Cannabis use is associated with an increased risk of psychotic symptoms in individuals from the general population. In patients with an established psychotic disorder, cannabis has a negative impact on illness course, as evidenced by more and earlier relapses, more frequent hospitalisations, and poorer psychosocial functioning. Other epidemiological work, however, has suggested that cannabis may reduce negative and affective symptoms in patients with schizophrenia. Studies using self-report questionnaires to investigate motives that maintain cannabis use among individuals with psychotic disorders indicate that the principal motives for use in this group are enhancement of positive affect, social acceptance and coping with negative affect. Additional evidence that patients may use cannabis to self-medicate distress was reported in a population-based study where vulnerability for psychosis (measured by means of a questionnaire) predicted future cannabis use in those who had never used cannabis before the onset of psychotic symptoms. Two earlier population-based studies, however, had found no evidence for such reverse causality. In order to design successful interventions, epidemiological designs may not be sufficient to provide full insight into the complicated dynamics of cannabis use and its varied effects in patients with psychosis. The current study was therefore designed to further examine, in the context of these complicated dynamics, the association between cannabis use and psychosis using a momentary assessment technique. The aims of the current study were to investigate whether:

(a) frequency of cannabis use in daily life fluctuates as a function of mood and psychotic symptom level (i.e. self-medication effects);
(b) cannabis use is associated with subsequent changes in mood and psychotic symptom level;
(c) patients with a psychotic disorder differ from healthy controls in their sensitivity to the psychosis-inducing effects of cannabis; and

(d) temporal dynamics of cannabis-induced symptoms are apparent (i.e. short- vs. long-lasting effects of cannabis can be discerned).

**Method**

The study sample consisted of 48 patients with a clinical diagnosis of a psychotic disorder and 47 healthy controls. The patients and controls were all frequent cannabis users (current use of at least three times per week). Patients were recruited through in-patient and out-patient mental health service facilities in South Limburg, The Netherlands, and controls were recruited from local ‘coffee shops’ (cafés where cannabis is sold and consumed legally). Participants were provided with a complete description of the study and written informed consent was obtained. The study was carried out in accordance with the World Medical Association’s (WMA’s) Declaration of Helsinki on Ethical Principles for Medical Research Involving Human Subjects, as adopted by the 52nd WMA General Assembly, Edinburgh, October 2000. The study was approved by the standing medical ethics committee of Maastricht University Medical Centre. Interview data were used to complete the Operational Criteria Checklist for Psychotic Illness (OPCRIT), yielding diagnoses according to Research Diagnostic Criteria (RDC) through the OPCRIT computer program for Windows. In addition, current symptomatology was assessed with the Positive and Negative Syndrome Scale (PANSS) and additional information with regard to current psychiatric illness, past and current substance use, and other demographic information was collected. Exclusion criteria were: respiratory, cardiovascular or neurological disease and alcohol use in excess of 5 units per day. Pregnant women were also excluded. A personal or family history of psychosis or use of

**Aims**

To examine the effects of cannabis on psychotic symptoms and mood in patients with psychosis and healthy controls.

**Method**

Patients with a psychotic disorder (n = 42) and healthy controls (n = 38) were followed in their daily lives using a structured time-sampling technique.

**Results**

Daily life cannabis use predicted subsequent increases in positive affect and in patients, but not in controls, decreases in negative affect. In patients, but not in controls, cannabis use predicted increased levels of hallucinatory experiences. Mood-enhancing properties of cannabis were acute, whereas psychosis-inducing effects were sub-acute. There was no direct evidence for self-medication effects in daily life.

**Conclusions**

Patients with psychosis are more sensitive to both the psychosis-inducing and mood-enhancing effects of cannabis. The temporal dissociation between acute rewarding effects and sub-acute toxic influences may be instrumental in explaining the vicious circle of deleterious use in these patients.

**Declaration of interest**

None.
antipsychotic medication was a further exclusion criterion for the control group.

**Experience sampling method**

The experience sampling method (ESM) is a pseudo-random time sampling self-assessment technique. Previous applications of ESM have demonstrated feasibility, validity and reliability in patients with schizophrenia.\textsuperscript{13,14} Participants received a digital wristwatch, and a paper-and-pen ESM booklet. Twelve times a day on six consecutive days, the watch emitted a beep at random moment in each 85-minute time block between 07.30h and 00.30h. After each beep, participants were asked to fill in a self-assessment form, collecting reports of affect, thoughts, severity of symptoms and activity rated on 7-point Likert scales at the moment of the beep. Participants were instructed to complete the form immediately after the beep to minimise memory distortion and to record the time at which they completed their report. During a briefing session, the ESM procedure was explained and a practice booklet was completed and discussed with the participant to confirm that the scale format was accurately understood. During the ESM week, participants were contacted by phone in order to ensure that they complied with the instructions. Participants were requested not to use illicit drugs other than cannabis during the 6 consecutive study days; however, no sanctions were involved. None of the participants was a regular user of drugs other than cannabis. During the telephone contacts, participants were asked about their use of other drugs and one person admitted to having used cocaine on the third day of the ESM week. Reports following this cocaine use were excluded from the analyses. At each beep when forms were completed, participants were asked to report the exact time. Reports completed more than 5 mins before and 15 mins after the beep were excluded from the analyses. This was done because previous research has shown that remote answers are less reliable and less valid than reports at the exact moment of the beep.\textsuperscript{15} Participants with less than 24 valid reports were excluded from the analyses, as previous work has shown that measures of individuals with less than 30% of completed reports are less reliable.\textsuperscript{15}

**Measures**

Measures regarding cannabis use, mood, and psychotic symptoms were derived from the ESM reports as described below.

