The role of the monoamine oxidase A gene in moderating the response to adversity and associated antisocial behavior: a review

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Abstract: Hereditary factors are increasingly attracting the interest of behavioral scientists and practitioners. Our aim in the present article is to introduce some state-of-the-art topics in behavioral genetics, as well as selected findings in the field, in order to illustrate how genetic makeup can modulate the impact of environmental factors. We focus on the most-studied polymorphism to date for antisocial responses to adversity: the monoamine oxidase A gene. Advances, caveats, and promises of current research are reviewed. We also discuss implications for the use of genetic information in applied settings.

Keywords: behavioral genetics, antisocial behaviors, monoamine oxidase A

Introduction

Our aim in the present paper is to provide an update of studies on the monoamine oxidase A (MAOA) gene as a moderator of adversity-induced antisocial behavior (ASB). After a brief introduction of some key behavioral and molecular genetic concepts, we move on to review selected evidence on the MAOA–adversity–ASB triad; unlike previous reviews that focused only on correlational approaches, we also include experimental work that has shed light on the specific mechanisms underlying the aforementioned relationships. Moreover, we also discuss current and future issues in the applied use of genotyping.

Genes moderate behavior

Can genes moderate our daily behavioral responses? If so, how do genes moderate them? The moderation of behavior by genes, namely genetic sensitivity to environmental stimuli, is known as gene–environment (G × E) interaction. More specifically, G × E interactions are said to occur when the effects of the environment on individuals vary by their genotype, or conversely, when environments modulate genetic effects.¹ Behavioral G × E interaction studies thus operate according to the notion that individuals may have distinct genetic sensitivity to environmental conditions in the determination of a given outcome. Under this apparently simple reasoning, colliding views have been proposed regarding how G × E interactions take place.² Does genetic makeup confer modified sensitivity to environmental stressors, or does it provide differential responsiveness to both favoring and detrimental environments? The former perspective corresponds to the diathesis–stress model, wherein genes are thought to determine individuals’ vulnerability to environmental risk factors. The latter view matches the differential susceptibility model, which assumes that
constitutional features shape general individual plasticity to environmental influences.\(^2,3\)

While most G × E studies assume implicitly or explicitly the diathesis–stress model,\(^2,3\) some research has provided disagreeing evidence. For instance, van Ijzendoorn et al\(^4\) found support for the differential susceptibility model on the serotonin transporter gene’s promoter region polymorphism (5HTTLPR) with a meta-analytic approach. Caucasian participants carrying the short allele – the risk variant – of the aforementioned polymorphism were more susceptible than individuals with the long allele to both positive environments \((r=0.21\) versus \(r=0.11\)) and negative environments \((r=0.22\) versus \(r=0.06\)) for different behavioral, psychiatric, and developmental measures. These results support the assertion that certain assumedly risk-related genetic variants may help determining the degree to which individuals can benefit from favorable environmental conditions.\(^5\) In fact, carriers of risk alleles often have the lowest outcome scores in absence of environmental stressors and the highest scores when those stressors are present.\(^2\) As commented later on in this paper, the differential susceptibility hypothesis deserves further investigation.

Despite the appeal of the approaches briefly outlined above, it must be noted that they conceive of individuals as passive recipients of environmental influences, which are considered as external and independent from the person. In this sense, a line of research led by Robert Plomin caused an upheaval in our understanding of behavioral genetics when he treated the environment as a dependent variable, only to find that environmental measures could be partially determined by genetic variation.\(^6\)

This led to the notion of gene–environment correlations \((r_{G \times E})\), which emerge when genetic and environmental factors covary synergistically.\(^7\) Active \(r_{G \times E}\) occurs if individuals’ genes drive the environments they select (niche-picking). Passive \(r_{G \times E}\) is said to happen when individuals receive genetic and environmental characteristics that reinforce mutually. Evocative \(r_{G \times E}\) takes place when individuals elicit responses in the environment in a way that matches their own inherited qualities. School achievement constitutes an excellent example to illustrate how \(r_{G \times E}\) can be relevant. Youngsters with greater intellectual ability are more likely to attend class (active \(r_{G \times E}\)), where they may be stimulated by an academic environment (passive \(r_{G \times E}\)) in which they evoke intellect-enhancing behaviors from their teachers (eg, pedagogical attention) and peers (eg, amusing discussions or debates). Given the ubiquitous influence of \(r_{G \times E}\), it has become customary to control for \(r_{G \times E}\) in interactional approaches,\(^8-10\) although this is not always accomplished properly.

Aiming to reunite the existing perspectives on G × E relationships, a new framework emerged under the name of gene–environment transactions,\(^6\) which paralleled the development of the person–environment transactions models in personality and social psychology.\(^11\) The neutrality of the term “transactions” is on purpose, to include both interactional and correlational approaches to G × E relationships. This seems to be the next direction in behavioral genetics to fully account for the boundaries, direction, and magnitude of gene-to-environment and environment-to-gene effects. In this section, we have reviewed how genes and environment relate to each other, either interacting or correlating in a number of ways.

### Specific genes involved in behavioral responses

In the previous section, we reviewed theoretical and empirical relationships between genes, environment, and behavior. But, how does this translate into specific findings? Some polymorphisms have been associated to psychological outcomes, ranging from basic processes (eg, working memory capacity,\(^12\) emotion perception,\(^13\)) to complex social behaviors (eg, altruism\(^14\)): as time goes by, the number of such polymorphisms increases. In this sense, some genes have received more attention than others, depending on the existing preliminary evidence (eg, animal models) and theory on the role of certain genes (eg, putative implication in response to pathogen) when selecting candidate polymorphisms.\(^1\)

The most extensively studied G × E relationships to date are 5HTTLPR and stressful life events in depression,\(^15\) and the \(MAOA\) gene and childhood maltreatment in antisocial behavior.\(^16\) Genes related to dopaminergic activity, such as the \(catechol-O-methyl\) transferase (\(COMT\)) gene,\(^17\) the dopamine transporter gene,\(^18\) or polymorphisms encoding for distinct dopamine receptors (eg, \(DRD4\)),\(^19\) have also been widely examined. Whereas infantile conduct disorders\(^19\) and addictions\(^20\) are prominent outcomes in such studies, perhaps the most renowned G × E association involving dopamine systems is the effect of adolescent cannabis consumption in psychosis, varying upon \(COMTVal158/108\)Met genotype.\(^21\)

One cautionary note must be stated here. Most of the research presented so far is based on association studies\(^1\) involving a single gene in conjunction with environmental measures; therefore, findings thus far have to be confirmed either via meta-analyses or complementary approaches that
include causal exploration and more-complex models, such as gene–gene interactions, neuroimaging genetics, or experimental G × E research.

