Continued cannabis use and risk of incidence and persistence of psychotic symptoms: 10 year follow-up cohort study

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ABSTRACT

Objective To determine whether use of cannabis in adolescence increases the risk for psychotic outcomes by affecting the incidence and persistence of subclinical expression of psychosis in the general population (that is, expression of psychosis below the level required for a clinical diagnosis).

Design Analysis of data from a prospective population based cohort study in Germany (early developmental stages of psychopathology study).

Setting Population based cohort study in Germany.

Participants 1923 individuals from the general population, aged 14-24 at baseline.

Main outcome measure Incidence and persistence of subthreshold psychotic symptoms after use of cannabis in adolescence. Cannabis use and psychotic symptoms were assessed at three time points (baseline, T2 (3.5 years), T3 (8.4 years)) over a 10 year follow-up period with the Munich version of the composite international diagnostic interview (M-CIDI).

Results In individuals who had no reported lifetime psychotic symptoms and no reported lifetime cannabis use at baseline, incident cannabis use over the period from baseline to T2 increased the risk of later incident psychotic symptoms over the period from T2 to T3 (adjusted odds ratio 1.9, 95% confidence interval 1.1 to 3.1; P=0.021). Furthermore, continued use of cannabis increased the risk of persistent psychotic symptoms over the period from T2 to T3 (2.2, 1.2 to 4.2; P=0.016). The incidence rate of psychotic symptoms over the period from baseline to T2 was 31% (152) in exposed individuals versus 20% (284) in non-exposed individuals; over the period from T2 to T3 these rates were 14% (108) and 8% (49), respectively.

Conclusion Cannabis use is a risk factor for the development of incident psychotic symptoms. Continued cannabis use might increase the risk for psychotic disorder by impacting on the persistence of symptoms.

INTRODUCTION

Cannabis is the most commonly used illicit drug in the world, particularly among adolescents.12 The use of cannabis is consistently associated with mental illness,4 in particular psychotic disorder.4-9 It remains a matter of debate, however, whether the association between cannabis and psychosis is causal, or whether early psychotic experiences might in fact prompt cannabis use as a means of self medication.10,11 This issue can be resolved only if incident cannabis use is investigated in relation to later incident psychotic symptoms or disorder. Rarely have studies been able to examine the longitudinal relation between cannabis use and psychosis in this fashion.

The issue of self medication was addressed by Henquet and colleagues, using data from the German prospective early developmental stages of psychopathology study.11,12 The authors investigated the association between cannabis use at baseline and subsequent development of psychotic symptoms at four year follow-up and reported that after adjustment for pre-existing psychotic symptoms, cannabis use at baseline still remained significantly associated with psychotic symptoms at follow-up. There was no evidence of an effect of self medication as pre-existing psychotic symptoms did not significantly predict later cannabis use. Ferdinand and co-workers investigated the role of pre-existing self reported psychotic symptoms and showed a bi-directional association between cannabis and psychotic symptoms over a 14 year follow-up study in the general population.13 They showed that cannabis use predicted later psychotic symptoms in individuals with no evidence of psychotic symptoms before starting to use cannabis and that the reverse was also true, in that psychotic symptoms predicted cannabis use in those who had not used cannabis before the onset of those symptoms.13 A prospective population based cohort study also found evidence for a self medication effect.14 Individuals with self reported hallucinations at the age of 14 had a higher risk of using cannabis on a daily basis at the age of 21. In a sibling pair analysis, however, this study also suggested an independent effect of cannabis use on self reported delusional ideation later in life.14 Thus, although the cannabis-psychosis link has been investigated in
many studies, results on the temporal association between cannabis use and psychotic symptoms remain conflicting. Longitudinal cohort studies with multiple repeated interview based measures of cannabis use and psychotic symptoms are needed to clarify this issue. The EDSP study\textsuperscript{12,13} which completed its recent 10 year follow-up representing the fourth assessment (assessments at baseline, T1, T2, and T3, see also fig 1), is uniquely suitable for the investigation of the temporal association between cannabis and psychosis.

