Title
Associations between parent anxiety and offspring internalising problems: A systematic review and initial meta-analysis of genetically informed research

Running title
Parent anxiety and offspring internalising

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Abstract

Objective

Parent anxiety is associated with offspring internalizing problems (emotional problems related to anxiety and depression). This may reflect causal processes, whereby exposure to parent anxiety directly influences offspring internalizing (and/or vice versa). However, parent-offspring associations could also be attributable to their genetic relatedness. We present a systematic review and meta-analysis to investigate whether exposure to parent anxiety is associated with offspring internalizing after controlling for genetic relatedness.

Method

A literature search in five databases identified 429 records. Publications were retained if they used a quasi-experimental design in a general population sample to control for participant relatedness in associations between parent anxiety and offspring internalizing outcomes. Publications were excluded if they involved an experimental exposure or intervention. Studies of pre- and post-natal anxiety exposure were meta-analysed separately. Pearson’s correlation coefficient estimates ($r$) were pooled using multilevel random effects models.

Results

Eight publications were retained. Data were drawn from four population cohorts, each unique to a quasi-experimental design: adoption, sibling-comparison, children-of-twins or in-vitro-fertilisation. Cohorts were located in northern Europe or America. Families were predominantly of European ancestry. Three publications ($N_{\text{families}}$>11,700; offspring aged 0.5—10 years) showed no association between prenatal anxiety exposure and offspring internalizing outcomes after accounting for participant relatedness ($r$.04, CI -.07,.14). Six publications ($N_{\text{families}}$>12,700; offspring aged 0.75—22 years) showed a small but significant association between concurrent symptoms in parents and offspring, after accounting for participant relatedness ($r$.13, CI .04,.21).

Conclusions

Initial literature, derived from homogenous populations, suggests that prenatal anxiety exposure does not cause offspring internalizing outcomes. However, postnatal anxiety exposure may be causally associated with concurrent offspring internalizing, via non-genetic pathways. Longitudinal stability, child-to-parent effects, and the role of moderators and methodological biases require attention.
Lay summary

Anxiety problems are known to run in families. This could be because parents and offspring directly influence the development of symptoms in one another, and/or because parents and offspring are genetically related. We conduct a systematic search for all studies examining associations between parent anxiety and offspring emotional (i.e., internalizing) problems, where authors have sought to account for the influence of genetic transmission in families. Together, data from three studies show that foetal exposure to maternal anxiety is not associated with child internalizing outcomes after accounting for genetic transmission. Together, data from six studies show that child exposure to parent anxiety after birth is associated with child internalizing problems at the same timepoint, even after accounting for the effects of genetic transmission. More research is needed to build on this initial literature, to examine whether non-genetic processes endure across time, and to test the extent to which child symptoms may influence parent symptoms.

Clinical guidance

- **When working with pregnant women who are experiencing anxiety symptoms:**

  Evidence to date suggests that mothers’ prenatal anxiety symptoms do not cause the development of similar symptoms in child offspring. This may be reassuring for parents to hear. Evidence suggests that genetic transmission can explain associations between mothers’ prenatal anxiety symptoms and offspring internalizing outcomes. Continued exposure to parent anxiety after birth may also play a role. As such, clinicians should consider ways to prevent or alleviate symptoms in offspring who inherit higher genetic risk propensity for internalizing problems; and consider ways to prevent or alleviate the continuation of parent symptoms postnatally.

- **When working with parents who are experiencing anxiety symptoms as children grow up:**

  Evidence to date suggests that parents’ anxiety symptoms may be causally associated with internalizing problems in their offspring at the same point in time. This contrasts with research on the prenatal period. However, we still cannot tell whether parents’ symptoms exert palpable, lasting influence on offspring internalizing outcomes. We encourage clinicians to take a holistic view: considering the influence of genetic transmission; the influence of child symptoms on parents; the influence of parent symptoms on children; and the influence of the shared family environment. To prevent the development of child symptoms, it may not be sufficient to only treat parent symptoms. Parents may be reassured to learn this.
• *Considering the generalisability of findings:*

All robust, genetically informed research to date on this subject has been conducted in northern Europe or America, with families who are of predominantly European ancestry. Furthermore, research has been predominantly conducted with mothers. It is important for clinicians to note that findings may differ across other samples and that we know very little about the role of fathers’ anxiety in families.
Introduction

Anxiety disorders are the most prevalent class of mental disorders worldwide.¹ They are characterised by pervasive emotional and physical distress that can substantially restrict daily functioning. The median age of onset for anxiety disorders is 11 years.² Anxiety symptoms and disorder diagnoses cluster within families, with disorder status among parents being a robust predictor of related internalizing problems among developing offspring.³⁻⁵ Internalizing problems encapsulate emotional symptoms characteristic of both anxiety and depression. Core internalizing symptoms include worry, fear, sadness, and withdrawal.⁶ Parent anxiety could pose an environmental risk for the development of offspring internalizing problems, for example via modelling processes and social learning,⁷⁻⁹ or via foetal programming during pregnancy.¹⁰⁻¹³ It is also possible that child symptoms influence parents’ symptoms, resulting in environmentally mediated transactional effects between parents and offspring.¹⁴⁻¹⁶ However, genetic transmission from parents to offspring is likely to at least partially account for symptom associations across generations. It is important to distinguish the potential environmental effects of familial exposure from associations attributable to genetic relatedness in families, to inform the development of successful intervention and prevention strategies.

Previous research on associations between parent anxiety and offspring internalizing outcomes has mostly relied upon observational studies, where researchers may adjust for measured confounding but cannot account for unobserved variables, including genetics. e.g., ¹⁷⁻²⁰ This is a major limitation that leads to ambiguous results, where potential causal pathways operating between parents and offspring (i.e., if parents’ symptoms directly influence child symptoms, and vice versa) are indistinguishable from influence by other common causes (e.g., if the same genes influence symptoms in both parents and offspring). Population variance in anxiety and related internalizing problems is attributable in part to genetic influences,²¹⁻²³ with the same genetic factors found to act in multiple internalizing phenotypes across the lifespan.²⁴⁻²⁶ It is therefore reasonable to expect that genetic factors influencing anxiety in adult parents may also act in their genetically related offspring, manifesting as similar problems during childhood. This results in ‘passive gene-environment correlation’, whereby the environment shared by parents and offspring is passively correlated with the genes that they share.²⁷ Across time, it is possible for genetically influenced behaviours in parents and offspring to evoke changes in one another, as exemplified by longitudinal studies showing dynamic, transactional processes between family members.¹⁴⁻¹⁶ Together, existing evidence from genetic and longitudinal research suggests that purely observational studies cannot provide robust conclusions as to whether and how parent anxiety might directly influence the development
of child internalizing problems. It is important to first control for confounding by shared genes, then ask questions about the direction of effects between generations.