Hence, some genes can interact with other genes at different loci in a process called epistasis, or gene–gene interaction. The extent to which epistatic phenomena are relevant is still unclear, but their underlying logic is straightforward: because most complex traits have polygenic influences, might one gene buffer or enhance the effect of another? Epistatic interactions have been described in mood and behavior disorders, heralding a thrilling area of inquiry.

Once associations have been established between genes, environment, and behavior, a more-complete understanding of these relationships can be achieved by measuring genes’ functional and biochemical correlates at a lower level of abstraction than the behavioral phenotype of interest. The rationale behind this strategy is that outcomes intermediate to behavior should be influenced by a lesser number of genes with more-distinguishable effects. For example, the COMT gene should have stronger effects on speed of performance than on general intelligence scores; in turn, concentration of the COMT enzyme in the frontal cortex should depend more on the COMT gene than on speed of performance.

Functional neuroimaging and other in vivo techniques can lead to the characterization of these intermediate phenotypes, which Gottesman and Gould labeled “endophenotypes.” They restricted the concept to heritable, state-independent, outcome-related, and co-segregated features. In the case mentioned above, one could test frontal levels of COMT as an endophenotype for intelligence. Although the concept of endophenotype is not universally accepted, confirming previously hypothesized – or unexpected – biochemical, neuropsychological, or cognitive pathways for gene action gives G × E relationship findings a sounder ground.

In addition, we can experimentally study the moderation of behavioral responses by the genotype in order to confirm the causal role of genes or the potential roles of expected moderators and to consequently bring these phenomena into the lab. In this sense, van Ijzendoorn and Bakermans-Kranenburg review different alternatives to study G × E interactions experimentally and they provide some specific examples, from the serotonin transporter gene (5-HTT) to the DRD4 gene. Although they do not mention any study involving the MAOA gene, they do provide some examples on substance abuse, which is closely related to ASB.

In the following, we will attempt to provide an update of studies on the MAOA gene as a moderator of adversity-induced ASB. Unlike previous reviews that focused only on correlational approaches, we also include experimental work that has shed light on the specific causal pathways through which genetic effects are exerted and discuss applied genotyping in behavioral settings.

The MAOA gene and its role in antisocial behavior

The MAOA gene was one of the first genes linked to antisocial behavior. Back in 1993, Brunner et al detected a rare mutation of the MAOA gene in three generations of a Dutch family. This mutation hinders the synthesis of MAOA enzyme, which is responsible for degrading neurotransmitters such as serotonin or dopamine. All males carrying this structural anomaly had borderline intelligence and consistently displayed severe antisocial conduct such as arson and rape attempts, whereas females remained unaffected. (Brunner’s finding must not be confused with another recently identified mutation in the MAOA gene associated with a similar phenotype). Located at the short arm of the X chromosome (position 11.23–11.4), the MAOA gene has ever since been central in genetic studies of antisocial behavior.

Given that the mutation described by Brunner et al was highly uncommon, researchers inquired whether normal variations of the MAOA gene could be relevant for ASB. Sabol et al found a polymorphic area in which a nucleotide sequence can be repeated 2, 3, 3.5, 4, or 5 times – although the authors did not report the 2-repeat variant in their original paper. Differences in sequence repeats are called “variable number of tandem repeats” (VNTR) and are generally associated to differential rates of transcription. In the case of the MAOA gene, the 3.5-repeat and 4-repeat alleles had a markedly higher transcriptional efficiency (ie, led to an increased production of the MAOA enzyme) than did the 3-repeat and 5-repeat alleles. Hence, individuals are usually classified as possessing high-efficiency (3.5-repeat or 4-repeat) or low-efficiency (2-repeat, 3-repeat, or 5-repeat) alleles for this gene. It must be noted that the MAOA gene is located at the X chromosome, and hence males are homozygous (ie, have only one copy of the allele). Females have two copies, and it is unclear how the expression of the MAOA gene is affected by the natural random inactivation of one of the X chromosomes.

Caspi et al provided the first evidence that common variants of the MAOA gene could modulate individuals’ antisocial tendencies in interaction with childhood maltreatment in a longitudinal cohort comprehensively – the Dunedin study. Specifically, severely maltreated participants carrying the low-activity allele of the MAOA gene displayed the
highest scores in disposition toward violence and antisocial personality disorder scores, and demonstrated the greatest proportions of adolescent conduct disorder and convictions due to violent behavior.

Since Caspi et al. published their milestone article, many studies have been conducted attempting to replicate and expand their findings to other domains of ASB. Given that Taylor and Kim-Cohen provided the last meta-analysis on the topic, we sought to review papers published in 2007 through 2013 tapping the MAOA gene–adversity–ASB triad. We conducted searches on PubMed and Web of Science, selecting only empirical (not theoretical) articles conducted on human subjects using at least one environmental measure of adversity. References were hand-searched for possible nonretrieved articles. Figure 1 depicts the search sequence; Table 1 shows the search syntax used for each source.

The studies reviewed are presented in Tables 2 and 3. A majority of the studies included in this review present statistically significant evidence for the interaction between the MAOA gene and environmental adversity measures (31 of 37). Overall, the pattern of results suggests that carriers of the low-activity variant of the MAOA gene are more likely to incur ASB when reared in adverse environments. This pattern holds across disparate outcomes such as delinquency, aggressive dispositions, or attention deficit hyperactivity disorder. Among the several environmental stressors that are labeled as adversity, such as neglect, sexual abuse, or family dysfunction, childhood maltreatment shows the strongest effects in predicting ASB. There is thus remarkable variability across studies regarding measurement, sampling, and analytical procedures. Even if this complicates comparison between studies, it also increases confidence in the robustness of the MAOA-by-adversity interaction for ASB, as it has been found in a considerable range of samples assessed in a number of ways.

Extrajamilial social environment appears to be a powerful moderator of the MAOA–adversity interaction. Material deprivation, neighborhood disadvantage, and adoption of the “street code” all seem to add up to the effect of maltreatment, with low-activity allele carriers being at slightly greater risk for ASB in such rearing environments. Indeed, maltreatment is more likely to occur in an unfavorable broad social environment, which in turn offers fewer resources to overcome its pernicious consequences.