Another issue is the mechanism by which cannabis might increase the risk of psychotic symptoms, particularly whether it might increase the risk by causing persistence of normally transitory developmental expression of psychotic experiences. For most individuals, subclinical expression of psychotic phenomena (that is, expression of psychosis below the level required for a clinical diagnosis) is transitory and never progresses to psychotic illness.\textsuperscript{15} Subthreshold psychotic experiences could, however, become abnormally persistent, depending on the degree of additional exposure to environmental risk factors,\textsuperscript{16-18} and progressively greater levels of persistence might be associated with a greater risk for transition to clinical psychotic disorder.\textsuperscript{19} Spauwen and colleagues showed that the persistence rate of psychotic experiences was much higher for individuals growing up in an urban rather than a rural environment.\textsuperscript{20} Similarly, Cougnard and co-workers provided evidence that childhood trauma, urban environment, and cannabis act additively in increasing the risk of persistence of psychotic experiences.\textsuperscript{17} The fact that cannabis use increases risk of psychosis in a dose-response fashion\textsuperscript{3-14,19,20} and that patients with psychosis who continue to use cannabis show more severe and persistent symptoms\textsuperscript{21} suggests that cannabis use might increase the risk for psychotic illness by impacting on the persistence rate of psychotic experiences that under normal circumstances (that is, without exposure to cannabis) would have remained transitory phenomena for most people. In a population based 10 year follow-up cohort study of adolescents and young adults, we investigated the association between incident cannabis use and true incidence of psychotic experiences (that is, after exclusion of individuals with lifetime pre-existing psychotic experiences) and risk of persistence of psychotic experiences.

**Table 1** Characteristics of participants by use of cannabis at T2 (3.5 years after baseline) (n=1923). Figures are numbers (percentages)

| Characteristics of participants by use of cannabis at T2 (3.5 years after baseline) (n=1923). Figures are numbers (percentages) | Used cannabis (n=393) | Did not use cannabis (n=1530) |
|---|---|---|
| Men | 119 (30) | 807 (53) |
| Women | 274 (70) | 723 (47) |
| Socioeconomic status*: | | |
| Lower | 30 (7) | 85 (6) |
| Middle | 220 (54) | 894 (59) |
| Upper | 137 (34) | 524 (35) |
| Other | 22 (5) | 6 (0.4) |
| Urban/rural environment‡: | | |
| Urban | 295 (75) | 1050 (69) |
| Rural | 98 (25) | 480 (31) |
| Childhood trauma*: | | |
| Yes | 97 (25) | 266 (17) |
| No | 296 (75) | 1264 (83) |
| Use of other drugs at baseline§: | | |
| Yes | 25 (6) | 11 (1) |
| No | 368 (94) | 1519 (99) |
| Use of other drugs at T2§: | | |
| Yes | 40 (10) | 3 (0.2) |
| No | 353 (90) | 1527 (99) |
| Any psychiatric disorder at baseline**: | | |
| Yes | 76 (19) | 180 (12) |
| No | 317 (81) | 1350 (88) |
| *Socioeconomic status: lower (lower class, lower middle class), middle (middle middle class), upper (higher middle class, upper class), other (none of the above or missing). Data missing for five participants. |
| ‡Urban (city of Munich, 10 559/km\(^2\)), rural (surroundings of Munich, 1432/km\(^2\)). |
| §Childhood trauma: any traumatic experience during childhood. |
| §On more than five occasions. |
| **Other than psychosis, according to M-CIDI diagnoses. |

**Fig 1** Study design. Top: testing association between incident cannabis use with onset in period from baseline to T2 and incident psychotic symptoms with onset in period from T2 to T3 in individuals who had not used cannabis at baseline and who had not reported any psychotic experience at T2 (that is, no lifetime psychotic experiences by T2). Bottom: testing association between different cannabis exposure states (combinations of cannabis use at baseline (lifetime), or T2 or both (interval) and persistence of psychotic experiences (that is, presence of psychotic experiences at both T2 (lifetime) and T3 (interval))]
obtained, and registration is strictly enforced by law and the police. More details on the sampling, representativeness, instruments, procedures, and statistical methods of the study sample can be found elsewhere.\textsuperscript{12,13}

The overall design of study is longitudinal and prospective, consisting of a baseline and three follow-up surveys. The first (baseline to T1) covered a mean of 1.6 years (SD 0.2), the second (baseline to T2) covered a mean of 3.5 years (SD 0.3), and the third (baseline to T3) covered a mean of 8.4 years (range 7.3-10.5, SD 0.7). Because our primary goal was to examine the incidence and developmental risk factors for psychosis, we sampled the younger group (age 14-15), presumed to have the highest incidence density, at twice the rate of people aged 16-21 and sampled the oldest group (age 22-24) at half this rate. For the same reason we examined people aged 14-17 at baseline at the four time points and those aged 18-24 only three times (baseline, T2, T3). The present study is based on the whole cohort assessed at baseline, T2, and T3. Figure 1 shows a schematic illustration of the overall design and the current analyses. Response rates were 84\% (2548) at T2 and 73\% (2210) at T3 (fig 2).