**Experimental research**

To explore the causal pathways linking parent anxiety and offspring internalizing, researchers can use either experimental or quasi-experimental methods. Experimental methods are used in medical research to test the effect of an exposure (or intervention) on an outcome, with randomised controlled trials (RCTs) typically labelled the ‘gold standard’. It is unfeasible and unethical to experimentally randomise children to be reared by parents who are versus are not experiencing anxiety symptoms. Instead, researchers can temporarily exacerbate anxiety in parents or offspring, using controlled experiments, to examine whether increases in symptoms in one generation predict the same in the other. For example, parents’ fear responses to a novel toy can predict offspring fear and avoidance behaviours.\(^{28}\) Another experimental approach involves researchers treating parent or offspring symptoms and examining whether symptom improvements in one generation predict improvements in the other. RCT data shows that treatment of child anxiety is associated with a reduction in parent anxiety.\(^{29,30}\) Treatment of parental anxiety may also help to improve child outcomes, in combination with child-focused treatment.\(^{31}\) although evidence is not always consistent.\(^{32,33}\) In sum, experimental research provides some evidence for causal effects between parent anxiety and offspring internalizing symptoms, but they cannot inform on the nature of the association outside of the experimental setting, nor on the influence of genetic transmission in naturalistic settings. In the context of epidemiological research, experiments are limited because they tell us how things can be, rather than how they are in a population.

Quasi-experiments provide an alternative approach to experimental trials, which can be used to test the possibility that causal mechanisms underlie associations between an exposure and an outcome. Within a quasi-experiment, the exposure of interest (e.g., parent anxiety) is naturally occurring and not manipulated by the investigator. Unique design features are used to account for unmeasured variables that are confounded with the hypothesised causal environment, such as genetic relatedness between parents and offspring, to strengthen causal inferences.\(^{34,35}\) A range of genetically informed quasi-experimental research designs have been developed for this purpose, comparing family members for whom genetic relatedness is known or can be approximated.\(^{36}\) Family types integral to these designs include: parents with adopted children; parents with children conceived via gamete or embryo donation; identical and nonidentical twin pairs with children; and parents with two or more children who are differentially exposed to the variable of interest. These designs help to tease apart the role of genetic and
environmental transmission effects across generations. When combined with longitudinal data, they can also shed light on the direction of effects between generations.

Genetically informed quasi-experiments

Adoption and in vitro fertilisation (IVF) designs

When used for quasi-experimental purposes, adoption designs require data from parents and offspring adopted at birth. In such studies, similarities between birth parents and the adopted child reflect only genetic and prenatal influences, because postnatal contact is absent (this means that passive gene-environment correlation cannot occur). Similarities between adoptive parents and their unrelated adopted child are free from confounding by genetic relatedness (again, passive gene-environment correlation cannot occur), so are used to examine influence of the rearing environment on child development, independent of genetic effects. The IVF (or ‘cross-fostering’) design follows a similar premise for offspring conceived via gamete or embryo donation, who are essentially ‘adopted at conception’. Where embryos are implanted into unrelated mothers, the IVF design can be used to distinguish the influence of both pre- and postnatal rearing environments from genetic effects. Longitudinal data using both ‘adoption at birth’ and ‘adoption at conception’ study designs allow for examination of environmentally mediated effects between generations, including parent-to-child and child-to-parent effects. Here, results can quantify the extent to which genetically influenced child traits evoke change in parent traits (referred to as ‘evocative gene-environment correlation’), which would increase parent-offspring trait associations. A major advantage of adoption designs is that they eliminate any effect of shared genes in associations between parents and offspring by design, because they use genetically unrelated parent-offspring dyads. However, a caveat to this is that participants in adoption and IVF studies may not be representative of the general population, so results may not be generalisable and sample sizes are usually small. Further, it is necessary to examine and control for adoption openness (i.e., the degree of contact and knowledge between birth and adoptive families) and the possibility of selective placement (i.e., when birth or donor parent characteristics are matched with adoptive parent characteristics), as these can violate the design assumptions and bias results by making adopted children more similar to their birth or donor parents.

Children-of-twins designs

Children-of-twins designs require data on identical (monozygotic, MZ) and fraternal (dizygotic, DZ) adult twin pairs with children. In MZ twin families, offspring are just as genetically related to their parent’s identical twin, who is their aunt or uncle, as they are to their own parent (genetic correlation=.50 with both individuals). In
DZ twin families, offspring are less genetically related to their aunt/uncle (genetic correlation=.25 on average) as compared to their own parent (genetic correlation=.50). As such, if offspring are more similar to their aunt/uncle for any given trait in MZ compared to DZ families, an effect of shared genes is indicated. Following this logic, researchers statistically estimate and control for the role of shared genes between generations (i.e., controlling for the effects of passive gene-environment correlation). Residual parent-child associations are unconfounded by genetic relatedness and comprise the environmental effect of parents on offspring, and vice versa, as well as any unmeasured confounding (i.e., additional confounding that is not captured by controlling for shared genetic influences). Extensions of the design can include more than one child per parent and different combinations of adult siblings, including half siblings and unrelated sibling-in-laws.\textsuperscript{44-46} It is also possible to model the influence of environments shared across all members of nuclear/extended families. Analyses are reliant upon an ‘equal environments assumption’, positing that the children of MZ twins are not exposed to their parent’s twin any more so than the children of DZ twins (which has been found to hold true in previous research).\textsuperscript{47} Children-of-twins data have not yet been modelled longitudinally in research and it is possible for the effects of evocative gene-environment correlation to inflate estimates of the parent’s causal influence on the child.

\textit{Sibling-comparison designs}

Sibling-comparison designs are unique in that they do not rely upon differentially related individuals, but instead on siblings who are differentially exposed to a given environment. Specifically, sibling-comparison designs require a sample of parents with at least two children, where sibling differences exist for the independent variable/exposure (e.g., prenatal maternal anxiety).\textsuperscript{48} Siblings are naturally matched into family units where they broadly share many potential confounding variables, including their parents, home and family environment, and genetic factors (genetic correlation between siblings=.50 on average). Researchers compare groups of differentially exposed, family-matched siblings on an outcome of interest (e.g., internalizing problems), to examine the exposure effect whilst eliminating within-family confounding. It can be assumed that genetic risk transmission is equal between siblings at a population level, on average, given the random nature of inheritance. As such, researchers simultaneously control for both unmeasured genetic and environmental confounding in families (this controls for more than just passive gene-environment correlation in families, as all aspects of the siblings’ shared environment is controlled for). Typically, in sibling comparison methodology no distinction is drawn between parent-to-child and child-to-parent effects and it is assumed that siblings do not significantly influence one-another.\textsuperscript{49} Again, it is possible for the effects of evocative gene-environment correlation to inflate estimates of the parent’s causal influence on the child.
Additional sources of confounding

The discussed quasi-experimental designs account for genetic confounding in different ways and require specific sub-populations of families on whom phenotypic data has been collected. These designs cannot control for all potential confounds and each design is characterised by a different set of methodological caveats and assumptions. Across all designs, shared method variance can arise when parents report on both their own and their offspring’s symptoms, thereby inflating estimates of intergenerational associations in research. Further, the length of time elapsed between measurement of the exposure (parent anxiety) and outcome (child internalizing) can influence results, with concurrent associations typically being stronger than those with a lagged outcome.\textsuperscript{50,51} Results may also differ depending on child age or developmental period, participant sex, socio-economic status, presence of comorbid diagnoses, or reliability of the measures used for data collection. As such, it is important that researchers consider both measured and unmeasured confounders, whilst drawing on a range of quasi-experimental research designs and protocols, to yield reliable and robust conclusions.\textsuperscript{52}

Aims

We conduct the first systematic literature review to identify all existing empirical research where authors have accounted for familial genetic confounding in associations between parent anxiety and offspring internalizing outcomes. We focus on quasi-experimental research. We exclude studies that involve experimental manipulation of anxiety state within families and observational research where controls are not included for unobserved sources of confounding. Results relating to prenatal and postnatal parent anxiety exposure are investigated separately, as they relate to distinct forms of anxiety exposure, with distinct hypothesised modes of transmission.\textsuperscript{10} We provide a narrative synthesis, critique, and meta-analysis of the retrieved literature to date. Our primary aims were to examine the following questions:

a) Is parent anxiety associated with offspring internalizing outcomes after accounting for familial genetic confounders?

b) If so, what can we tell about the direction of effects between parents and offspring?

c) If extracted data permits further analysis of moderator terms, to what extent is the magnitude of the parent-child association affected by methodological (e.g., study design, reporter, time-lag between exposure and outcome) and/or observed (e.g., sex, age, socio-economic status, comorbid parent depression, obstetric complications) covariates?

