Cannabis use and childhood trauma interact additively to increase the risk of psychotic symptoms in adolescence

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Background. Adolescent cannabis use has been shown in many studies to increase the risk of later psychosis. Childhood trauma is associated with both substance misuse and risk for psychosis. In this study our aim was to investigate whether there is a significant interaction between cannabis use and childhood trauma in increasing the risk for experiencing psychotic symptoms during adolescence.

Method. Psychiatric interviews using the Schedule for Affective Disorders and Schizophrenia for School-Age Children (K-SADS) semi-structured instrument were carried out with 211 adolescents aged between 12 and 15 years and their parents as part of a population-based study. The interview enquired about early traumatic events, cannabis use and psychiatric symptoms in adolescence.

Results. In separate analyses both cannabis use and childhood trauma were significantly associated with risk of experiencing psychotic symptoms. However, the presence of both childhood trauma and early cannabis use significantly increased the risk for psychotic symptoms beyond the risk posed by either risk factor alone, indicating that there was a greater than additive interaction between childhood trauma and cannabis use.

Conclusion. Our finding of a greater than additive interaction between childhood trauma and cannabis use may have implications for the identification of individuals at high risk of experiencing psychotic symptoms. For example, measures to actively discourage or intensively treat cannabis use in children and adolescents who have experienced abuse may help to prevent the development of psychosis in this vulnerable group. Our findings require replication in larger samples to confirm this interaction effect.

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Introduction

Cannabis use has been increasing over the past few decades (Gledhill-Hoyt et al. 2000) and age at first use has been decreasing in Western societies (Degenhardt et al. 2000). There are particular concerns about the use of cannabis in adolescence as the brain is still developing (Giedd et al. 1999) and may be vulnerable to the potentially toxic effects of cannabis (Arseneault et al. 2002; Cannon et al. 2006). Cannabis use has been found to be associated with mental health problems, most notably psychotic disorders (Arseneault et al. 2004; Moore et al. 2007), although it is a matter of some debate whether the relationship is causal in nature (Macleod et al. 2004, 2006).

Recent studies have shown that childhood trauma is a risk factor for the development of schizophrenia (Read et al. 2005; Morgan & Fisher, 2007; Bendall et al. 2008) and psychotic symptoms (Bak et al. 2005; Kelleher et al. 2008). It has also been shown that childhood trauma increases the risk of substance misuse in adolescence and adulthood (De Bellis, 2002; Gordon, 2002). Taken together, these strands of evidence point to the possibility that childhood trauma may have a role to play in the association between cannabis use and psychosis (Cougnaud et al. 2007; Houston et al. 2008).
There is increasing interest in the role of interactions (gene–environment, gene–gene and environment–environment) in the aetiology of psychiatric disorders (van Os & Sham, 2003; Moffitt et al. 2005; Caspi & Moffitt, 2006; van Os et al. 2008; Clarke et al. 2009). However, there is some debate about the usefulness of such findings and the correct model to use for the investigation of psychiatric outcomes (Risch et al. 2009; Zammit et al. 2009). Predictions can be modelled on either additive or multiplicative scales. A multiplicative model assumes that risks multiply in their effect. An additive model assumes that individuals can develop the outcome of interest from either of the risk factors acting alone and tests for positive deviations from additivity (superadditivity) that indicate the presence of synergy. The additive model is considered to be the more meaningful model in psychiatric epidemiology, where diseases usually have complex multifactorial aetiologies (Darroch, 1997; van Os & Sham, 2003; Schwartz & Susser, 2006).

In this study we aimed to investigate whether the presence of both childhood trauma and early cannabis use increases the risk of experiencing psychotic symptoms in adolescence beyond that expected if each risk factor were working independently. The importance of psychotic symptoms in childhood and adolescence lies in mounting evidence that they are a risk marker for later psychotic illness (Poulton et al. 2000; Scott et al. 2008).

