Misconceptions about gene–environment interactions in psychiatry

Stanley Zammit,1,2 Michael J. Owen,1 Glyn Lewis2

Misunderstandings surrounding the study of gene–environment interactions are very common. Given the large increase in the number of studies examining interactions in recent years, this raises serious concerns about the value of time and resources spent in these endeavours. In this article we discuss why, despite frequent claims to the contrary, studies of gene–environment interactions are very unlikely to enhance our understanding of disease aetiology or prevention.

PSYCHIATRIC DISORDERS ARE OF MULTIFACTORIAL COMPLEX AETIOLOGY

For multifactorial complex disorders, risk factors are neither necessary nor sufficient to cause disease. In other words, for any given case of disease, multiple risk factors will have acted to bring about the onset of disease in that person. Many different combinations of risk factors will exist for a disease in a population, as described in the sufficient-component-cause model of disease.1 Some of these risk factors are likely to be genetic, and some non-genetic, and both types almost certainly play a contributory role in causing most, if not all, cases of disease. This is sometimes referred to as gene–environment interaction (definition 1) and few scientists would dispute that this is a real phenomenon.

However, this is not the same definition of gene–environment interaction as that reported by studies following analysis of data, which always refers to a measure of statistical interaction. Here data are compared to predictions from statistical models in order to study how exposure to two risk factors in combination affects disease risk. Statistical interaction occurs when the risk of disease if exposed to both the gene (A) and the environmental exposure (B) is different from that predicted by the statistical model being used (definition 2; henceforth referred to as G×E).

However, these predictions can be modelled on either additive or multiplicative scales, and it is important to appreciate that interactions under these models mean very different things (see Appendix 1). For example, statistical interaction between A and B under an additive model (eg, when studying risk differences) means that the risk of disease if both A and B are present is not equal to the additive effect (sum) of the risk if exposed to A only and that if exposed to B only (ie, risk from joint exposure could be greater or less than this). Similarly, statistical interaction between A and B under a multiplicative model (eg, studying risk ratios using logistic regression) means that the risk of disease if both A and B are present is not equal to the multiplicative effect (product) of the risk if exposed to A only and that if exposed to B only (ie, risk from joint exposure could be greater or less than this).

This highlights an important, but often overlooked issue. If both A and B are associated with the disease, then assuming a study is adequately powered, one can always find evidence of statistical interaction by testing both additive and multiplicative models (eg, looking at both risk differences and risk ratios within the same data). In this case, simply providing statistical evidence of G×E has no intrinsic value. Furthermore, hypothesizing that an environmental exposure will have a ‘greater effect’ in the presence of a risk allele compared to absence of that allele is also ambiguous unless the statistical model is specified. For example, the hypothetical data in table 1 show that, under an additive model, the effect of E on risk of disease is greater when G is also present compared to G being absent. However, under a multiplicative model, the same data show that the effect of E on risk of disease is smaller when G is also present. Both models show evidence of G×E, but with opposite conclusions.

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as causal components in any one causal model of disease. Such relationships seem rather unlikely for multifactorial complex diseases, though perhaps are more plausible for studies of adverse effects in pharmacogenetic studies.

As the patterns of interaction become less extreme, gains in power from studying interaction over that of studying main effects decreases, and under most scenarios, simulations indicate that gains from studying G×E over studying main effects only are likely to be minimal, if any. Even where gains in power can occur from studying G×E, this has to be offset against the potential problem of multiple testing and Type I errors. A major problem in the field of genetics has been that of multiple testing, particularly now with the availability of vast arrays of genotypes from genome-wide studies. Examining not only main effects of genes but also their interactions (potentially under different statistical and genetic models) with a multitude of environmental exposures, or on a number of different outcomes, means that spurious results are likely to be extremely common. This problem of multiple testing is further exacerbated by studies reporting interactions in sub-group analyses, or after testing 3-way, 4-way, or more complex interactions.

The reason for this is that the relationships between risk factors for multifactorial complex diseases are always likely to be greater than additive at an epidemiological level, as risk factors are neither necessary nor sufficient to bring about disease. We would rarely expect to observe additivity between risk factors as it is very unlikely that risk factors never co-participate in any causal models of disease. Indeed, empirical evidence shows that where combined effects of established risk factors on disease have been examined, findings support greater than additive patterns of joint risk that is also reflected by findings in other fields of medicine as well as psychiatry. Indeed, the programme of interventions aimed at reducing cardiovascular disease by specific targeting of high-risk groups is based on the assumption of multiplicative (greater than additive) models of combined effects on risk. Where very strong interactions occur (eg, where joint exposure effect is much greater than multiplicative), the case for selective interventions may be strengthened, whereas qualitative interactions clearly have the most important implications for targeting high-risk groups. For less extreme patterns of interaction, however, implications of G×E for informing targeted interventions and prevention of disease are likely to be very limited.