**Instruments**

We used the computerised version of the Munich composite international diagnostic interview (DIA-X/M-CIDI),\textsuperscript{22} an updated version of the World Health Organization’s CIDI version 1.2.\textsuperscript{23} The DIA-X/M-CIDI is a comprehensive fully standardised diagnostic interview that assesses symptoms, syndromes, and diagnoses of various mental disorders in accordance with the definitions and criteria of ICD-10 (international classification of diseases, 10th revision) and DSM-IV (Diagnostic and Statistical Manual of Mental Disorders, fourth edition), along with information about onset, duration, severity of symptoms, and psychosocial impairment. The CIDI has been primarily designed for use in epidemiological studies of mental disorders and can also be used for clinical purposes. It is divided into 16 sections; one sociodemographic section, 12 sections assessing 288 symptoms of groups of mental disorders (including somatoform and dissociative, phobic and other anxiety, depressive and dysthymic, manic and bipolar affective, schizophrenia and other psychotic, eating, dementia and other cognitive, post-traumatic stress, as well as tobacco, alcohol, and substance related disorders), and three final sections containing concluding questions, interviewers’ observations, and interviewers’ ratings. The instrument, designed for use by trained interviewers who are not clinicians, has shown high inter-rater\textsuperscript{24,25} and test-retest reliability.\textsuperscript{22,26} The assessment of psychosis with CIDI by lay interviewers is not considered reliable so trained clinical interviewers at the level of clinical psychologist, who were allowed to probe with follow-up clinical questions, conducted the interviews in the respondents’ homes. At baseline, the DIA-X/M-CIDI lifetime version was used. At each of the follow-up assessments, the interval version was used to assess the period from the last interview until the next. Data on the G section concerning psychosis and its clinical relevance were collected only at T2 (lifetime version) and T3 (interval version). As the assessment of substance use was part of the diagnostic interview with the DIA-X/M-CIDI, psychologists who did the interviews were not blinded for cannabis use.

**Assessment of psychotic symptoms**

Data on psychotic experiences were collected at time T2 (lifetime version) and T3 (interval version) with the G section of the DIA-X/M-CIDI. As the primary objective of the EDSP study was to investigate the early stages of substance misuse in adolescents and young adults, data on the occurrence of psychotic symptoms as assessed with the G section were added at T2 (measuring lifetime experience of psychotic symptoms) and T3 (measuring interval experience of psychotic symptoms).\textsuperscript{13} As in previous work\textsuperscript{6,9} presence of psychotic experiences was broadly defined as any rating of present on any of the 20 DIAx/M-CIDI core psychosis items (G1, G2a, G3-G5, G7-G13, G13b, G14, G17, G18, G20, G20C, G21, and G22a), including 14 delusion items, five hallucination items, and one item on passivity phenomena. Items relate to classic psychotic symptoms involving, for example, persecution, thought interference, auditory hallucinations, and passivity phenomena. The

**Table 2 Patterns of cannabis use in relation to presence of psychotic symptoms**

| Cannabis use‡ | Psychotic symptoms at T2 | Psychotic symptoms at T3 |
|---------------|--------------------------|--------------------------|
|               | Yes          | No          | Yes          | No          |
| Baseline      |              |             |              |             |
| Yes           | 81 (4)       | 166 (9)     | 42 (2)       | 205 (11)    |
| No            | 355 (18)     | 1321 (69)   | 189 (10)     | 1487 (77)   |
| T2            |              |             |              |             |
| Yes           | 126 (7)      | 267 (14)    | 69 (4)       | 324 (17)    |
| No            | 310 (16)     | 1220 (64)   | 162 (8)      | 1368 (71)   |

*Any psychotic symptom lifetime (T2) and interval (T3) as assessed with M-CIDI (G) section.
†Some percentages do not total 100 because of rounding.
‡On more than five occasions as assessed with M-CIDI (L) section.
psychologist invited participants to read a list of all the psychotic experiences and then asked them whether they ever experienced such symptoms (list and phrasing available on request). All these psychosis items were rated as absent or present. Thus, all the psychosis DIA-X/M-CIDI items used in the present study were coded in a dichotomous manner without intermediate levels. Psychotic symptoms at T2 accordingly referred to lifetime occurrence of at least one positive rating on any psychosis item in the G section; T3 psychotic symptoms denote interval occurrence of at least one positive rating on any psychosis item in this section.