Method

Participants

The ‘Challenging Times’ study (Lynch et al. 2004, 2006; Mills et al. 2004) was established to investigate the prevalence of psychiatric disorders among Irish adolescents aged 12–15 years in an urban environment. The study was carried out in the geographical catchment area of a Child and Adolescent Mental Health Team in North Dublin with a population of 137,000. Participating schools were selected using a stratified random sampling technique, stratified according to the approximate socio-economic class of the school to approximate to the geographical area population. In brief, 743 pupils in eight mainstream schools (52% of the total school population in that area) were screened for psychiatric symptoms using the Strengths and Difficulties Questionnaire (SDQ), which assesses emotional symptoms, conduct problems, hyperactivity/inattention, peer relationship problems, and prosocial behaviour (Goodman, 1997; Goodman et al. 2000). The Children’s Depression Inventory (CDI), which assesses cognitive, affective and behavioural signs of depression, was also used (Kovacs, 1985). After complete description of the study to the subjects, written informed consent was obtained from their parent or guardian. One hundred and forty adolescents scored above threshold on these instruments, indicating high risk of having mental health problems. Adolescents were included in this ‘at-risk’ category if they scored in the clinical range on the CDI and/or if they scored in the clinical range on the SDQ, and/or if they ticked ‘I want to kill myself’ on item 9 of the CDI. Of these 140 adolescents, 117 (83.6%) agreed to attend for full psychiatric interview. A comparison group of 173 adolescents, matched for gender and school, were also invited to attend, of whom 94 (54%) agreed.

Ethical approval for the study was granted by the Medical Ethics Committee of the Mater Misericordiae University Hospital, Dublin, Ireland. The study was supported by the multi-disciplinary Child and Adolescent Mental Health Service covering the geographical area. The protocol ensured that any adolescent who was deemed to be in need of a clinical service could be referred to the appropriate team.

Interview instrument

The interview schedule used in this study was the Schedule for Affective Disorders and Schizophrenia for School-Age Children, Present and Lifetime Versions (K-SADS-PL; Kaufman et al. 1996). The K-SADS is a well-validated semi-structured research diagnostic interview for the assessment of all Axis-I psychiatric disorders in children and adolescents. Children and parents were interviewed separately, both answering the same questions about the child. Interviews were conducted by one psychiatrist or two psychologists who were trained in the use of the K-SADS. Inter-rater reliability for the K-SADS was estimated as over 90% in this study (Lynch et al. 2006).

Psychotic symptoms

Axis I psychiatric diagnoses according to DSM-IV (APA, 1994) were determined using the K-SADS semi-structured interview with adolescents and parents or guardians. The Psychosis section of the K-SADS asks about the child’s experience of hallucinations and delusions not occurring during acute intoxication with a substance. Responses to these questions were recorded on the interview sheet. Three psychiatrists (M.H., F.L. and M.C.) and one psychologist (I.K.) examined these responses and concurred that the symptoms seemed genuine in content.

Measurement of substance misuse

During the interview, in the Substance Misuse section of the K-SADS, both adolescent and parent were asked...
whether the adolescent had ever used cannabis. If the answer was yes, then frequency of use in the past 6 months and frequency ever were recorded. Cannabis use was defined as ever having used cannabis.

**Measurement of childhood trauma**

As part of the K-SADS interview the following childhood traumatic experiences were assessed. Interviews were conducted with parents and adolescents separately.

**Childhood abuse.** Both child physical abuse and child sexual abuse were assessed as part of the K-SADS interview. Adolescents were asked a series of questions in relation to physical or sexual abuse. Parents were asked the same questions appropriately modified. A disclosure of physical or sexual abuse from the parent was taken as evidence of a history of child abuse, regardless of whether it was also disclosed by the child. There were no cases where the parent disputed the occurrence of abuse that had been disclosed by the child.

**Domestic violence.** Exposure to domestic violence was assessed in the Post-Traumatic Stress Disorder section. Adolescents and their parents were separately asked the same questions. A disclosure of domestic violence from the parent was taken as evidence of violence between parents/step-parents. There were no cases where the parent’s report of the occurrence of domestic violence disagreed with that disclosed by the child or vice versa. Childhood trauma was defined as a history of physical abuse, sexual abuse, witnessing domestic violence or any combination of these.

**Socio-economic status (SES) and family history**

SES of each study participant was determined using parental occupation assessed according to the Irish Social Class Scale (Central Statistics Office, 1996). We divided the sample into two major groups according to social class: the first group contained SES groups 1 and 2 (professional/managerial) and the second group contained SES groups 3–7 (non-manual skilled, skilled manual, semi-skilled manual, unskilled manual, and unemployed). The K-SADS interview includes a routine screening section for family history of psychiatric illness, which was used in the present study. Family psychiatric history included history of affective disorders, anxiety disorders, psychotic and substance use disorders in a first- or second-degree relative.