Method
Search strategy

Our methods were registered in advance using the International Prospective Register of Systematic Reviews (PROSPERO; protocol number: CRD42019134977). Our search was conducted between July – September 2019, using Web of Science and Ovid (Embase, MEDLINE, Global Health, PsycINFO). The search was restricted to articles published in English. The following search terms were used to identify papers examining parent anxiety (NB., parent terms and anxiety terms were combined to restrict the number of search results, see Supplement 1 for full search strategy, available online):

- mother* or matern* or father* or patern* or parent* or *natal
  AND
- anx* or phobi* or “social* anx*” or “general* anx*” or neurotic* or obsessive* or panic or agoraphobi*

The following terms were included to identify papers that examined offspring outcomes and those that used a quasi-experimental design to control for potential genetic confounding (NB., to ensure that we identified all possible internalizing outcomes examined to date, we did not restrict the search to pre-defined internalizing outcomes):

- child* or adolescen* or teen* or youth* or young or offspring or infan*
  AND
- twin or twins or sibling* or adoption or adopted or “in vitro fertilization” or “assisted conception” or “cross-fostering” or “instrumental variable” or “quasi-experiment*” or causa* or genes or genetic or geno* or heritab*

The abstracts of all returned papers were screened independently by YA and MH. Studies were excluded if there was clear evidence that criteria were not met, with agreement from both researchers. The reference lists of relevant review papers were screened to identify any articles that were missed from the search, and further searches were made to identify published manuscripts from the authors of relevant conference abstracts. Full text screening for all retained studies was conducted by YA and MH, independently, to confirm eligibility. Disagreement was resolved through discussion with the senior researcher, TM. Data extraction from studies to be included in the meta-analysis was conducted by YA, checked by TM and TS.
Study selection

Published studies presenting empirical research were included if they:

- involved a population of human parents and offspring (no sex or age restrictions)
- examined associations between parent anxiety (measured at trait, symptom or disorder level) and offspring internalizing outcome/s (relating to withdrawal, somatic complaints, anxiety, depression)\(^5^3\)
- used a natural quasi-experimental research design to account for genetic relatedness in associations between parents and offspring (i.e., intergenerational genetically informed research designs, which enable researchers to control for participant relatedness)\(^3^6\)

Our search terms identified some studies of parental ‘stress’. We considered these to meet inclusion criteria if exclusively measuring feelings of stress (i.e., anxiety symptoms), not stressful life circumstances. Further, our search terms identified publications using ‘candidate gene’ approaches (using parent and child DNA) to control for the influence of specific shared genes in parent-offspring associations. Mental health phenotypes are typically classified as complex traits; i.e., they are polygenic, influenced by hundreds of thousands of genetic variants across the genome, each exerting a very small effect.\(^5^4-5^6\) Therefore, studies accounting for the transmission of only a handful of genes (i.e., ‘candidate genes’) between generations are insufficient to control for genetic confounding in parent-child associations for mental health phenotypes.\(^5^7\) The only genomic studies that thus met our inclusion criteria would be those taking a genome-wide, polygenic approach to quantifying intergenerational genetic relatedness.

Studies were excluded if they:

- focused exclusively on populations with specific physical health problems (e.g., cancer, seizures, low gestational age) or a diagnosed developmental disorder (i.e., communication or learning disorders, motor disorders, attention-deficit/hyperactivity disorder, or autism spectrum disorders)
- involved an experimental exposure or intervention

Figure 1 presents the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flowchart. Of the 441 records screened, eight publications met our inclusion criteria.\(^3^9,4^0,5^1,5^8-6^2\) Of the excluded
publications, seven included design features that accounted somewhat for bias by genetic confounding.63-69 These studies and our reasons for their exclusion are outlined in Supplement 2, available online. In brief, they were studies of prenatal anxiety exposure that used paternal anxiety symptoms as a ‘negative control’ for environmentally mediated prenatal transmission; and/or examined child exposure to parent state-level (i.e., current, transitory) anxiety symptoms, while controlling for parent trait-level (i.e., stable, longer-term) symptoms. We determined that they did not meet our criteria for a robust method to “account for genetic relatedness in associations between parents and offspring”.

Data extraction

Data relating to sample characteristics, measurement protocol and statistical analyses were extracted from each publication that met our inclusion criteria. All genetically informed effect estimates were initially extracted. Where authors published multiple effect sizes for the same set of variables (e.g., where multiple analytic strategies were explored), we then had to decide which estimates to include in the meta-analysis to derive a meaningful pooled result. For the three publications examining prenatal anxiety exposure, one examined both continuous and binary coded data.59 We selected results based on continuous scores, to be consistent with the two other publications.51,58 Two publications examining prenatal anxiety exposure reported separate effect estimates from analyses before and after adjusting for postnatal anxiety exposure,58,59 while the third reported only adjusted results.51 All effect estimates were retained, to explore comparison of results that did versus did not attempt to isolate the effects of prenatal from postnatal exposure.

For the six publications of postnatal anxiety exposure, two reported both bivariate correlations and beta estimates from structural equation models (i.e., partial correlations) for the same set of variables, involving both cross-sectional and longitudinal datapoints.39,40 It was not possible to extract data from the structural equation models in an informative way for inclusion in the meta-analysis, because saturated results were not presented in either publication (i.e., some paths had been trimmed from the models). Of the bivariate correlation analyses, longitudinal correlations (i.e., between parent anxiety exposure and future internalizing outcomes in offspring) were not informative on their own as prospective associations, because they did not include correction for concurrent exposure to parental symptoms. As such, only the cross-sectional bivariate correlations were selected from these publications for inclusion in the meta-analysis. Cross-sectional effect estimates were available for inclusion in three of the four remaining publications examining postnatal anxiety exposure. For the one publication
using only longitudinal data, the effect size with the shortest exposure-outcome time-lag was selected (9 months), to conserve consistency in the meta-analysis. Two remaining longitudinal effect estimates (derived from two publications) were subsequently excluded, given that so few longitudinal estimates would be uninformative in the meta-analysis.