Consistent with the rationale above, G × E effects for convictions are not as strong as for milder indices of ASB. This suggests that the types of ASB that lead to imprisonment may be more strongly influenced by macrosocial variables than solely by individual or familial features. However, failures to confirm this G × E interaction have also been reported. Perhaps the most outstanding among such findings are the results obtained by Haberstick et al., who placed formidable effort into replicating the original study by Caspi et al. and did not find any interactive effect in a large sample.

Other studies have found G × E interactions with discordant results, associating the high-activity variant with greater ASB scores in the presence of early adversity; the reasons for these results are unclear. Even more intriguing, this trend is usual in female samples or subsamples.

Table 1: Search syntax to locate articles conducted on human subjects with at least one environmental measure of adversity

<table>
<thead>
<tr>
<th>PubMed</th>
<th>Web of Science</th>
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<tbody>
<tr>
<td>(((maoa&gt;Title/Abstract)) OR (monoamine oxidase a&gt;Title/Abstract)) AND (antisocial&gt;Title/Abstract) OR agress&gt;Title/Abstract) OR viol&gt;Title/Abstract)) AND (advers&gt;Title/Abstract) OR abus&gt;Title/Abstract) OR maltreat&gt;Title/Abstract)) AND (&quot;2007&quot;[Date – Publication]:&quot;2013&quot;[Date – Publication])</td>
<td>TOPIC: (MAOA OR monoamine oxidase a) AND TOPIC: (antisocial OR agress OR viol) AND TOPIC: (advers OR abus OR maltreat). Timespan:2007-2013. Search language:English.</td>
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</table>
Table 2 Recent studies replicating the relationship between MAOA-uVNTR, adversity, and at least one kind of antisocial outcome in humans

<table>
<thead>
<tr>
<th>Reference</th>
<th>Sample</th>
<th>Gene(s) studied</th>
<th>Environmental variables</th>
<th>Outcome</th>
<th>Selected results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nilsson et al (2007)</td>
<td>66 males</td>
<td>MAOA</td>
<td>Family function, maltreatment</td>
<td>Alcohol-related problem behavior</td>
<td>MAOA interacted with family function and maltreatment, such that carriers of the low-activity allele had the greatest alcohol-related problem behavior scores when maltreated or reared in poor-functioning families. The interaction was only marginally significant in a regression model, but was reaffirmed with nonparametric interaction tests.</td>
</tr>
<tr>
<td>Reif et al (2007)</td>
<td>184 males referred for forensic assessment</td>
<td>MAOA, SHHTTLPR, DAT</td>
<td>Childhood adversity, history of substance abuse</td>
<td>Violent behavior</td>
<td>Significant main effect of MAOA on violent behavior, short allele carriers being 2.3 times more likely than long allele carriers to engage in violence. Interaction between SHHTTLPR and MAOA, such that MAOA-L subjects with at least one copy of the short SHHTTLPR allele had a 0.78 probability of being classified as violent if reared in an adverse environment.</td>
</tr>
<tr>
<td>Sjöberg et al (2007)</td>
<td>119 females</td>
<td>MAOA</td>
<td>Sexual abuse</td>
<td>Stealing, vandalism, violence, total criminality</td>
<td>Sexual abuse and MAOA interacted in predicting total criminality and vandalism, but not violence or stealing. 4R homozygotes displayed the greatest criminality scores.</td>
</tr>
<tr>
<td>Vanyukov et al (2007)</td>
<td>148 males</td>
<td>MAOA</td>
<td>Parenting (separately for mothers and fathers)</td>
<td>Disruptive behavior</td>
<td>Paternal, but not maternal, care interacted with genotype in predicting ADHD. Relationships between parenting and disruptive behavior disorders were observed in 4-repeat allele carriers but not in participants with the 3-repeat allele for all outcomes. The 4-repeat allele was thus associated with greater disruptive behavior scores as a function of parenting.</td>
</tr>
<tr>
<td>Ducci et al (2008)</td>
<td>291 American Indian women</td>
<td>MAOA</td>
<td>Child sexual abuse</td>
<td>Alcoholism, ASPD</td>
<td>Greater presence of the low-activity allele in alcoholics and antisocialalcoholics compared to controls within abused group, but not within nonabused group. Also differences in several MAOA and MAOB SNPs.</td>
</tr>
<tr>
<td>Nilsson et al (2008)</td>
<td>119 females</td>
<td>MAOA</td>
<td>Quality of family relations, physical/sexual abuse</td>
<td>Alcohol-related problem behavior</td>
<td>Quality of family relations interacted with MAOA, while the effect of maltreatment varied depending on which groups were compared and was only marginally significant. Homozygous MAOA-L subjects displayed the greater problem scores if family relations were poor.</td>
</tr>
<tr>
<td>Hart et al (2009)</td>
<td>865 males</td>
<td>MAOA</td>
<td>Neighborhood child saturation, poverty, urbanicity, racial heterogeneity</td>
<td>Self-reported aggression</td>
<td>MAOA interacted with child saturation (ie, number of children &lt;15 years of age living in a neighborhood) in predicting aggression, but not with other social indexes. MAOA-L subjects had the greater increase in aggression scores.</td>
</tr>
<tr>
<td>Kinnally et al (2009)</td>
<td>158 females</td>
<td>MAOA</td>
<td>Parental care and SLEs</td>
<td>Impulsivity, trait aggression, depressive symptoms</td>
<td>Three-way interaction between parental care, SLEs, and MAOA. Favoring rearing practices buffered the detrimental effect of SLEs on impulsivity/aggression in MAOA-L subjects, who had similar mean levels as MAOA-LH carriers and way lower than MAOA-HH participants. There were no differences in genotype when subjects suffered both poor parental care and a severe stressor.</td>
</tr>
<tr>
<td>McDermott et al (2009)</td>
<td>78 males</td>
<td>MAOA</td>
<td>Experimental monetary provocation</td>
<td>Experimental aggression (amount of hot sauce allocated against the provocateur)</td>
<td>MAOA-L subjects displayed more aggression than their counterparts in the high-provocation condition, but not in the low-provocation condition.</td>
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**Table 2 (Continued)**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Sample</th>
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<th>Environmental variables</th>
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</thead>
</table>
| Prom-Wormley et al (2009) | 721 females  | MAOA    | Childhood adversity (not abuse), parental ASPD   | Conduct disorder                              | MAOA had a direct effect and an interactive effect with childhood adversity on CD. MAOA-L/L females had a greater proportion of CD in medium levels of exposure to adversity than did carriers of other genotypes, but were protected against CD at the highest adversity levels. The interaction between MAOA and trauma exposure predicted inattention but not rule-breaking or aggression, and differences in aggression between genotype groups only existed at moderate levels of trauma. The MAOA-L variant was related to greater inattention scores at low to moderate levels of trauma, but high-activity allele carriers scored higher when exposed to extreme adversity.
| Weder et al (2009)   | 114 male children (73 maltreated, 41 controls) | MAOA    | Childhood adversity                              | Aggression, rule-breaking, inattention        | Genotype interacted with sexual assault in predicting ASB and CD, but only a small proportion of participants (3%) had suffered this stressor. Low-activity carriers displayed the greater scores in ASB and CD. Harsh discipline increased alcohol and tobacco abuse, ASPD and CD, but did not interact with genotype. Significant G × E interaction for delinquency with multiple and maternal reports, but not with teacher or self reports. Higher antisocial scores for MAOA-L participants.
| Beach et al (2010)   | 538          | MAOA    | Childhood maltreatment                           | Depressive symptoms and ASPD                 | Females carrying low-activity alleles were more susceptible than their counterparts to SLEs for hyperactivity at 4 years and 7 years of age. SLEs at 1.5 to 2.5 years had a greater effect for MAOA-L boys at age 7. Interaction between MAOA genotype and heavy drinking, and between MAOA and childhood abuse, in predicting violent recidivism. Abuse and alcohol consumption heightened risk for violent behavior in high-activity allele carriers, but not in MAOA-L subjects.
| Edwards et al (2010) | 186 males    | MAOA    | Childhood physical discipline                    | Delinquency, aggression, child externalizing problems | MAOA interacted with prenatal exposure to cigarettes in predicting CD. CD symptoms were higher at greater prenatal smoking exposure in female carriers of the high-activity allele, whereas, for males, the low-activity allele was the risk variant. For hostile attribution bias, a milder MAOA × prenatal smoking exposure was found on females, with the high-activity allele associated with greater anger-face identification.
| Enoch et al (2010)  | 6,129 Children 4 to 7 years old | MAOA    | SLE and family adversity                        | Behavioral disinhibition                      | Main and interactive G × E effect on global delinquency index. MAOA-L maltreated boys and MAOA-H/H maltreated girls exhibited the highest delinquency rates. Significant G × E interaction between MAOA and risk-protective factors for anger and hostility in males. Almost significant G × E interaction for incarceration. The MAOA-L allele was associated with higher anger and hostility in unfavorable environments, and with lower scores in favoring environments, when compared with MAOA-H.
| Tikkanen et al (2011) | 174 male alcoholic offenders | MAOA    | Childhood physical abuse                         | Violent behavior                              |                                                                                                                                                                                                                 |
| Wakschlag et al (2010) | 176          | MAOA    | Prenatal exposure to cigarettes, harsh parenting, maternal antisocial behavior | CD, hostile attribution bias (judging nonangry faces as angry) |                                                                                                                                                                                                                 |
| Asklund et al (2011) | 1,825        | MAOA    | Childhood maltreatment                           | Delinquency, vandalism, stealing, violence Incarceration, anger, hostility |                                                                                                                                                                                                                 |
| Beaver et al (2011)  | 493          | MAOA    | Compound measure of protective (eg, social support, religiosity) and risk (eg, neighborhood disadvantage, maternal negativity) factors | bloginfo:www.dovepress.com/dovepress/190/190.html |                                                                                                                                                                                                                 |
Fergusson et al\textsuperscript{10} (2011) & 398 males & MAOA, TPH1, SERT & Childhood abuse (sexual and physical) & Property and violent offending, convictions, conduct problems, hostility & Significant G \times E interactions for most outcomes, especially for self-reported property offending, but not for convictions. When taken separately, sexual abuse had a more significant effect on all outcomes than physical abuse. MAOA-L participants had markedly higher ASB scores than MAOA-H carriers if exposed to early abuse. &

