mediating factors can be examined using causal mediation analysis (Daniel et al., 2015; Imai et al., 2010), and moderators could be tested using a recently proposed framework for causal moderator analysis based on potential outcomes (Dong et al., 2022). In the following sections, we outline some future directions and promising, potential methodological advances for intergenerational causal inference.

**New Data for Intergenerational Research**

Valuable data for intergenerational research are increasingly evolving, with multigenerational data on a national scale becoming available in terms of birth registries (Van Der Wel et al., 2019), electronic health records (Friedman et al., 2013), or data on education or criminal records. Combining these data provides a tremendous opportunity to gather information about parental characteristics and child outcomes. Furthermore, new data (e.g., through genotyping) are emerging and could be matched with other registries, providing an important basis for the application of MR or the estimation of genetic nurture effects. Other collaborative efforts using large within-family cohorts will result in publicly available summary statistics that can be used for such purposes (Howe et al., 2022). Combining different registry data implies that comprehensive data in a “natural setting” can be used, which is precisely timed and documented, while being representative of the target population. Hence, these data can further be linked to certain events or time periods, such as policy changes, natural disasters such as flooding or pandemics such as COVID-19. However, such detailed data on individuals raise important questions of data privacy, which have to be carefully addressed (e.g., Behrendt et al., 2018).

**Spillover From Related Fields**

We have seen that certain causal inference methods are underrepresented in intergenerational psychiatry and psychology, such as regression discontinuity or instrumental variable designs, although both are strong designs for causal inference (Kim & Steiner, 2016). This may result from a cautious attitude of researchers in psychology and psychiatry toward causal inference (Grosz et al., 2020; Rutter, 2007), potentially coupled with a lack of statistical training in such methods. It has been argued that causal relations are often implicitly assumed by researchers but not explicitly expressed; however, the expression of such assumptions is essential to choose adequate methods and to address their specific biases (Goetghebeur et al., 2020; Grosz et al., 2020; Pingault, Richmond, & Smith, 2022). As mentioned, applying these designs can also be challenging as appropriate instruments might not be available for certain research questions.

Regression discontinuity designs following policy changes were used to examine causal effects of education or alcohol consumption on mental health outcomes (Courtin et al., 2019; Ertan Yörük & Yörük, 2012). Future studies could expand the analysis to the offspring generation of those individuals to study causal effects of parental characteristics on child psychological outcomes as in other designs (Costello et al., 2003; Maurin & McNally, 2008).

**Blurring Boundaries Between Classifications**

Although in this review we described methods based on distinct categories (Table 1), such categories are not exclusive. First, methods capitalizing on random variation in the exposure (Column 1 of Table 1) may further statistically adjust for confounders (Row 2, Column 2). This has been suggested, for example, for natural experiments (Dunning, 2008) and is common in research using instrumental variables (Vansteelandt & Didelez, 2018). For example, multivariate MR can also be seen as a method to control explicitly for suspected pleiotropic pathways (Burgess & Thompson, 2015). Second, methods using naturally occurring settings to control for confounding (Row 1) may further adjust for confounders to approach conditional exchangeability. This can be done, for example, to control for measured nonshared environmental confounders in twin comparison studies, in addition to the adjustment permitted by the natural design (Row 1, Column 2). Furthermore, researchers may be involved more tightly in the planning phase of policy changes (Column 1) to assess data on potential confounders of individuals likely to be affected by the “natural experiment” in a pre-post design.

**Advances in Genetically Informed Designs**

There is growing interest in and potential of using genetic data in observational studies to strengthen causal inference (Pingault et al., 2018). For instance, a sensitivity analysis has been introduced which
controls for genetic confounding in observational studies with available genotype information of the participants (Pingault et al., 2021). Causal inference methods like the regression discontinuity design could be used to study causal effects of selective prevention programs for children, using parental risk factors as assignment variable (e.g., indexed by genetic instruments). Comparing individuals whose parents score just above or below the threshold (say, “mean genetic risk”) would allow us to infer a causal effect of the preventive intervention. Unlike in an RCT, individuals of higher risk would thus all be eligible for the program, instead of using a random assignment.

Alternatively, instead of contrasting individuals with similar values on the assignment variable, recall-by-genotype designs compare individuals at the extreme ends of low versus high genetic risk for a disease. Similar to MR, these designs exploit the random allocation of genetic variants at conception for causal inference (Corbin et al., 2018). Unlike MR, recall-by-genotype studies only focus on a smaller subsample, which can be examined in detail to understand the causal pathway to disease. A translation to the intergenerational setting could be to screen parents regarding their genetic risk toward a particular disorder, and to follow up on their children’s outcomes. Thus, one could examine the effects of parental risk factors on the offspring’s outcomes in a smaller subsample, allowing deep phenotyping of parents and offspring. Similar to intergenerational MR, adjusting for offspring genotype would be necessary to exclude genetic transmission.

There is a large variety of possible methodological extensions using genetic data to study causal environmental effects. However, as genetics represent sensitive individual data, recall-by-genotype, and other designs using genetic screening need careful study design, high transparency, and good communication of the study aims, the meaning of genetic risk, and consequences for the individuals (Beskow et al., 2010; Corbin et al., 2018), especially when children are involved.

Conclusion

The rich toolbox of causal inference methods for observational studies can also be leveraged in intergenerational settings and so far, has successfully contributed to distinctions between putative causal risk factors and confounded associations. Researchers should consider the numerous available designs and methods when conceiving studies to investigate parental factors in observational research and exploit methods from related fields beyond their current repertoire. Besides the broad range of existing methods, promising new approaches to strengthen causal inference are to be expected. Eventually, observational research on intergenerational effects may inform the design of appropriate preventive interventions. Converging evidence suggesting that observed effects of putative parental risk factors are explained by confounding will have practical implications, for example, redesigning interventions to target factors other than those previously considered.

References


Seeman, M. V. (2016). Schizophrenogenic mother. In J. Lebow, A. Chambers, & D. C. Breunlin (Eds.), *Encyclopedia of couple and family therapy* (pp. 1–2). Springer International Publishing. https://doi.org/10.1007/978-3-319-15877-8_482-1


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