Table 1

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<thead>
<tr>
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<th>E–</th>
<th>E+</th>
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<tbody>
<tr>
<td>G–</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>G+</td>
<td>2</td>
<td>8</td>
</tr>
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</table>

Additive model: Risk difference for E is 4 where G is absent (5–1), and is 6 where G is present (8–2); the effect of E on disease is greater when G is also present

Multiplicative model: Risk ratio for E is 5 where G is absent (5/1), and is 8 where G is present (8/2); the effect of E on disease is smaller when G is also present.

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QUESTION 2: CAN WE RELIABLY DESCRIBE PATTERNS OF RISK FOR JOINT EXPOSURE (REPLICATION)?

Nowhere are the misunderstandings surrounding the study of G×E more apparent than in the literature claiming replication of reported interactions. Interpreting replication of interactions is much more complex than that for main effects.
An essential requirement for claiming replication of G×E is that the pattern of disease risk where there is joint exposure to both risk factors compared to where there is exposure to only one or no risk factors is the same as in the original study. Unfortunately, this is frequently ignored, and unjustified claims of replication are often made simply on the basis of finding evidence of G×E.

Problems of multiple testing in studies of G×E are further compounded by a temptation for authors to conduct subgroup analyses, or to test 3-way or 4-way interactions in what often appear to be increasingly desperate efforts to find evidence of G×E and thereby (incorrectly) claim to replicate an original finding. We illustrate the problems associated with replication of reported G×E by using three high-profile examples.

5-HTT x SLE on depression

An interaction reported between SLEs and variation within a polymorphism in the serotonin transporter gene (5-HTTLPR) on risk of depression received substantial media attention and has undoubtedly been influential in the large increase in studies examining G×E over recent years. In the original report, presence of SLEs was associated with increased risk of depression in subjects homozygous for the s-allele (OR for depression per SLE = 1.68, 95% CI 1.25 to 2.30), with a weaker effect in heterozygotes (OR = 1.48, 95% CI 1.24 to 1.76), and especially l-allele homozygotes (OR = 1.17, 95% CI 0.91 to 1.51). Risk of depression when exposed to both SLEs and 5HTT s-allele was greater than multiplicative (and hence also greater than additive).

Of 34 studies that have examined this relationship since then (see online supplement), 17 have examined whether this relationship is indeed greater than multiplicative. Of these, only 2 find a pattern of interaction consistent with that in the originally study (1 of these in a sub-group only), 3 find evidence of interaction but with patterns inconsistent with that originally reported (4 of these in the opposite direction) and 7 fail to find evidence of any interaction (5 of these with sample sizes substantially larger than the original). Of 15 studies that examined additive models, 6 reported interactions with patterns consistent with the original study (2 in sub-groups only), 3 found patterns that were inconsistent with the original (1 opposite) and 6 failed to find evidence of interaction, though sample sizes were on average, 3-times smaller than the original study. Two of these 6 studies reported G×E in subsequent subgroup analyses.

MAOA x maltreatment on antisocial or violent behaviour

An interaction between variation in the monoamine oxidase A gene (MAOA) and childhood maltreatment on antisocial and violent behaviour in males was reported, such that the effect of maltreatment was greater with the low-activity variants of MAOA compared to those with high activity. Again, this relationship was greater than multiplicative.

Of 11 studies that have since examined this in males (see online supplement), only 4 have examined whether this relationship was indeed greater than multiplicative. Of these, 3 found no evidence for such a relationship, while 1 found support support only when reporting a 1-tailed p value. Of 10 studies that examined additive models, 5 reported evidence of a greater than additive effect (2 of these in subgroup analyses only), 4 found no evidence and 1 found evidence of an interaction but in the opposite direction to that originally reported. In samples that also included females, 1 study examined a multiplicative relationship and found an interaction in the opposite direction. Of 6 studies that examined additive models, 2 found evidence of a greater than additive effect, 3 failed to do so and 1 found an effect in the opposite direction. Studies finding, or failing to find evidence of interaction were equable in size.