**Effect size calculations**

Pearson’s correlation coefficient, $r$, was used as the uniform effect size across all studies, with confidence intervals computed for each estimate using the R package `compute.es`. Pearson’s $r$ is an appropriate effect size to use for associations between continuous variables and results are easily interpretable. Non-independent effect sizes derived from the same or overlapping samples in a single publication (e.g., effect sizes at different child ages for families in a single cohort) were aggregated using the R package `MAd` (for meta-analysis with mean differences), to account for their correlation. Aggregation of correlated (i.e., non-independent) effect sizes within publications is required to prevent over-estimation of the precision of the pooled effect size in meta-analysis, which occurs when findings based on the same data are incorrectly treated as unique. Aggregation of effect sizes within publications also prevents studies with more effect estimates from being given more weight in meta-analysis. Multilevel models are not appropriate to account for non-independence of effect size estimates within a single publication.

Non-independent effect sizes within each publication were aggregated first to create pooled effect sizes per publication ($r_{\text{publication}}$). In sensitivity analyses we aggregated non-independent effect sizes within each cohort, to create pooled effect sizes per cohort ($r_{\text{cohort}}$; see Figure S1 [available online] for an example, depicting the aggregation of 14 effect sizes across four publications, all derived from one cohort). Aggregation of non-independent effect sizes requires specification of their correlation. The magnitude of their correlation depends on the degree of overlap in the population sample, measures and timepoints used for each estimate. As is typically the case in a meta-analysis, the correlations between dependent effect sizes within each publication, and between publications, were unknown, meaning that we had to specify a likely value. Given the potential for this specified correlation to impact results, we conducted three sensitivity analyses for each aggregation, testing results using a full range of possible correlations for the association between dependent effect sizes: $r=.10$, $.50$ and $.90$.

**Random Effects Models (REMs)**
Meta-analytical models were conducted as multilevel random effects models (MREMs) using the R package \textit{metafor}.\footnote{75} MREMs allow for between-study heterogeneity and can be used to test for moderating effects when data permits (i.e., methodological and observed covariates). First, an MREM was used to pool Pearson’s $r$ effect sizes from each publication ($r_{\text{publication}}$) examining concurrent anxiety exposure. In this model, a source of variation was introduced for each cohort, to account for random variance (i.e., higher order clustering) between cohorts, and for each publication within each cohort. Next, in a sensitivity analysis, we conducted a standard REM to pool aggregated, independent Pearson’s $r$ effect sizes from each cohort ($r_{\text{cohort}}$). These results are not biased by non-independent effect sizes, although they eliminate any information on the effects of moderating terms on the magnitude of associations within cohorts, relating to study design and sample characteristics in each publication.

Heterogeneity between effect sizes was assessed using the $I^2$ statistic, to examine whether study characteristics moderated the pooled effect size.$^76$ The $I^2$ statistic is the percentage of total variation in study estimates that is due to heterogeneity, or between-study variability (values $<25\%$ indicate low; $25\%$-$75\%$ moderate; and $>75\%$ considerable heterogeneity). Publication bias was evaluated visually using funnel plots, plotting effect sizes against their standard errors. Symmetrically distributed data points indicate absence of publication bias. The low number of included studies yielded insufficient statistical power to test for asymmetry using Egger’s linear regression.$^77$

\section*{Results}

\subsection*{Study descriptions}

The eight retrieved papers were published between 2010 – 2019, using data derived from four independent cohorts located in northern Europe and America ($N_{\text{total}}=12,990$).\footnote{39,40,51,58-62} Each cohort had one quasi-experimental research design applied (adoption,$^39,40,60,61$ IVF,$^58$ children-of-twins,$^62$ sibling-comparison$^{51,59}$) and was mostly restricted to the study of one developmental period (infancy,$^40,51,59,61$ middle childhood$^39,58$ or adolescence,$^62$ Tables 1–2). The sibling-comparison sample (derived from the Norwegian Mother, Father and Child Birth Cohort Study; MoBa) was far larger than all other samples combined. As shown in Tables 1–2, available information suggested that $>90\%$ of the rearing parents examined across publications were of European ancestry. No information was available for participant ancestry in the MoBa, although data from Statistics Norway suggest that approximately $90\%$ of the Norwegian population had Norwegian born parents and grandparents the year that
MoBa recruitment ended (2009), indicating that MoBa participants would be predominantly of European ancestry. Ancestry data were ambiguous for participants in the children-of-twins sample (the Twin and Offspring Study in Sweden, TOSS; see footnote of Table 2), although again evidence suggests that participants were predominantly of European ancestry.

Parent anxiety and offspring internalizing symptoms were measured along continuous scales in all analyses extracted for the meta-analyses. All publications used correlation coefficients and/or beta estimates to evaluate intergenerational covariance (see Tables 1–2). Parent anxiety was measured by self-report, using five different measures of adult anxiety across publications. Seven child internalizing constructs were assessed across the publications (e.g., combined internalizing, negative affect, anxiety, social inhibition). Parents contributed at least partially to child symptom scores in all publications except one (where child social inhibition was measured solely by researcher observations). Results derived from the sibling-comparison or IVF datasets were subject to the greatest risk of shared method variance, because only mothers’ reports were used to construct variables in these cohorts. Two publications (each using the same adoption sample at different developmental stages) examined the directionality of effects between generations and analysed mother-child and father-child associations separately.

A range of different measured covariates were accounted for in analyses, each attenuating the crude parent-offspring correlation to varying degrees.

**Meta-analysis**

**Prenatal anxiety exposure**

MREM results showed a negligible and nonsignificant pooled effect size between prenatal anxiety exposure and infant internalizing outcomes, using data from three publications (Nfamilies >11,700; offspring aged 0.5—10 years) that were corrected for genetic confounding and exposure to postnatal anxiety (r=.04, 95% CI -.07,.14; Figure 2A). Pooled estimates were equivalent in REM analyses using aggregated cohort data (Figure 2B). Two publications provided results that were unadjusted for postnatal anxiety exposure. REM analyses of these estimates revealed the pooled effect size to be larger than those using adjusted estimates, but still non-significant (r=.11, 95% CI -.05,.28). Because there were only three publications examining prenatal anxiety exposure, statistical power was insufficient to test for heterogeneity of effect sizes.

**Postnatal anxiety exposure**
MREM results showed a significant pooled effect size between concurrent anxiety exposure and offspring internalizing outcomes, using data from six publications (N\textsubscript{families} > 12,700; offspring aged 0.75—22 years) that were corrected for genetic confounding (r=.13, 95% CI .04, .21; Figure 3A). Results were comparable in analyses using effect sizes aggregated by cohort (Figure 3B). Results showed substantial levels of heterogeneity between publications (F=90, suggesting that 90% of the x\textsuperscript{2} statistic was explained by variation between studies of postnatal anxiety exposure). Assessment of relevant moderators to identify sources of heterogeneity was not feasible, because the cohorts used were largely dissimilar in their sample and design characteristics. They could not be grouped and compared in meaningful ways and statistical power would have been insufficient to explore variance explained by higher-order clustering (i.e., MREMs to examine moderation by covariates require meaningful variance between covariates).

Of note, most publications were conducted using an adoption design, meaning that authors could not report estimates that were free from adjustment by genetic confounds (i.e., all adoption results are ‘adjusted by design’ for genetic relatedness, because parents and offspring are not genetically related). Therefore, we were unable to compare effect sizes across levels of adjustment (i.e., adjusted versus unadjusted for genetic confounds).