**Statistical analysis**

An additive model of interaction assumes that risks add in their effects so that results over and above additivity indicate synergy. However, when examining the interaction of two risk factors, it is important to note that some individuals can develop disease from either one of the two risk factors under study alone; this phenomenon is termed parallelism and it reduces the measure of the exposures’ combined effect. An additive model of interaction therefore includes the supperladditive effect of the interaction of two causal partners, and the subadditive effect of parallelism. The details of the analysis are as follows: if we have two risk factors for psychosis A and B, there are four possible exposure states, each carrying a specific risk. Therefore, the risk in those exposed to neither A nor B is R; the risk in those exposed to A only is R(A), the risk in those exposed to B only is R(B), and the risk in those exposed to both is R(AB). On the additive scale, the effect of a risk factor is expressed as a risk difference. We can therefore express the effect associated with A as R(A) − R, with B as R(B) − R and with AB exposure as R(AB) − R. The excess of the combined effect over the sum of the solitary effects of A and B is R(AB) − R(A) − R(B) + R; this represents the statistical additive interaction.

We wanted to examine for additive interaction rather than multiplicative interaction in this study (Darroch, 1997; van Os & Sham, 2003; Cougnard et al. 2007). Because of the case-control design of the Challenging Times study, however, it was not possible to estimate the baseline risk of disease and use risk differences in this analysis. Schwartz & Susser (2006) have described a method whereby estimates from a multiplicative model can be used to assess interaction on an additive scale. This adaptation allows the use of odds ratios (ORs) instead of risk differences. In this study, we used logistic regression for the initial analyses and translated these to look at interaction on an additive scale using an interaction contrast ratio method (Rothman & Greenland, 1998; Schwartz & Susser, 2006). This ratio is calculated as: R(AB) − R(A) − R(B) + 1.

In brief, we stratified our data into four categories: (1) no exposure to cannabis use or childhood trauma [R], (2) exposure to cannabis use only [R(A)], (3) exposure to childhood trauma only [R(B)], and (4) exposure to both cannabis use and childhood trauma [R(AB)]. The outcome measure was self-reported psychotic symptoms. The ORs for groups 2, 3 and 4 were calculated using group 1 as baseline. The ORs were then used in the standard formula for calculating an interaction contrast ratio. The interaction contrast ratio divided by the OR in those exposed to both risk factors
can be interpreted as the proportion of disease among those with both risk factors that is attributable to the interaction.

All analyses were carried out using Stata Statistical Software release 9.2 (Stata Corporation, USA).

Results

Of the 211 children in the interviewed sample, 83 (39%) received a DSM-IV Axis I diagnosis following the K-SADS interview. The most common disorders diagnosed were depressive disorders (17.5%, n = 37). No participant received a formal diagnosis of a psychotic illness. Fourteen participants (6.6%) reported experiencing psychotic symptoms, primarily auditory and visual hallucinatory experiences.

Substance use

No adolescents in this sample met criteria for a cannabis dependence syndrome. Of the adolescents interviewed, 8.5% (n = 18) reported having ever used cannabis. Only five adolescents reported ever having used any illicit substance (ecstasy, ‘poppers’, snorting deodorant or petrol) other than cannabis and four of these had also used cannabis. Adolescents who had used cannabis were five times more likely to have experienced at least one psychotic symptom (see Table 1).

Childhood trauma

Childhood trauma was reported in 11.3% of participants interviewed (n = 24). Those adolescents who had experienced childhood trauma were almost five times more likely to use cannabis [OR 4.86, 95% confidence interval (CI) 1.63–14.51, p = 0.005] than those who had not experienced trauma. Participants who had experienced childhood trauma were also five times more likely to develop psychotic symptoms (OR 5.20, 95% CI 1.58–17.13, p = 0.007) compared to those who had not (Table 1).

Examining the relationship between cannabis, childhood trauma and psychotic symptoms

Fig. 1 shows the prevalence of psychotic symptoms in each of the four exposure categories: No trauma or cannabis use; Cannabis use only; Trauma only; Trauma and cannabis use. Using these ORs in the formula for additive interaction [R(AB) = R(A) × R(B) + R] gave a value of 17.4 [20.9 − 1.9 − 2.6 + 1]. The statistical additive interaction is positive, indicating that childhood trauma and cannabis use interact on the additive scale. We can estimate from our data that 83% (17.4/20.9) of the occurrence of psychotic symptoms among those exposed to both cannabis use and childhood trauma is attributable to the interaction between these factors.