Assessment of cannabis use
Cannabis use was assessed with the L-section of the DIA-X/M-CIDI at all three assessments. The question “Have you ever used cannabis five times or more?” was used to define cannabis exposure. As with previous work in this sample, the CIDI cut off of use of five times or more was used to define cannabis exposure (yes versus no). Thus, at baseline, cannabis use was defined as lifetime use of cannabis of five times or more (“baseline cannabis use,” dichotomously defined yes versus no). At T2, and similarly at T3, cannabis use was defined as interval use of cannabis of five times or more, during the intervals of baseline to T2 and T2 to T3, respectively (T2 cannabis use, T3 cannabis use, both dichotomously defined yes versus no). As assessment of frequency of cannabis use differed between time points (for instance, at baseline: How often do you use cannabis on a five point scale ranging from less than monthly to almost daily?; at T2: How many times have you used cannabis since the last assessment?), the frequency variable was not included in the analyses.

Statistical analyses
We used Stata, release 10.0 (StataCorp, College Station, TX) for data analysis and expressed associations as odds ratios from logistic regression models. All analyses were a priori adjusted for age at baseline (in years), sex (0=female, 1=male), socioeconomic status (lower, middle, upper, other, as assessed at baseline), use of other drugs at baseline and T2, childhood trauma (that is, trauma before the age of 14 as assessed at baseline),27 and urban/rural environment.28 29 The variable use of other drugs included psychostimulants, sedatives, opiates, cocaine, phencyclidine, and psychedelic drugs and refers to CIDI rated use of five times or more (lifetime or interval use at baseline and T2 respectively) of any of these drugs, as assessed with the L section of the M-CIDI. Childhood trauma refers to experience of any traumatic experience during childhood as assessed with the N section of the M-CIDI at baseline and was dichotomously defined as present or absent. Fisher and colleagues have shown that self report questionnaires constitute a valid method of assessing childhood trauma in individuals presenting with psychotic features.30 Urban/rural environment was assessed at baseline by retrieving data on place of residence from the population registry offices of the city and the 29 counties of Munich and was defined as either living in the city of Munich (10559 people per km²) or in the rural surroundings (1432 people per km²) at baseline.

To examine whether associations between cannabis use and psychotic symptoms were independent of other psychiatric diagnoses at baseline, we additionally adjusted analyses for any psychiatric diagnosis other than psychosis at baseline. We used a dichotomous variable of any psychiatric diagnosis (present versus absent), summarising main psychiatric diagnoses other than psychosis at baseline (such as major depressive disorder, hypomania, dysthymic disorder, bipolar disorder, panic disorder, obsessive compulsive disorder, post-traumatic stress disorder, conversion disorder, somatisation disorder, eating disorders).

We carried out sensitivity analyses to investigate whether attrition occurred at random and to assess potential bias introduced by missing data. This was done by multiple imputation of missing values with the ICE routine in Stata 11.1. This method imputes several alternative versions of the complete dataset from the available data.

Incident use of cannabis and incidence of psychotic symptoms
We used logistic regression analyses to investigate the association between incident cannabis use from baseline to T2 and incident psychosis outcome from T2 to T3 (see fig 1, top). To investigate the true incidence of psychotic symptoms after cannabis use, rather than persistence of pre-existing psychotic experiences, we excluded from the analysis individuals who had admitted to lifetime presence of any psychotic symptom at T2 (n=574, 23%). We also excluded all individuals with cannabis use at baseline, thus including only individuals with new cannabis exposure between baseline and T2.

We investigated reverse causality (that is, self medication) by testing the association between psychosis at baseline-T2 and cannabis use at T2-T3 in those who had not used cannabis at both baseline and T2.

Cannabis use and risk of persistent psychotic symptoms
We calculated a dichotomous persistence variable (no persistence versus persistence), with no persistence referring to experience of psychotic symptoms either never or only once (at T2 or T3) and persistence
referred to experience of psychotic symptoms at both T2 and T3. To investigate whether different levels of exposure to cannabis affected persistence of psychotic symptoms differentially, we calculated a categorical cannabis continuation variable (0 = never used cannabis; 1 = cannabis use at baseline but not at T2; 2 = cannabis use at T2 but not at baseline, 3 = cannabis use at both baseline and T2). We used logistic regression analyses to investigate the association between cannabis continuation and psychosis persistence (fig 1 bottom).