Lee et al\textsuperscript{11} (2011) & 672 males & MAOA & Deviant peer affiliation & Overt and covert ASB & Interaction between deviant peer affiliation and MAOA for overt but not for covert ASB. The high-activity allele was associated with greater problematic alcohol consumption and overt ASB scores. &

Nilsson et al\textsuperscript{12} (2011) & 1,586 & MAOA & Quality of family relations, sexual abuse & Alcohol use disorder & MAOA genotype interacted with sexual abuse in predicting alcohol use disorder (controlling for socioeconomic factors), but not with family relations. MAOA-L males had the highest mean levels of alcohol abuse, whereas MAOA-H/H was the risk allele for females. &

Pardini et al\textsuperscript{13} (2011) & 197 males & MAOA & Brain injury (prefrontal versus non-prefrontal versus controls) & Trait aggression & Genotype \times group interaction, such that, among non–prefrontal-lesion patients, high activity carriers had greater aggression scores than low activity carriers, with the opposite trend in healthy controls and no differences in genotype in prefrontal-lesion subjects. Positive correlations between aggression, early traumatic experiences, and posttraumatic stress disorder symptoms within healthy and non–prefrontal-lesion participants, but not in prefrontal-damaged subjects. &

Cicchetti et al\textsuperscript{14} (2012) & 672 low income children (10–12 years of age) & MAOA, TPH1, SHTTLPR & Childhood maltreatment & ASB & Maltreated MAOA-L children had higher lifetime ASB scores, as well as during the past 6 months. These G \times E interactions were found only for self-reported ASB, but not for peer or adult ratings. &

Fergusson et al\textsuperscript{15} (2012) & 399 males & MAOA & IQ at age 8–9 years, childhood maltreatment, family material deprivation & Property and violent offending, convictions & G \times E interactions for self-reported violent offending and self-reported property offending, but not as strong for convictions. MAOA-L carriers displayed greater antisocial scores than MAOA-H subjects if exposed to early adversity, with childhood maltreatment outscoring other risk factors. &

McGrath et al\textsuperscript{16} (2012) & 192 females (oversampled for prenatal smoking) & MAOA & Childhood maltreatment & Conduct problems & Maltreatment had a stronger effect on conduct problems for carriers of the high-activity allele than for the low-activity variant, but conduct problems were more related to other antisocial dimensions (eg, impulsiveness, interpersonal aggression) in the latter group. &