Discussion

In this study we found that both cannabis use and childhood trauma increased the risk for psychotic symptoms in adolescence in main effect analyses. However, when we examined both risk factors under an additive model we found evidence of a greater than additive interaction between childhood trauma and cannabis use, whereby the effect of the joint presence

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**Table 1. Risk of psychosis associated with childhood trauma and cannabis use as main effects**

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Total no.</th>
<th>Psychosis n (%)</th>
<th>Unadjusted OR (95% CI)</th>
<th>Adjusted OR a (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trauma</td>
<td>24</td>
<td>5 (21)</td>
<td>5.20 (1.6–17.1)</td>
<td>6.16 (1.65–23.1)</td>
</tr>
<tr>
<td>Cannabis</td>
<td>18</td>
<td>4 (22)</td>
<td>5.23 (1.4–18.8)</td>
<td>4.32 (1.1–17.3)</td>
</tr>
</tbody>
</table>

OR, Odds ratio; CI, confidence interval.

a Adjusted for gender, age, socio-economic status and family psychiatric history.
Table 2. Risk of psychotic symptoms in each exposure category

<table>
<thead>
<tr>
<th>Risk exposure category</th>
<th>Total no.</th>
<th>Psychosis, n (%)</th>
<th>OR (95% CI)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>No cannabis use/no trauma</td>
<td>175</td>
<td>8 (4.6)</td>
<td>1.0 (baseline)</td>
<td></td>
</tr>
<tr>
<td>Cannabis use/no trauma</td>
<td>12</td>
<td>1 (8.3)</td>
<td>1.9 (0.04–16.5)</td>
<td>0.55</td>
</tr>
<tr>
<td>Trauma/no cannabis use</td>
<td>18</td>
<td>2 (11.1)</td>
<td>2.6 (0.25–14.6)</td>
<td>0.23</td>
</tr>
<tr>
<td>Trauma/cannabis use</td>
<td>6</td>
<td>3 (50.0)</td>
<td>20.9 (2.3–173.5)</td>
<td>0.000</td>
</tr>
</tbody>
</table>

OR, Odds ratio; CI, confidence interval.

What are the possible biological mechanisms of an interaction effect?

There are several possible biological, genetic and psychosocial factors that could be involved in the association between cannabis use, trauma and psychotic symptoms. The most parsimonious explanation involves effects on the dopamine neurotransmitter system. First, all effective antipsychotic medications affect dopamine and dopamine sensitization has been proposed as a mechanism explaining psychotic symptoms (Laruelle, 2000; Kapur et al. 2005). Second, it has been demonstrated that cannabis affects dopamine release and that the direction of effect depends on the chronicity of use (Verrico et al. 2003; Luzi et al. 2008; Bossong et al. 2009). Third, there is evidence from both animal (Plotsky & Meaney, 1993; Ladd, 1996; Liu et al. 2000; Kalinichev et al. 2002) and human studies (Heim et al. 2000; Carpenter et al. 2007) that response to severe stress, such as childhood abuse early in life, can lead to permanent changes in the hypothalamic–pituitary–adrenal (HPA) axis and dopamine systems (for discussion see Read et al. 2001).

Limitations and strengths of study

Although the ORs in this study are high, the CIs are wide, reflecting a relatively small sample size when subgroups are compared. The Challenging Times study was designed as a general epidemiological study of mental health in adolescents, not specifically to test the current hypotheses. Although this may have led to less detailed information obtained specifically in relation to cannabis use and psychotic symptoms, it is likely that it minimized interviewer bias. Another limitation was that information on the timing of events (first use of cannabis, first experience of childhood trauma) was limited and relied on retrospective recall. However, recent data suggest that the temporal ordering of childhood trauma and cannabis use does not affect the risk of developing psychosis (Shewlin et al. 2009). Rates of cannabis use were relatively low in this sample. This may be due to under-reporting of...
substance use, although participants were assured of the confidentiality of their data and collateral information was obtained from at least one parent/guardian. It should also be noted that the measure of sexual abuse used in this study is a conservative one, which may explain the low rates of sexual abuse reported.

The strengths of this study include the use of an epidemiological community-based sample, whereby schools were selected using a randomized stratified sampling method so that the study population is representative of the area population. A well-validated standardized psychiatric interview was used in this study and administered by trained researchers with a clinical background. Information was also obtained from parents/carers so that information on psychiatric symptoms, on environmental risk factors, and on a variety of potential confounding factors was available from two sources.

Conclusions

In this study, we report a greater than additive interaction between childhood trauma and cannabis use in increasing the likelihood of developing psychotic symptoms. Our findings should be replicated in a larger sample of adolescents using a prospective design to clarify the temporal relationship between risk factors and symptoms. Future studies examining links between cannabis use and psychosis should consider the effects of childhood trauma as an important potential effect modifier. The findings reported here may have important practical implications for the prevention of psychosis. Young people with a history of childhood trauma could be conceptualized as an ‘at-risk’ group and could be targeted for psychotherapeutic intervention, if warranted, or psycho-educational interventions concerning risks of substance use, particularly during adolescence.

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Declaration of Interest

None.

References