RESULTS
Sample
A total of 2210 individuals completed the T3 assessment; information on substance use and psychotic symptoms was missing for 287 participants, resulting in a final risk set for analysis of 1923 individuals, of whom 926 (48%) were men (fig 2). Mean age was 18.3 (SD 3.3) at baseline, 21.8 (SD 3.4) at T2, and 26.6 (SD 3.3) at T3. Table 1 gives further characteristics of participants.

At baseline, 247 participants reported using cannabis (13% lifetime use). Of those, 56 participants (23%) used cannabis almost every day, 69 (28%) reported weekly use, 57 (23%) used cannabis monthly, and 65 (26%) reported using cannabis less than once a month. At T2, 392 participants reported using cannabis (20% interval use). The mean frequency of those who reported cannabis use at T2 was 130 times within the period from baseline to T2 (range 5-997). Psychotic symptoms were reported by 436 participants (23%) at T2 (lifetime) and by 231 participants (12%) at T3 (interval). Table 2 summarises patterns of cannabis use in relation to psychotic symptoms.

Incident cannabis use and incidence of psychotic symptoms
The incidence rate of psychotic symptoms over the period from baseline to T2 was 31% (152) in exposed individuals and 20% (284) in non-exposed individuals; over the period from T2 to T3 these rates were 14% (108) and 8% (49), respectively.

Lifetime cannabis use as assessed at T2 significantly increased the risk of psychotic experiences at T3 (adjusted odds ratio 1.5, 95% confidence interval 1.1 to 2.1; P = 0.018; tables 3 and 4). After we excluded all individuals who had used cannabis at baseline and had reported psychotic experiences at T2, incident cannabis use over the period from baseline to T2 similarly increased the risk for incident psychotic experiences between T2 and T3 (1.9, 1.1 to 3.1; P = 0.021).

Results were unchanged after additional adjustment for any psychiatric diagnosis other than psychosis at baseline (1.9, 1.1 to 3.1; P = 0.020).

There was no evidence for self medication effects, as psychotic experiences at T2 did not predict incident cannabis use between T2 and T3 (0.8, 0.6 to 1.2; P = 0.3).

Cannabis use and persistence of psychotic experiences
Analyses showed a significant association between continuation of cannabis use and risk of persistence of psychotic experiences (adjusted \( \chi^2 = 16.22, P = 0.001 \); tables 4 and 5). After adjustment for age, sex, socioeconomic status, use of other drugs at baseline and T2, urban/rural environment, childhood trauma, and occasional cannabis use (cannabis use at either baseline or T2, exposure states 1 and 2), effect sizes were attenuated, with significance only for the combination of cannabis use at both baseline and T2 (exposure state 3; adjusted odds ratio 2.2, 1.2 to 4.2; P = 0.016). Additional adjustment for any psychiatric diagnosis other than psychosis at baseline occasioned a slight reduction in the association between cannabis use at both baseline and T2 and the risk for persistence of psychotic symptoms (2.0, 1.0 to 3.8; P = 0.037).

Missing data
Imputation of 20 datasets with the ICE routine in Stata 11.1, which imputes multiple alternatives based on available data, showed that the association between continued cannabis use and the risk of persistence of psychotic symptoms remained significant (adjusted \( \chi^2 = 16.65, P = 0.001 \)). Again, the strongest association was found for exposure state 3 (that is, cannabis use at both baseline and T2), with an adjusted odds ratio of 2.0 (1.1 to 3.7; P = 0.028).

DISCUSSION
This 10 year follow-up study showed that incident cannabis use significantly increased the risk of incident psychotic experiences. The association was independent of age, sex, socioeconomic status, use of other drugs, urban/rural environment, and childhood trauma; additional adjustment for other psychiatric diagnoses similarly did not change the results. There was no evidence for self medication effects as psychotic experiences did not predict later cannabis use. The results thus help to clarify the temporal association between cannabis use and psychotic experiences by systematically addressing the issue of reverse causality, given that the long follow-up period allowed exclusion of all individuals with pre-existing psychotic experiences or pre-existing cannabis use. In addition, cannabis use was confirmed as an environmental risk factor impacting on the risk of persistence of psychotic experiences (fig 3).
The mechanism behind the association

We investigated the association between cannabis use and risk of psychosis by analysing the expression of psychotic experiences. Psychotic experiences share many characteristics with clinically relevant psychosis, such as demographic, environmental, and genetic risks, and are thought to represent a behavioural marker for psychosis liability. Psychotic experiences are a common and generally transitory phenomenon in the general population, that, nevertheless, might become abnormally persistent and progress to clinical psychotic disorder if combined with exposure to environmental risks. Our study confirmed cannabis as an environmental risk factor, impacting on risk of psychosis by increasing the risk of incident psychotic experiences, and, if use continues over time, increasing the risk of persistent psychotic experiences.