All presented models used aggregate effects sizes within publications/cohort assuming a median correlation of r=.50, as suggested elsewhere.\textsuperscript{72-74} Results were consistent across the three sensitivity analyses run for each effect size aggregation (r=.10/.50/.90, by publication and by cohort; see Figure S2, available online).

Additional observations

Within the publications examining prenatal anxiety exposure, only the IVF design yielded a significant, non-genetic association, for offspring anxiety in middle childhood (standardised β=.21).\textsuperscript{58} However, this effect was attenuated and no longer significant after postnatal anxiety exposure was controlled for (standardised β=.11). In the two publications using sibling-comparison designs, researchers found no significant associations in any of their reported analyses with offspring during early childhood (standardised β range -.03–.07).\textsuperscript{51,59}

As shown in Table 2, all effect sizes involving postnatal anxiety exposure were weak (standardised β and r values ranged .00–.25). However, structural equation models in two publications using adoption designs showed that parent and child symptoms could prospectively predict one-another across time, highlighting intergenerational, non-genetic, transactional effects during early and middle childhood.\textsuperscript{39,40} Results from these publications showed differences for mother-child versus father-child effects. For example, stronger evidence for
an effect of child symptoms on fathers’ compared to mothers’ anxiety was observed during infancy;\textsuperscript{39} while an
effect of child symptoms during middle childhood was only observed for mothers’, not fathers’ anxiety.\textsuperscript{39} The
only publication using a sibling-comparison design for postnatal analyses showed that mothers’ symptoms did not
prospectively predict offspring internalizing symptoms, after controlling for genetic relatedness.

**Publication Bias**

Studies with significant findings are more likely to be published in scientific journals, which increases risk of
incorrect conclusions from systematic reviews of published literature and risk of false positive or negative findings
in meta-analytic results.\textsuperscript{39} For example, non-significant intergenerational associations are unlikely to be published,
meaning that parents may appear more similar to their offspring when judging by the published literature alone.
Funnel plots for our data are shown in Figure S3 (available online), providing preliminary, albeit non-significant,
evidence for publication bias. Only one publication reported null findings for any association between parent
anxiety and offspring internalizing, however the main focus of that study was on other phenotypes not relevant to
this review, for which they had significant findings.\textsuperscript{61}

**Discussion**

Following a systematic literature search we found only eight publications where authors used a quasi-experimental
research design to control for genetic confounding in associations between parent anxiety exposure and offspring
internalizing outcomes. These used data from four independent cohort studies located in northern Europe or
America, where each cohort had a different quasi-experimental research design applied. Low homogeneity
between publications from different cohorts yielded low statistical power to test for moderation by methodological
(e.g., study design) or observed (e.g., child age) covariates. Results highlight a striking need for new research,
without which we remain ill-equipped to identify the pathways underpinning why parent anxiety symptoms are
associated with the development of offspring internalizing problems.

**Mother’s prenatal anxiety symptoms were not associated with offspring internalizing symptoms after
controlling for genetic relatedness**

Results from three publications, using data from two cohorts, indicated that prenatal exposure to maternal anxiety
is not associated with offspring internalizing symptoms via non-genetic mechanisms. Quasi-experimental research
examining prenatal depression symptoms shows similar findings (also derived from the MoBa cohort evaluated in the present study). As such, quasi-experimental findings to date contradict existing literature on foetal programming in the context of familial risk for internalizing problems, which has been derived mostly from observational and/or animal studies. We emphasise the need for new genetically informed investigations to produce a robust evidence base, looking across child development and into adulthood, including data from more diverse samples. Until new research is available, we encourage researchers and clinicians to consider the importance of genetic transmission and postnatal exposure in their work on maternal anxiety during pregnancy.

**Concurrent associations between parent anxiety and offspring internalizing symptoms remained significant after controlling for genetic relatedness**

A small but significant association was found for concurrent anxiety exposure and child internalizing symptoms, in quasi-experimental studies that accounted for parent-child genetic relatedness. This finding is consistent with a causal interpretation, potentially reflecting at least some direct, environmentally mediated influence between parents and offspring. However, this result is limited to cross-sectional data. It cannot inform on the direction of effects between parents and offspring, nor on the stability of associations across time. Meta-analyses of concurrent versus longitudinal associations were not feasible given the scarcity of available data.

Mixed findings were reported in the few publications that did include longitudinal analyses within their quasi-experimental design. Adoption data showed evidence consistent with parent anxiety predicting child internalizing symptoms, within two year periods during early and middle childhood. The same data also showed evidence for child-to-parent effects, mirroring results from longitudinal studies that do not control for genetic relatedness between parent and child. However, researchers using a sibling-comparison design found that mothers’ postnatal anxiety symptoms did not prospectively predict offspring internalizing symptoms within a five-year-period. It is clear that further research is needed. Although we did not restrict our search by offspring age, we only found publications conducted during childhood. Genetically informed research on familial depression in Sweden suggests maintenance of parent-offspring associations into adulthood. It is unknown whether the same pattern holds for anxiety.

In sum, the data retrieved in our systematic search provide some evidence for non-genetic pathways between parent anxiety and concurrent offspring internalizing symptoms during childhood, however longitudinal research is lacking and so the direction of effects between generations remains unclear. This is an important message for clinicians working with parents experiencing anxiety symptoms: we currently cannot tell with
confidence whether parents’ symptoms exert palpable, lasting influence on offspring internalizing outcomes. Furthermore, research has been limited to very homogenous participant groups and information is lacking as to the generalisability of results across populations.

**Considering the role of methodological confounding**

*Bias by quasi-experimental design*

More research is required before we can test the extent to which effect estimates were biased by each quasi-experimental design used. The largest parent-offspring association that we found was derived from the only publication to examine adolescent offspring – also the only publication to use a children-of-twins design. We cannot tell whether this reflects influence of the developmental period, research design, and/or other factors. In children-of-twins research, the influence of genetic relatedness on a parent-offspring correlation will be underestimated if statistical power is low, thereby inflating the unconfounded residual estimate. As such, statistical power issues could explain the relatively large effect size derived from the only children-of-twins publication. Conversely, the role of genetic relatedness in families can be overestimated in sibling-comparison research, thereby deflating the adjusted estimate. This is because confounding by genetic and environmental family factors are simultaneously corrected for, while assuming that symptoms in the exposed sibling do not influence symptoms in the non-exposed sibling. This could explain the relatively small effect sizes reported in the two sibling-comparison publications we included in meta-analyses.

It is also possible for both children-of-twins and sibling-comparison designs to overcorrect for genetic relatedness if genetic factors comprise an integral part of the causal pathway in parent-to-child environmental transmission, rather than acting as confounders across generations. Further, in both designs it is possible for the effects of evocative gene-environment correlation to inflate estimates of the parent’s causal influence on the child.