Nikulina et al\textsuperscript{17} (2012) & 575 & MAOA & Physical abuse, sexual abuse, neglect, multiple maltreatment & Dysthymia, major depressive disorder, alcohol abuse & The low-activity allele protected females from the effects of physical abuse on dysthymia. The 3 or 4 allele form protected Whites against dysthymia, major depressive disorder, and alcohol abuse symptomatology if sexually abused, while the 4 allele genotype protected non-Whites. &

Simons et al\textsuperscript{18} (2012) & 224 & MAOA, SHTTLPR, DRD4 & Hostile/demoralizing rearing environment (including harsh parenting) & Delinquency, aggression, and adoption of the "street code" & Interaction between cumulative plasticity alleles (ie, number of “risk” alleles) and hostile environment on adoption of the street code and aggression. Mediation analyses revealed that the best-fitting model was one in which the G \times E predicted aggression through adoption of the street code. Number of plasticity alleles (including the MAOA-L variant) predicted antisocial scores as a function of environmental adversity – lowest ASB in favoring environments, highest in detrimental ones. &

Gallardo-Pujol et al\textsuperscript{19} (2013) & 57 males & MAOA & Experimental social exclusion & Experimental aggression (point subtraction aggression paradigm) & Significant G \times E (MAOA \times social exclusion) effect. Excluded MAOA-L participants scored more than twice on experimental aggression than their MAOA-H counterparts.
<table>
<thead>
<tr>
<th>Sample</th>
<th>Outcome</th>
<th>Environmental variables</th>
<th>Genotype(s)</th>
<th>Gene(s)</th>
<th>Reference (year)</th>
</tr>
</thead>
</table>

Notes: Studies included here were published in 2007 or later, when the last meta-analysis on the topic was published by Taylor and Kim-Cohen (2007). Only prospective studies were included. Antisocial outcomes include problematic alcohol consumption. A variant of Taylor et al terms is alphabetical for papers published in the same year. Samples include participants of both sexes unless otherwise specified. Antisocial outcomes include problematic alcohol consumption. Abbreviations: 5HTTLPR, serotonin transporter linked polymorphic region; ADHD, attention-deficit hyperactivity disorder; ASB, antisocial behavior; ASPD, antisocial personality disorder; CD, conduct disorder; CSF, cerebrospinal fluid; DAT, dopamine transporter; DAT1, dopamine transporter; E, environment; G, gene; G×E, gene–environment interaction; MAOA, monoamine oxidase A; MAOA-L, monoamine oxidase A low-activity allele; MAOA-H, monoamine oxidase A high-activity allele; ODD, oppositional defiant disorder; SNP, single nucleotide polymorphism; TPH1, tryptophan hydroxylase 1; uvNTR, upstream variable number of tandem repeats.

The latter results may be reflective of a gene–by–sex interaction, such that the low activity (MAOA-L) variant would be the predominant risk or plasticity variant in males, whereas the high activity (MAOA-H) variant would act as such in females. Unfortunately, the answer will remain subject to speculation until we have a clearer picture of the MAOA gene’s action in females. One of the main sources of general discrepancy across studies is the aforementioned methodological variability; the use of longitudinal versus cross-sectional approaches or the selection of instruments may influence results dramatically, let alone the ubiquitous measurement and sampling error. Furthermore, other artifacts such as population stratification (ie, differences in allelic frequencies between subpopulations) might have also biased some results in multiethnic samples. Such bias could be supported by reports from Lea and Chambers, who stirred controversy claiming that the increased percentage of 3-repeat alleles in a Maori population could represent a positive selection for behavioral aggression. However, their claims are not related to modern violence and are hardly generalizable to other populations, but they could explain a certain degree of population stratification. In fact, different allelic frequencies and variants could certainly alter the association between a gene and a complex genotype, which is the case of the oxytocin receptor gene and the serotonin transporter gene.

Among noninteractive effects of the MAOA gene, the 2 repeat (2R) allele has been linked to enhanced delinquency, as well as to strikingly lower levels of transcriptional activity, in comparison to all other variants. Nonetheless, mono- genic effects tend to yield relatively low effect sizes, and should not be considered separately from the environment in applied work.

ASB, as well as aggression, shows an exaggeratedly sex-specific prevalence. In fact, being a male dramatically increases the risk of displaying ASB, and one of the reasons that has been pointed out as an important determinant of these differences is the homozygosity of males for the MAOA gene. Females’ heterozygosis may help to compensate the detrimental effects of the MAOA-L allele through developmental deactivation of the X chromosome that carries the low-activity allele. There is some evidence suggesting that the MAOA-H allele puts females in greater risk for overt ASB. In any case, females have been shown to predominantly manifest ASB in subtler ways than males (eg, verbal rather than physical aggression). Relational, nonphysical aggression is a field of research that should yield relevant results in the future. Interestingly, the few studies that have
Table 3 Recent studies failing to confirm the relationship between MAOA-uVNTR, adversity, and at least one kind of antisocial outcome in humans