The finding that longer exposure to cannabis was associated with greater risk for persistence of psychotic experiences is in line with an earlier study showing that continued cannabis use over time increases the risk for psychosis in a dose-response fashion. This is also in agreement with the hypothesis that a process of sensitisation might underlie emergence and persistence of psychotic experiences as an indicator of liability to psychotic disorder. Sensitisation refers to the phenomenon that repeated exposure to an environmental stressor leads to progressively greater responses over time. In rats, repeated exposure to THC (delta-9-tetrahydrocannabinol, the main psychoactive component of cannabis) induces behavioural sensitisation: rats pre-treated with increasing doses of THC show greater behavioural (locomotor) responses to a THC challenge after a 14 day washout period than THC naive rats. In humans, however, direct evidence for cannabis sensitisation is lacking. As our study showed that the risk of persistent psychotic experiences increases with longer periods of cannabis exposure, we suggest that a process of sensitisation underlies the association between cannabis and psychosis.

Methodological issues

The results should be interpreted in the light of several limitations. Firstly, information on substance use and psychosis outcome was acquired with the DIA-X/M-CIDI, which essentially provides self reported information. The interview was conducted face to face by clinical psychologists, however, who were allowed to follow up with clinical questioning to ensure systematic and valid assessment of outcomes and can therefore be assumed to yield better and more valid results than a self report questionnaire. Secondly, the analyses were not directly adjusted for the possible confounding effects of a family history of psychosis as this information was not available in the EDSP data. Previous research has shown that associations between cannabis use and psychotic symptoms are not reducible to family history of psychosis and that genetic liability for psychotic disorder does not predict cannabis use. In addition, individuals with a family history of psychosis report more positive symptoms than individuals without such predisposition. As we excluded all individuals with at least one T2 lifetime psychotic symptom from the analysis, the possible confounding effect of family history for psychosis was indirectly adjusted for to a degree. Furthermore, we used a rather broad outcome measure, defined as a minimum of one positive rating on a G section item, representing psychotic experiences rather than clinically relevant psychotic disorder. It has been shown, however, that psychotic experiences show continuity with psychotic disorders such as schizophrenia. In addition, given that fact transient psychotic experiences might, under certain circumstances, become abnormally persistent, giving rise to clinical psychotic disorder, psychotic experiences represent an important phenotype for the investigation of mechanisms and pathways by which environmental risk factors such as cannabis impact on psychosis risk. A further limitation concerns the use of the G section of the DIA-X/M-CIDI. This section was administered at T2 to assess lifetime occurrence of symptoms, which represents a long period for retrospective assessment of psychotic phenomena, possibly resulting in false negative results. As we excluded participants with T2 lifetime experience of psychotic symptoms from the analyses, under-reporting would have resulted in false negative results being incorrectly retained in the analyses. It is unlikely that
Continued use of cannabis might increase the risk for psychotic disorder by impacting on persistence (of normally transitory) psychotic symptoms in young people.

under-reporting would have occurred as a function of cannabis use, which could have resulted in biased estimates. In addition, 23% of participants reported lifetime subclinical psychotic symptoms at T2, which is in keeping with the estimated 15-28% rate of subclinical psychotic symptoms in the general population. Therefore, the influence of under-reporting is probably limited. Finally, as the time between follow-up visits was four years on average, selective recall could have influenced the results. Spurious findings could have arisen if those with psychotic symptoms had better recall of earlier cannabis use. Given the well-known link between psychosis liability and cognitive alterations, including impaired memory, any influence of selective recall would probably have been conservative rather than anti-conservative.

Contributors: H-UW and RL were the principal investigators of the study. RK analysed the data in collaboration with CH, JvO, and MH. RK drafted the paper. All authors contributed to subsequent drafts of the paper and the final version. JvO is guarantor.

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Ethical approval: The EDSP project was approved by the ethics committee of the Medical Faculty of the Technische Universitaet Dresden (No EK-13811).

Data sharing: No additional data available.

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