The limitations associated with statistically controlling for genetic effects are bypassed in adoption and IVF designs, where parents and offspring are not genetically related. These designs make it easier to distinguish parent-to-child and child-to-parent causal effects. However, they come at the cost of smaller and potentially less-representative samples. For example, both parents of adopted and donor-conceived offspring go to great lengths to have a child, which may lead to differences in parent-offspring relationships compared to families raising naturally-conceived, biological offspring. Further, adoptive parents typically have more advantaged socio-economic backgrounds compared to the birth parents of adopted children and in comparison with non-adoptive parents. Parents in IVF samples may be older and experience higher levels of antenatal risk.
at birth are at higher risk for having experienced prenatal adversity and inheriting genes associated with psychopathology; while the experience of being raised by parents to whom you are not genetically related (as a result of adoption or donor conception) may also influence child development.87,88 As such, conducting new research using a range of quasi-experimental designs should help to balance the strengths and limitations of each, yielding more reliable and robust conclusions.52

Measurement bias

It is likely that measurement bias accounts at least partially for the heterogeneity observed across our reported effect estimates. When working with the large samples required for genetically informative quasi-experiments, it can be methodologically and/or logistically impractical to include lengthy assessments and more than one reporter per family. For example, prenatal symptoms in the IVF study were reported by mothers using a single item, several years after pregnancy, alongside mothers’ reports of offspring internalizing.58 Recall bias and shared method variance may have inflated the parent-offspring correlation in this sample. In the only publication to eliminate risk of shared method variance, parents’ self-reports of anxiety were not associated with child symptoms (measured by researcher observations).61 However, researcher observations of young offspring in artificial situations may not have been as reliable as parents’ reports. Indeed, data from multiple reporters do not always converge. For example, in the adoption cohort we see low agreement between parent reports of offspring anxiety, with father-child anxiety associations only observed when using fathers’ reports for both child and self.39 When new research becomes available, it will be informative to test for moderation by aspects of publication measurement protocol, to investigate influence on pooled results in MREM analyses. In the meantime, it will be important for researchers to consider the perspectives of multiple reporters where possible and maintain clarity as to the potential impact of measurement bias on results.

Use of observed covariates

In each publication, the nature, number and combination of observed covariates will have influenced the strength and meaning of the results. Authors attempted to correct their analyses in a range of ways across publications (e.g., regressing out the effects of age, sex and socio-economic-status, see Tables 1 and 2). Some effect estimates included in our meta-analyses included no correction for measured covariates and were arguably ‘under-corrected’ (e.g., adoption results that did not include correction for perinatal complications). In contrast, analyses in one publication using a sibling-comparison design involved use of several covariates.51 When our meta-analysis of concurrent anxiety exposure was computed without results from the sibling-comparison analyses (which
comprised by far the biggest sample), it was reassuring to find that the pooled effect estimate only increased by $r = .03$ (see Supplement 3, available online). Further, controls for anxiety exposure at different developmental stages requires consideration. In the case of chronic parental anxiety, collinearity becomes an issue for statistically differentiating exposure effects at different periods (e.g., prenatal versus postnatal anxiety effects). That is, if anxiety symptoms before and after the child’s birth are highly correlated, controlling for variance in one period will remove variance in the other. This could explain why the prenatal anxiety association in the IVF study became non-significant after controlling for postnatal anxiety exposure (although analyses of postnatal exposure that included correction for prenatal symptoms did not find the same phenomenon, as residual postnatal symptoms remained predictive of offspring internalizing). Going forwards, we encourage researchers to report both unadjusted and adjusted results, as Bekkhus et al. did, alongside information on the variance explained by each covariate, to help in future research efforts to combine results.

**Further avenues for research**

*Expanding analyses beyond parent-offspring dyads*

The majority of research used in this review is focussed on mother-child dyads. Where possible, it will be informative to take a more holistic approach to intergenerational research, considering fathers, siblings, extended family members and the myriad of social, economic and societal factors that can influence participants’ mental health. For example, modelling both mother-child and father-child associations concurrently across time shows transactional influences between all individuals. Going forward, researchers could also include sibling effects in research and avoid the bias associated with selecting only one child per family for analyses (or two differentially exposed siblings). This could be possible in the Early Growth and Development Study (EGDS), where data are now collected on both birth and adoptive siblings. Information on multiple children per parent is also available in the Norwegian Mother, Father and Child Birth Cohort Study (MoBa), where siblings can be included in multiple-children-of-twins models. These can be used to examine moderation by environments shared within families (e.g., family composition and social support) and between families (e.g., cultural and societal factors), while also including data on two parents, to address issues surrounding assortative mating. The consequences of parents’ resemblance in anxiety has not yet been considered in genetically informed, intergenerational research. In sum, researchers should strive to move beyond analyses of only mother-child dyads, to ensure validity and generalisability of results across families.

*Cohorts that were not designed for quantitative genetic research*
The quasi-experimental designs used in this review require highly specific, large-scale family-samples. That we only identified eight publications, using data from only four cohorts, is telling of the challenges associated with collecting these data. Several publications that we excluded in our systematic search used data from large-scale population studies (e.g., Generation R and the Avon Longitudinal Study of Parents and Offspring) that are rich in phenotypic information but lacking the targeted recruitment required for traditional, pedigree-based genetic research (e.g., adoptive parents or twins with children). Rapidly evolving methods in genomic research may soon provide novel opportunities for these cohorts, using participant DNA to examine intergenerational genetic transmission.

At present, genomic research for complex traits remains limited by a ‘ceiling effect’, whereby results reflect only the additive effects of genetic variants tagged on DNA arrays, excluding non-additive effects or rare variants. Until this is addressed, genomic methods cannot adequately control for genetic relatedness when examining associations between parent and child traits in a way that is comparable with the control achieved in adoption, sibling-comparison or children-of-twins research. When whole-genome methods become possible, analyses can involve use of polygenic scores in parents and offspring, to examine the role of transmitted versus non-transmitted genetic variants in phenotypic associations across generations. The principals of Mendelian randomisation can also be used to examine environmentally-mediated, parent-to-child causal pathways, using parent genes as instrumental variables. Further, genomic variance decomposition methods can be used to partition the influence of parent and offspring genetic influence on traits when genome-wide single nucleotide polymorphism (SNP) data have been collected from family members (e.g., using m-GCTA, Trio-GCTA or Relatedness Disequilibrium Regression). We may soon be able to decompose covariance in traits across generations using estimates of SNP-based heritability. With rapid advances in genomic research, we may be on the brink of a new era for advancing our understanding of familial risk for anxiety and internalizing.

Some limitations of our methodology require emphasis. To pool together all available data, we combined a mix of bivariate and partial correlations. This limited our ability to directly compare estimates between publications, where different adjustments were made for observed covariates. We did not distinguish different types of internalizing problems among offspring, but instead pooled available data relating to child anxiety, negative affect, social inhibition and other emotional difficulties. We cannot tell whether findings would differ by child disorder subtype. This results from lack of available data, meaning we could not test for moderating terms in MREM analyses. All reviewed publications included a quasi-experimental design to account for genetic transmission effects in parent-offspring associations (i.e., controlling for passive gene-environment correlation),
and in our discussion of their findings we consider the possibility of child-to-parent evocative effects. However, we do not consider the possible action of gene-by-environment interaction in families, whereby genetic effects on traits vary in relation to individuals’ contexts or environments, and vice versa. Gene-by-environment interactions are not modelled in any of the publications that we review and thus represent an important avenue for future research in the context of exposure to parental anxiety. Finally, all data used in our meta-analysis were derived from participants located in northern Europe or America. Ancestry data suggested that participants were predominantly of European descent. We highlight the need for new research in more representative samples, in terms of geographical regions and participant ancestry. Without efforts to improve diversity in research participation, we risk preserving a cycle of scientific evidence, and subsequent evidence-based policy, based on groups who are, on average, privileged members of society.