<table>
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<tr>
<th>Reference</th>
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<tr>
<td>Prichard et al(^a) (2008)</td>
<td>1,002 males</td>
<td>MAOA</td>
<td>General childhood adversity (including abuse)</td>
<td>Indicators of ASB</td>
<td>MAOA × adversity interaction only when comparing participants who suffered &lt;2 adversity exposures with those who experienced 2–5 exposures. Carriers of the high activity allele showed the highest ASB scores. The interaction between MAOA and maltreatment almost reached significance for delinquency, but not for aggression or the overarching externalizing scale. Main effect for MAOA in all measures, so that carriers of the high-activity allele obtained higher scores than MAOA-L participants.</td>
</tr>
<tr>
<td>Van der Vegt et al(^c) (2009)</td>
<td>239 adopted boys</td>
<td>MAOA</td>
<td>Childhood maltreatment</td>
<td>Externalizing behavior (including aggressive and delinquent behavior)</td>
<td>No interaction between number of early stressful events and genotype in any brain region. Interaction between genotype and sex in the processing of emotional-valenced faces. MAOA-L males had clearly reduced responses in several parietal, frontal, and occipital brain areas when presented with angry faces, whereas for females, genotype differences existed only in the superior occipital gyrus, and with an opposite pattern. MAOA had a dose-dependent effect on the aggression reactivity facet of depression sensitivity in women, such that MAOA-H/H females scored higher than MAOA-H/L ones, who, in turn, had greater scores than MAOA-L/L. No genotype effects were found on trait or state anger. Childhood adversity did not interact with MAOA genotype for any antisocial outcome. Significant though mild main effect of MAOA gene on convictions for the whole sample, and on disposition toward violence in White males. Neither main nor interactive effects in any other score. The absence of G × E effects was not due to lack of statistical power.</td>
</tr>
<tr>
<td>Williams et al(^d) (2009)</td>
<td>210</td>
<td>MAOA</td>
<td>Early life stress</td>
<td>Electroencephalographic activity in a face-processing task</td>
<td>No interaction between number of early stressful events and genotype in any brain region. Interaction between genotype and sex in the processing of emotional-valenced faces. MAOA-L males had clearly reduced responses in several parietal, frontal, and occipital brain areas when presented with angry faces, whereas for females, genotype differences existed only in the superior occipital gyrus, and with an opposite pattern. MAOA had a dose-dependent effect on the aggression reactivity facet of depression sensitivity in women, such that MAOA-H/H females scored higher than MAOA-H/L ones, who, in turn, had greater scores than MAOA-L/L. No genotype effects were found on trait or state anger. Childhood adversity did not interact with MAOA genotype for any antisocial outcome. Significant though mild main effect of MAOA gene on convictions for the whole sample, and on disposition toward violence in White males. Neither main nor interactive effects in any other score. The absence of G × E effects was not due to lack of statistical power.</td>
</tr>
<tr>
<td>Verhoeven et al(^e) (2012)</td>
<td>432</td>
<td>MAOA</td>
<td>Childhood adversity</td>
<td>Anger, depression sensitivity</td>
<td>MAOA had a dose-dependent effect on the aggression reactivity facet of depression sensitivity in women, such that MAOA-H/H females scored higher than MAOA-H/L ones, who, in turn, had greater scores than MAOA-L/L. No genotype effects were found on trait or state anger. Childhood adversity did not interact with MAOA genotype for any antisocial outcome. Significant though mild main effect of MAOA gene on convictions for the whole sample, and on disposition toward violence in White males. Neither main nor interactive effects in any other score. The absence of G × E effects was not due to lack of statistical power.</td>
</tr>
<tr>
<td>Habersstick et al(^f) (2014)</td>
<td>4,316 males</td>
<td>MAOA</td>
<td>Childhood maltreatment</td>
<td>Adolescent conduct problems, adult antisocial behavior, convictions for violent crimes, disposition toward violence</td>
<td>MAOA had a dose-dependent effect on the aggression reactivity facet of depression sensitivity in women, such that MAOA-H/H females scored higher than MAOA-H/L ones, who, in turn, had greater scores than MAOA-L/L. No genotype effects were found on trait or state anger. Childhood adversity did not interact with MAOA genotype for any antisocial outcome. Significant though mild main effect of MAOA gene on convictions for the whole sample, and on disposition toward violence in White males. Neither main nor interactive effects in any other score. The absence of G × E effects was not due to lack of statistical power.</td>
</tr>
<tr>
<td>Sadeh et al(^g) (2013)</td>
<td>237 males</td>
<td>MAOA, 5HTTLPR</td>
<td>Childhood maltreatment</td>
<td>Psychopathy</td>
<td>Significant although small difference between MAOA-L and MAOA-H subjects in the impulsive–antisocial psychopathy dimension, higher for low-activity carriers. No interaction of the MAOA gene with childhood maltreatment. MAOA × 5HTTLPR interaction was not tested, but 5HTTLPR showed effects on the affective and interpersonal psychopathy dimensions.</td>
</tr>
</tbody>
</table>

Notes: Studies included here were published in 2007 or later, when the last meta-analysis on the topic was published by Taylor and Kim-Cohen\(^{20}\) (2007), except if not listed in such article. Articles are presented chronologically; the order is alphabetical for papers published the same year. Samples include participants of both sexes unless otherwise specified.

Abbreviations: 5HTTLPR, serotonin transporter linked polymorphic region; ASB, antisocial behavior; G × E, gene–environment interaction; MAOA, monoamine oxidase A; MAOA-H, monoamine oxidase A high-activity allele; MAOA-L, monoamine oxidase A low-activity allele; uVNTR, upstream variable number of tandem repeats.

examined the role of MAOA in psychopathy—characterized by instrumental aggressive tendencies—have yielded small or nonexistent effects, thus reinforcing the implication of this gene in reactive, rather than proactive, aggression.\(^{57,66}\)

So far we have presented correlational research in which outcomes are often aggregated or self-reported measures.\(^{1}\) As in the case of other genes, as mentioned previously, some authors have brought this question into the lab and have shown that the MAOA gene also moderates specific, pinpointed behaviors. In this sense, McDermott et al\(^{17}\) showed that after provocation, reactive aggression was higher among carriers of the low-activity allele of the MAOA gene.
than among carriers of the high-activity allele, a recently replicated finding. Social exclusion has been usually put forward as a major source of provocation that leads to personal distress and eventually to aggression. In this sense, Gallardo-Pujol et al experimentally replicated the finding from Caspi et al using an ad hoc procedure in which they combined behavioral genetics and social psychology. More specifically, they found that under conditions of social exclusion, MAOA-L carriers were twice as likely to engage in aggressive behaviors than were MAOA-H carriers. Another important contribution is that they provided guidelines for experimental research on G × E interactions. Interestingly, they suggest that experimental G × E research should follow these steps:

1) confirm that there is evidence of a given G × E interaction from epidemiological genetic research; 2) search for analogs of environmental risk and target behavior; 3) search for independent effects of these analogs upon the dependent variable (behavioral task); 4) check the plausibility of the effect of the environmental analog on the biological systems involved in the task; 5) check the evidence of an association for the candidate gene with similar laboratory tasks; 6) control for any possible confounding variables (by blocking, covariates, etc); 7) test for G × E interaction; and finally 8) perform independent replication and meta-analysis.

Once we have correlational and experimental evidence on the interaction of environmental adversity and the MAOA gene, what is the next step? What do we know about the causal mechanisms from the gene to behavior and its interaction with environmental stimuli? Some clues may lie in the neural correlates of the MAOA gene, as well as in brain areas responsive to adversity. If the cerebral circuitry associated to genetic variation couples with that observed in response to adversity, a clearer picture of the MAOA–adversity relationship may emerge. As commented next, this approach has yielded interesting results.

Eisenberger et al found that carriers of the low-activity MAOA allele displayed greater activity in the anterior cingulate cortex as a response to experimentally induced social exclusion, as compared to MAOA-H and MAOA-H/L participants. This evidence suggested that carriers of the MAOA-L variant would experience greater distress when confronted with adverse conditions.