COMT x cannabis on psychosis

An interaction between variation in COMT Val158Met and cannabis on psychosis was reported, such that the effect of cannabis was greater in individuals with the valine (high activity) variant compared to those with the methionine allele. Again, this relationship was more than multiplicative, with cannabis use associated with a 10-fold increase in psychosis risk in valine homozygotes, but with no increase in risk in methionine homozygotes. If such an interaction effect size was correct, then we would expect main effects to be observed for both exposures without the need to study interactions given the frequencies of the valine allele and of cannabis use in the population; nevertheless, even in large meta-analyses the association between COMT Val158Met and schizophrenia remains uncertain, despite more than adequately sized samples to identify a main effect of COMT if an interaction with cannabis existed as described above. Three studies that have subsequently examined this relationship (see online supplement) found no evidence to support a relationship as originally reported. In one of these studies, an interaction was reported but only from a subgroup (and 3-way interaction) analysis.

Overall, as can be seen, there is little consistent evidence to support the patterns of disease risk described by any of the original three studies discussed previously. A recent meta-analysis, for example, failed to support a greater than multiplicative relationship between 5-HTT and SLEs on risk of depression. Furthermore, there was no evidence to support a main effect of 5-HTT in this or another recent meta-analysis, though both had adequate power to detect one if the G×E exists as originally reported.

It is possible to introduce bias in meta-analyses by only including studies from which data can be extracted, while debate about inclusion criteria for meta-analyses, as well as about potential reasons for non-replication, often centre on the use of different measures or different sample characteristics. Measures that are less likely than others to accurately reflect an environmental exposure or outcome may reduce the power to observe main effects and interactions, although we wouldn’t expect the pattern of interaction to be substantially different. Similarly, sample characteristics such as age might result in weaker evidence of effects (eg, if time between E and disease increases with age), but is unlikely to substantially alter the pattern of risk for joint exposure to E and G.

Reviews and meta-analyses of putative G×E can be informative if done adequately. However, it is a weak argument to conclude that there is evidence of replication of a specific relationship between two risk factors based merely on the number of studies showing G×E, without also considering the patterns of interaction reported.

SUPPORTING EVIDENCE

Finally, it has been argued, quite reasonably, that in assessing the putative G×Es reported, other lines of evidence should be taken into account. However, conclusions from reports of G×E in, for example, animal models, are equally limited. For example, evidence of greater than additive relationships between 5-HTT and stressful events does not tell us anything about the underlying biology of depression in animals other than both stress and 5-HTT play contributory roles in aetiology.
of the statistical models used and patterns of interaction reported are equally important, and literature in this field also requires careful scrutiny. For example, animal studies cited as supporting findings in humans may be based on selective reporting of subgroup analyses or show patterns of interaction not consistent with those expected21, whilst a more recent, large study of 5-HTT x stress shows a pattern of interaction opposite22 to that postulated in humans.

**REFERENCES**


**SUMMARY**

Genes and environment clearly play contributory causal roles in psychiatric disorders. However, claims about the benefits to be gained from studying G×E are usually made on the basis of misunderstandings surrounding the concepts of interaction. Although possibly greater for studies of pharmacokinetics and more direct studies of cell or system biology, the practical implications from studying G×E in epidemiological studies are likely to be minimal. A number of highly publicised interactions have been reported, but claims of replication have been largely unwarranted. Even if replicated though, the value of finding such interactions is very limited, and the current focus on studying G×E runs the risk of diverting resources from areas of research that can make more of a difference to our understanding of aetiological mechanisms and prevention of disease.

**Appendix 1**

<table>
<thead>
<tr>
<th>Statistical model</th>
<th>Relationship</th>
<th>Definition</th>
<th>Interactiona</th>
<th>Comment</th>
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<tbody>
<tr>
<td>Additive</td>
<td>Additive</td>
<td>Risk (A and B) = Risk (A only) + Risk (B only) – Risk (Neither A nor B)</td>
<td>No (Null hypothesis)</td>
<td>This is also less than multiplicative</td>
</tr>
<tr>
<td>Greater than additive</td>
<td>Risk (A and B) &gt; Risk (A only) + Risk (B only) – Risk (Neither A nor B)</td>
<td>Yes</td>
<td>This could be greater than multiplicative, multiplicative, or less than multiplicative</td>
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<tr>
<td>Less than additive</td>
<td>Risk ratio (A and B) &lt; Risk ratio (A only) x Risk ratio (B only)</td>
<td>Yes</td>
<td>This is also less than multiplicative</td>
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<tr>
<td>Multiplicative</td>
<td>Multiplicative</td>
<td>Risk ratio (A and B) = Risk ratio (A only) x Risk ratio (B only)</td>
<td>No (Null hypothesis)</td>
<td>This is also greater than additive</td>
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<tr>
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*aAssuming adequate statistical power.*