Summary

Quasi-experimental designs can help to control for the effect of genetic relatedness in similarities between parents and offspring. We sought to investigate whether associations between parent anxiety symptoms and offspring internalizing symptoms can be explained via non-genetic mechanisms. We found the existing literature to be limited, with only eight genetically informed studies published, using data from only four cohorts. In a meta-analysis of the available data, we found no evidence to suggest that maternal prenatal anxiety symptoms exert influence on the development of offspring symptoms via non-genetic mechanisms. However, we show that during childhood parent anxiety symptoms are associated with concurrent internalizing symptoms in offspring via non-genetic mechanisms.

References


Brooker RJ, Neiderhiser JM, Kiel EJ, Leve LD, Shaw DS, Reiss D. The Association Between Infants' Attention Control and Social Inhibition is Moderated by Genetic and Environmental Risk for Anxiety. *Infancy*. 2011;16(5):490-507.


Table 1. Studies of Prenatal Anxiety Exposure: Extraction of Quasi-Experimental Data

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Cardiff In Vitro Fertilisation (IVF) study</th>
<th>Norwegian Mother, Father and Child Birth Cohort Study (MoBa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region</td>
<td>United Kingdom</td>
<td>Norway</td>
</tr>
<tr>
<td>Available information on offspring ancestry</td>
<td>Offspring: 91.5% European</td>
<td>Information not available</td>
</tr>
<tr>
<td>Quasi-experimental design</td>
<td>IVF “cross-fostering”</td>
<td>Sibling-comparison</td>
</tr>
<tr>
<td>N individuals per family unit</td>
<td>2 (1 parent, 1 child)</td>
<td>3 (1 parent, 2 child)</td>
</tr>
<tr>
<td>Reference</td>
<td>Rice, 2010 58</td>
<td>Bekkhus, 2018 59</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Gjerde, 2018 51</td>
</tr>
<tr>
<td>N family units in quasi-experimental sample</td>
<td>205</td>
<td>5935</td>
</tr>
<tr>
<td></td>
<td></td>
<td>11553</td>
</tr>
<tr>
<td>Exposure period (gestational weeks)</td>
<td>Prenatal (31-40)</td>
<td>Prenatal (17-30)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Prenatal (30)</td>
</tr>
<tr>
<td>Exposure measure (reporter)</td>
<td>Anxiety/stress: 1 item completed retrospectively, 11-point response scale (self)</td>
<td>Anxiety: 5 and 8 item Hopkins Symptom Checklist, 4-point scale (self)</td>
</tr>
<tr>
<td>----------------------------</td>
<td>---------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------</td>
</tr>
<tr>
<td>Parent relationship to child</td>
<td>Mothers</td>
<td>Mothers</td>
</tr>
<tr>
<td>Outcome period (child age years)</td>
<td>Middle childhood (4-10)</td>
<td>Infancy (0.5, 3)</td>
</tr>
<tr>
<td>Outcome measure (reporter)</td>
<td>Anxiety: 6 items based on DSM-IV, 3-point response scale (mother)</td>
<td>Infant difficulties: 9 item Infant Characteristic Questionnaire, 7-point scale (mother)</td>
</tr>
<tr>
<td>Genetically informative analyses (estimates included in the meta-analysis)</td>
<td>(1) Multiple regression (standardised $\beta=.21$, longitudinal)</td>
<td>(1) Multiple regression (standardised $\beta=.07$, .02, longitudinal)</td>
</tr>
<tr>
<td>Measured covariates considered in the</td>
<td>(1) Child age, child sex, family social occupational class, antenatal complications (vaginal bleeding; admission to hospital for</td>
<td>(1) None</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>estimate extracted for meta-analyses *</td>
<td>high blood pressure/oedema; maternal cigarette smoking; maternal alcohol use; infant plurality)</td>
<td>(2) Child age, child sex, family social occupational class, antenatal complications (vaginal bleeding; admission to hospital for high blood pressure/oedema; maternal cigarette smoking; maternal alcohol use; infant plurality), maternal postnatal anxiety/depression</td>
</tr>
<tr>
<td>Direction of effects assessed in the publication</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

Note: DSM = Diagnostic and Statistical Manual of Mental Disorders

a. Numbering indicates separate, genetically informative analyses, where authors did versus did not adjust for postnatal anxiety exposure.

b. Regression coefficient standardised using the reported standard deviations (s) for the independent (x) and dependent (y) variables \[ z\beta = \beta(s_x/s_y) \].
<table>
<thead>
<tr>
<th>Cohort</th>
<th>Early Growth and Development Study (EGDS; Cohort I only)</th>
<th>Twin and Offspring Study in Sweden (TOSS)</th>
<th>Norwegian Mother, Father and Child Birth Cohort Study (MoBa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region</td>
<td>United States of America</td>
<td>Sweden</td>
<td>Norway</td>
</tr>
<tr>
<td>Available information on participant ancestry</td>
<td>Adoptive mothers: 91.4% European; 3.6% African; 2.5% Latino; 2.5% Other/Mixed. Adoptive fathers: 90.2% European; 5.0% African; 1.7% Latino; 3.1% Other/Mixed. Birth Mothers: 71.7% European; 11.4% African; 6.7% Latino; 10.8% Other/Mixed. Birth Fathers: 74.6% European; 8.7% African; 8.7% Latino; 8.0% Other/Mixed</td>
<td>Participants: 100% European a</td>
<td>Information not available</td>
</tr>
<tr>
<td>Quasi-experimental design</td>
<td>Adoption</td>
<td>Children-of-twins</td>
<td>Sibling-comparison</td>
</tr>
<tr>
<td>N individuals per family unit</td>
<td>2 (1 parent, 1 child)</td>
<td>4 (2 parent, 2 child)</td>
<td>3 (1 parent, 2 child)</td>
</tr>
<tr>
<td>N family units in quasi-experimental sample</td>
<td>361</td>
<td>361</td>
<td>349</td>
</tr>
</tbody>
</table>