Alia-Klein et al confirmed the functional relevance of the MAOA enzyme by showing moderate negative correlations of brain MAOA activity with trait aggression. However, no differences in enzymatic activity were found regarding genotype, in line with other research failing to find such relationship. These puzzling findings suggest that MAOA genotype does not determine basal enzymatic action, in apparent contradiction with differences detected in structural and functional measures. Speculations converge in pointing out that MAOA genotype may be especially crucial during early stages of development, such that the effects of genotype would only be relevant upon the action of a developmental or environmental disruption of this enzyme’s activity. In line with this formulation, Huizinga et al found no G × E interaction when victimization was limited to adolescence.

Another study reported that MAOA-L males displayed structural reductions in emotional processing areas (ie, bilateral amygdala and cingulate cortex) but increased functional activity in these same areas when evaluating angry versus fearful faces.

Drawing upon this work, Buckholtz and Meyer-Lindenberg posited that the low-activity allele of the MAOA gene conferred a more labile sociocognitive processing system, characterized by an increased tendency to respond hostilely to aggression cues – even if ambiguous. Biochemically speaking, these differences may translate into an excess of amygdalar reactivity that demands greater frontal regulatory action in MAOA-L subjects; supporting this view, functional connectivity between ventromedial prefrontal cortex and amygdala has been associated with high levels of harm avoidance and angry hostility, as well as low reward dependence, in MAOA-L (but not MAOA-H) males. By default, this pattern would only be manifest in the shape of temperamental variations within the normal range. Nevertheless, early disruptions in the serotonin and epinephrine circuits, such as those caused by maltreatment or other forms of environmental adversity, would render MAOA-L participants more susceptible to reacting aggressively in social interactions. The serotonin deficiency hypothesis, which states that low serotonin levels would be associated with greater aggression, has been demonstrated to be an oversimplification, but it appears clear nonetheless that the MAOA gene influences reactive aggression mainly through serotonergic imbalance. Note that the brain mechanisms mentioned above apply to structurally intact brains; however, a lesion study with war veterans has shown that an intact prefrontal cortex may be a prerequisite to detect genetic effects. See Table 2.

Taken together, these studies shed light about the causal mechanisms by means of which environmental adversity is moderated by the genotype, therefore providing intervention targets if they are needed.
**Personalized genomics in the management of antisocial behavior**

In this section, we will focus on interventions for ASB, since it has been the most important outcome associated to the MAOA gene, and how they could be improved by taking into account existing knowledge. Personalized genomics in this particular case could be fitted into the broader framework of neuroprediction. Neuroprediction is the use of biological data to predict future behavior. This term recently gained prominence in the realm of risk management after the publication of a longitudinal study in which the use of neuroimaging data successfully predicted rearrest. As we will see by the end of this section, taking genomics into account can significantly improve risk management. The incremental validity of including genomics in applied behavioral management remains to be tested, although this practice has been routinely incorporated into cancer protocols. Genetic effects are often small, and therefore it is very unlikely that a single gene would yield a large increment in validity. Nonetheless, the more we know about multiple genes in relation to ASB and their interactions with other risk factors, the more prepared we will be to properly manage them.

Most interventions to prevent or treat ASB are based on single psychosocial risk factors. For instance, training in peer-group skills or mentoring programs are aimed at reducing the influence of deviant peers, a well-established risk factor. However, coming together with delinquent partners may well reflect active rG × E, as youngsters might seek out such environments partially due to their own inherited tendencies. Hence, it is no surprise that success rates obtained by programs of this kind are inconsistent. Further quantitative genetics studies are required to draw clearer delineation between true criminogenic factors and those confounded with genetic influence. An accurate targeting of the factors to be modified could benefit not only society as a whole (eg, avoiding inefficient preventive policies), but also individuals (eg, by helping to adapt treatments for specific patients). Noteworthy, there is a growing interest in targeting specific risk factors as the most efficient way to cure a disorder, or as some authors suggest, “cause should inform cure”,.

Following this line of reasoning, Collins among others, argued that diagnoses based on etiology rather than symptomatology should be more reliable and should allow for better-tailored environmental and behavioral treatments. This constitutes the core idea of what he called “personalized medicine”. Hence, the identification of specific genetic variants probabilistically associated with certain outcomes directly taps into Collins’ prophecy by allowing the assessment of constitutional risk and protective factors. In fact, this is so-called “therapygenetics”, similar to pharmacogenetics.

The only study where the MAOA gene has been tested with therapeutic outcomes found that carriers of the high-activity allele had a worse response than low-activity carriers to cognitive–behavioral treatment for panic disorder with agoraphobia. More specifically, participants with the high-activity variant displayed less improvement than their counterparts in self-reported anxiety, avoidance (only in females who completed the whole study), and clinician-rated global severity. Furthermore, participants’ responses in a behavioral avoidance test (BAT) revealed that subjects bearing the high activity allele had a faster heart rate in all BAT phases, benefited less from repeated exposure, and reported more anxiety in anticipation and throughout the BAT than did low-activity allele carriers. This latter group also showed a neural activity pattern in a fear-conditioning paradigm suggestive of a better discrimination between anxiogenic stimuli. The implications of this study, pending replication, are manifold: carriers of the high-activity MAOA variant may benefit more from treatments focused on autonomic arousal (eg, relaxation, beta-adrenergic blockers) and might have an increased tendency to generalize learned fear responses, whereas individuals with the low-activity variants may have better response to usual cognitive–behavioral therapies, and thus might require less intensive pharmacological interventions. Similar studies with ASB as the main outcome would bring findings on the MAOA–adversity–ASB relationship into clinical practice, allowing the retest of epidemiological, correlational, and experimental evidence in a still-unexplored ambit.

Variations in the 5-HTT gene and in two DRD4 polymorphisms were tested in an early intervention program devised to improve mother–child attachment in maltreated and nonmaltreated 2 year olds. Whereas no genetic effects were found in maltreated children, there was a greater proportion of risk gene carriers among insecurely attached, nonmaltreated controls. This result is suggestive that the possible genetic influences in the development of mother–child relationships in normative populations may be diminished in presence of maltreatment. Considering the strong associations between maternal attachment and all kinds of psychopathology, it would be of great value to test the role of the MAOA gene in attachment processes, and especially in the response to preventive interventions.

Overall, mapping patients’ genotypes seems to be especially useful in the prediction of stimulus-contingent outcomes, but it may not translate well to the prediction of general dispositions. Genetic effects for broad personality...
dimensions were found to be inconsistent in an exhaustive meta-analysis, with effect sizes of specific polymorphisms on personality being predominantly small to moderate. However, as already commented, the MAOA gene may predispose toward reactive aggression after provocation; it is therefore interesting to consider MAOA genotype as a predictive tool in forensic or penitentiary settings, where hostile interpersonal interactions are common and estimating the risk of recidivism is crucial. We further argue that inquiring about the past maltreatment history of inmates or indicted individuals, complemented with their genetic makeup, can give important clues when it comes to assessing the likelihood of future ASB arising from social interactions.

It is almost compulsory to cite, in this regard, the work by Bernet et al., who depicted a series of cases in which genetic information was presented as evidence in murder trials. The authors first reviewed precedent cases, in two of which they raised the possibility that conviction reductions could have been influenced by genotypic evidence. Subsequently, the authors related their own experience in the use of genotype in murder trials, where they exposed the extenuatory influence of either the MAOA or the 5-HTT risk allele in combination with past history of harsh discipline or maltreatment. It is noteworthy, however, that the authors selectively presented the genetic markers that most favored the defendants, obviating putative gene–gene interactive effects – only one case had the risk allele in both measured genes. Furthermore, they alleged a general predisposition toward violence conferred by genes instead of relating it to specific instances of the situation in which murder occurred. Nonetheless, genotypic evidence has also been presented in an Italian murder trial, in which the defense alleged genetic risk for aggressive reactions to provocation, without considering the defendant’s upbringings; in this case, the indicted saw his conviction reduced.

At this point we want to draw attention to the fact that utilizing specific genes in trials is a relatively novel practice, but the allegation of “genetic predisposition” – usually on the basis of familial aggregation – is more established; Pioro et al have recently found this term in 468 different legal decisions when reviewing Canadian judicial databases. Farahany and Coleman encouraged caution regarding the use of genetic factors as evidence in court, considering that behavioral genetic evidence was still poorly suited to apprehend individual cases, and that the prevailing concepts on liability – such as legal free will – drained a great deal of importance from genetic factors in law; see Baum for further discussion. Although we agree on the need for wariness until more-thorough studies come out, the recently observed effects of the MAOA gene in experimental situations, obtained with very small samples and in controlled conditions, permit greater confidence in G × E interaction results at an individual level of analysis. Therefore, we suggest that the use of genotype in trials should be limited to cases of impulsive (rather than premeditated) crimes, with clearly demonstrable risk factors (eg, childhood severe adversity), and offering explicit and specific relationships between genetic makeup and the circumstances in which the offense occurred – such as provocation – that might have served as a trigger.

Finally, although internalizing psychopathology is not our present focus, we will introduce a brief note on the topic due to its relevance in forensic practice. In this ambit, it is worth mentioning a recent meta-analysis that has discarded the implication of the MAOA gene on suicidal behavior. While the high activity variant of the MAOA gene seems to protect males from ASB, it may heighten liability to anxiety and depression in females. This has led some researchers to label the MAOA as the “warrior–worrier gene”, but more evidence is needed to test the validity of this somewhat simplistic tag. It appears wise, in future research, to explore how the MAOA gene may relate differently to the internalizing and externalizing spectra, or even with the recently derived general psychopathology factor, or p.

Conclusions and therapy implications
Future G × E studies should place special effort in sampling not only adverse environments, but also beneficial ones. The fact that presumed risk alleles have been related to greater positive responses to protective factors questions several results presented here, as only adverse environmental factors are considered in the majority of G × E studies. Although great effort is expended to make samples representative of the different socioeconomic strata, a more-comprehensive assessment of identifiable positive factors would allow testing of whether some alleles are indeed simple risk variants, or if they map into greater behavioral variability in the presence of both detrimental and reinforcing environments.

Even if we only focus on the pathological side, multidimensional measures of environmental features can help uncover differential response patterns; Kinnaly et al., for instance, used a self-report of parent care and a separate questionnaire of stressful life events (eg, death of a close relative, physical or sexual abuse), and, as explained in Table 2, found different effects for each environmental variable. Indeed, studies that go beyond the usual two-level or
three-level scale for adversity often find subtle effects that cannot be captured in dichotomic measures.49,56,101 These are just a few examples of how the impact of environmental variables can be diverse. This is of crucial importance for those theories of crime that focus only on environmental aspects, as disentangling the relative importance of each level of environmental factors can lead to more successful environmental interventions.

As we have seen so far, the MAOA gene, in conjunction with environmental adversity, seems to play a central role in the genesis of ASB and related outcomes. Even though there is still some controversy, epidemiological, clinical, and experimental research converges in providing evidence for it.

All in all, the take-home messages about incorporating G × E research in applied settings are three. First, with respect to ASB management, incorporating genetic data into current risk assessment procedures, such as HCR-20 or Offender Group Reconviction Scale,102 could improve risk management protocols, especially in forensic settings.103,104 Second, taking genetic data into account could be useful in order to detect individuals at a greater risk of victimization,49 or at least, the most vulnerable ones. And last but not least, the use of genetic data could be useful to compare the performance of specific psychological treatments versus specific pharmacological treatments, as Lester and Eley suggested.105

Future research in this area should seek to fill the gaps in four main points. First, research testing the incremental validity of genomics testing in risk management settings must be conducted, especially using longitudinal designs. Neuroprediction106 has proven successful in spite of criticisms concerning the reliability of neuroimaging data. Genetic data is often considered more reliable, though their effects are smaller. Also, modeling their interactions with environmental risk factors should lead to better risk management. Second, research on factors predisposing to victimization has been considered taboo, as some nonacademic organizations consider this as blaming the victims. However, this type of research would help focus our interventions on them. In turn, this would lead to a rationalization in the allocation of resources. Actually, personality allows identification of those individuals that lack most resources to cope with severe forms of maltreatment, such as childhood sexual abuse,11 and we should not forget that personality traits are highly heritable.106 Third, little is known about why different genotypes confer differential sensitivities to psychological treatments, although some hypotheses have been suggested. These hypotheses are yet to be tested. Finally, as some other researchers have pointed out, understanding the MAOA–adversity–ASB triad can lead to unraveling the causal mechanisms and therefore allow more-successful interventions.

In conclusion, the moderating role of the MAOA gene on the development of antisocial behavior is not only useful for basic research and advancing in the ethiological knowledge of ASB, but also for managing risk and interventions in applied clinical and forensic settings.

Disclosure
The authors report no conflicts of interest in this work.

References