a European
<table>
<thead>
<tr>
<th>Exposure period (child age years)</th>
<th>Infancy (0.75)</th>
<th>Infancy (0.75)</th>
<th>Infancy (0.75, 1.5, .2.25)</th>
<th>Middle childhood (6, 7, 8)</th>
<th>Adolescence (11-22)</th>
<th>Infancy (0.5, 1.5, 3, 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure measure (reporter)</td>
<td>Anxiety: 21 item Beck Anxiety Inventory, 4-point scale (self)</td>
<td>Anxiety: 21 item Beck Anxiety Inventory, 4-point scale (self)</td>
<td>Anxiety: 21 item Beck Anxiety Inventory, 4-point scale (self)</td>
<td>Anxiety: 20 item State-Trait Anxiety Inventory for Adults, 4-point scale (self)</td>
<td>Anxiety: 20 item Karolinska Scales of Personality, 4-point scale (self)</td>
<td>Anxiety: 8 item Hopkins Symptom Checklist, 4-point scale (self)</td>
</tr>
<tr>
<td>Parent relationship to child</td>
<td>Unspecified</td>
<td>Unspecified</td>
<td>Mothers, Fathers</td>
<td>Mothers, Fathers</td>
<td>Unspecified</td>
<td>Mothers</td>
</tr>
<tr>
<td>Outcome period (child age years)</td>
<td>Infancy (0.75)</td>
<td>Infancy (1.5, 2.25)</td>
<td>Infancy (0.75, 1.5, .2.25)</td>
<td>Middle childhood (6, 7, 8)</td>
<td>Adolescence (11-22)</td>
<td>Infancy (1.5, 3, 5)</td>
</tr>
<tr>
<td>Outcome measure (reporter)</td>
<td>Social inhibition: Observational tasks (researcher)</td>
<td>Internalizing: 36 item Child Behaviour Checklist, 3-point scale (mother, father)</td>
<td>Negative affect composite: 11 item Infant Characteristics Questionnaire, 7-point scale; 36 item Infant Behaviour Questionnaire, 7-point scale; 19 itemToddler Behaviour Assessment</td>
<td>Anxiety: 13 item Child Behaviour Checklist, 3-point scale (mother, father)</td>
<td>Anxiety: 7 items from Child Behaviour Checklist, 3-point scale (mother, father); 7 items from Child Behaviour Checklist, 3-point scale (self)</td>
<td>Internalizing: 13 item Child Behaviour Checklist, 3-point scale (mother)</td>
</tr>
<tr>
<td>Measured covariates considered in the estimate extracted for meta-analyses</td>
<td>Genetically informative analyses (estimates included in the meta-analysis)</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>Bivariate correlation (r=.00, cross-sectional)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>None</td>
<td>Bivariate correlation (r=.23, longitudinal)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>Bivariate correlation (r=.03, .02, .00, .19, .07, .08; cross-sectional)</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>None</td>
<td>Bivariate correlation (r=.16, .15, .20, .24, .11, 10; cross-sectional)</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>None</td>
<td>Children-of-twins structural equation model (standardised β=.25; cross-sectional)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>Multilevel regression (standardised β=.05; cross-sectional)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Questionnaire, 7-point scale (mother, father), Observational tasks (researcher)

Genetically informative analyses (estimates included in the meta-analysis)

- Bivariate correlation (r=.00, cross-sectional)
- Bivariate correlation (r=.23, longitudinal)
- Bivariate correlation (r=.03, .02, .00, .19, .07, .08; cross-sectional)
- Bivariate correlation (r=.16, .15, .20, .24, .11, 10; cross-sectional)
- Children-of-twins structural equation model (standardised β=.25; cross-sectional)
- Multilevel regression (standardised β=.05; cross-sectional)

Measured covariates considered in the estimate extracted for meta-analyses

- None
- None
- None
- None
- Parent age, parent sex
- Child age, child sex, parity, maternal depressive symptoms at each assessment (including prenatal), exposure and outcome at each
<table>
<thead>
<tr>
<th>Direction of effects assessed in the publication</th>
<th>No</th>
<th>No</th>
<th>No</th>
<th>No</th>
<th>No</th>
<th>No</th>
<th>No</th>
</tr>
</thead>
</table>

**Note:**

a. The data to support this statistic were ambiguous. In their manuscript overviewing this cohort, Neiderhiser and Lichtenstein (2008) report that participants were “in principle 100% Caucasian … consistent with the population of Sweden”.

b. Regression coefficient standardised using the reported standard deviations (s) for the independent (x) and dependent (y) variables \[ z\beta = \beta(s_x/s_y) \].
Figure titles and captions (NB the authors are copyright holders for all figures included)

Figure 1. Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) Flowchart

- Records identified through database searching (k=752)
- Records after duplicates removed (k=429)
- Additional records identified through other sources (k=12)
- Titles/abstracts screened (k=441)
- Full text articles assessed for eligibility (k=55)
- Articles included in qualitative and quantitative synthesis (k=8)

Records excluded (k=386), with reasons used to filter records:
- a. No published empirical research (k=169)
- b. Not general population sample (k=141)
- c. No parent anxiety and/or offspring internalising measure (k=32)
- d. No genetically sensitive research design (k=44)

Full text articles excluded (k=47), with reasons used to filter records:
- a. No published empirical research (k=3)
- b. No parent anxiety and/or offspring internalising measure, or association between these not tested (k=12)
- c. No genetically sensitive research design (k=32)
Figure 2. The Association Between Prenatal Anxiety Exposure and Offspring Internalizing Outcomes

A. Estimates pooled by publication, with multilevel clustering by cohort

<table>
<thead>
<tr>
<th>Publication / Cohort</th>
<th>Estimate [95% CI]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bakkhus2018 / MoBa</td>
<td>-0.02 [-0.04, 0.01]</td>
</tr>
<tr>
<td>Gjerde2018 / MoBa</td>
<td>0.01 [-0.01, 0.03]</td>
</tr>
<tr>
<td>Rice2010 / Cardiff-IVF</td>
<td>0.11 [-0.02, 0.25]</td>
</tr>
</tbody>
</table>

Multilevel Random Effects Model: 0.04 [-0.07, 0.14]

B. Estimates pooled by cohort

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Estimate [95% CI]</th>
</tr>
</thead>
<tbody>
<tr>
<td>MoBa</td>
<td>-0.01 [-0.03, 0.01]</td>
</tr>
<tr>
<td>Cardiff-IVF</td>
<td>0.11 [-0.02, 0.25]</td>
</tr>
</tbody>
</table>

Random Effects Model: 0.03 [-0.08, 0.15]

Note: CI = Confidence Intervals; Cardiff-IVF = Cardiff In Vitro Fertilisation study; MoBa = Norwegian Mother, Father and Child Cohort Study.
Figure 3. The Association Between Postnatal Anxiety Exposure and Offspring Internalizing Outcomes

A. Estimates pooled by publication, with multilevel clustering by cohort

<table>
<thead>
<tr>
<th>Publication / Cohort</th>
<th>Estimate [95% CI]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brooker2011 / EGDS</td>
<td>0.00 [-0.10, 0.10]</td>
</tr>
<tr>
<td>Gjerde2018 / MoBa</td>
<td>0.05 [ 0.04, 0.07]</td>
</tr>
<tr>
<td>Brooker2015 / EGDS</td>
<td>0.07 [-0.01, 0.14]</td>
</tr>
<tr>
<td>Ahmadzadeh2019 / EGDS</td>
<td>0.16 [ 0.06, 0.26]</td>
</tr>
<tr>
<td>Brooker2014 / EGDS</td>
<td>0.23 [ 0.13, 0.33]</td>
</tr>
<tr>
<td>Eley2015 / TOSS</td>
<td>0.25 [ 0.19, 0.31]</td>
</tr>
</tbody>
</table>

Multilevel Random Effects Model: 0.13 [ 0.04, 0.21]

---

B. Estimates pooled by cohort

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Estimate [95% CI]</th>
</tr>
</thead>
<tbody>
<tr>
<td>MoBa</td>
<td>0.05 [ 0.04, 0.07]</td>
</tr>
<tr>
<td>EGDS</td>
<td>0.11 [ 0.03, 0.20]</td>
</tr>
<tr>
<td>TOSS</td>
<td>0.25 [ 0.19, 0.31]</td>
</tr>
</tbody>
</table>

Random Effects Model: 0.14 [ 0.02, 0.25]

Note: CI = Confidence Intervals; EGDS = Early Growth and Development Study; MoBa = Norwegian Mother, Father and Child Cohort Study; TOSS = Twin Offspring Study in Sweden.