**Assessment of cannabis use**

Cannabis use, reported after each beep, referred to the period between the previous beep and the current beep (cannabis use, a binary variable). Similarly, cannabis use\textsubscript{previous} referred to cannabis use during the period between the previous beep and the beep before that. Alcohol use, reported after each beep, referred to the period between the previous beep and the current beep (alcohol use, a binary variable).

**Assessment of mood**

Mood states were assessed with 11 mood adjectives rated on a 7-point Likert scale (1 = not at all, 7 = very) reported after each beep, referring to the mood state at that moment (e.g. ‘at this moment I feel anxious’). In previous ESM studies\textsuperscript{1,3,14} with similar populations (patients with psychosis and healthy controls), a positive and a negative mood scale were identified with factor analysis on the raw within-participant scores of the mood items. For the current analyses, therefore, the mean of the adjectives ‘cheerful’, ‘relaxed’, ‘happy’, ‘satisfied’, ‘enthusiastic’, ‘overall good’ formed the positive affect scale (a continuous variable, Cronbach’s $\alpha = 0.89$), and the mean of ‘insensitive’, ‘lonely’, ‘anxious’, ‘blue’, ‘guilty’ formed the negative affect scale (a continuous variable, Cronbach’s $\alpha = 0.80$). Positive affect\textsubscript{previous} and negative affect\textsubscript{previous} referred to mood states reported at the previous beep.

**Assessment of psychosis**

Positive psychotic symptoms were assessed with seven items rated on a 7-point Likert scale (1 = not at all, 7 = very) reported after each beep, referring to the psychotic experiences at that moment. In order to allow self-reporting of psychotic experiences, the items assessing delusional ideation include aspects that can be associated directly with concrete positive psychotic experiences, rather than interpretations of these experiences (e.g. ‘at the moment my thoughts are being controlled by others’). Guided by previous studies\textsuperscript{6,17} the mean of ‘preoccupied thoughts’, ‘racing thoughts’, ‘difficulty expressing thoughts’, ‘thoughts controlled by others’ and ‘suspicious’ formed the delusions scale (a continuous variable, Cronbach’s $\alpha = 0.72$). Hallucinatory experiences were assessed directly (e.g. ‘at the moment I’m hearing voices’) as it has been shown in previous studies that patients can distinguish between hearing real voices and verbal hallucinations.\textsuperscript{18} The mean of ‘seeing things’ and ‘hearing things’ formed the hallucinations scale (a continuous variable, Cronbach’s $\alpha = 0.70$). Delusions\textsubscript{previous} and hallucinations\textsubscript{previous} referred to psychotic symptom severity reported at the previous beep.

**Statistical analyses**

As ESM data have a hierarchical structure with multiple reports (level 1) nested within participants (level 2), multilevel random regression analyses were conducted to account for the variability associated with the two different levels. The odds ratios (for dichotomous variables) and betas (for continuous variables) of these regression analyses are the associations between the independent and dependent variables in the multilevel model. The STATA (version 10 for Windows) multilevel regression XTGEE routine was used for dichotomous variables, and the XTREG routine for continuous variables.

**Overall symptom levels**

Differences between patients and controls in overall symptom level were investigated using group (0 = controls; 1 = patients) as independent and negative affect, positive affect, delusions and hallucinations as dependent variables. To investigate whether patients and controls differed in overall symptom level, multilevel regression analyses were conducted using group as the independent variable, and using negative affect, positive affect, delusions and hallucinations as dependent variables in consecutive models.

**Self-medication effects**

Self-medication effects were examined using negative affect\textsubscript{previous}, positive affect\textsubscript{previous}, delusions\textsubscript{previous} and hallucinations\textsubscript{previous} as independent variables, and cannabis use as the dependent variable (Fig. 1, analysis A).

**Cannabis effects in patients v. controls**

Main effects of cannabis use on subsequent symptom levels were investigated with cannabis use as independent, and negative affect, positive affect, delusions and hallucinations as dependent variables (Fig. 1, analysis B). In order to test the hypothesis that patients and controls differed in their sensitivity to cannabis effects,
multilevel regression analyses were conducted with cannabis use and group as well as their interaction term as independent variables and with negative affect, positive affect, delusions and hallucinations as dependent variables. Regression model: ESM symptom level = β0 + β1(cannabis use) + β2(group) + β3(ESM cannabis use × group). This cannabis × group interaction was then fitted to allow estimation of cannabis effect sizes for both groups separately by calculating the appropriate linear combinations with the LINCOM command in STATA.

Temporal dynamics of cannabis effects

Based on the results of the previous analyses, post-hoc analyses were conducted to further investigate the duration of cannabis effects on mood and positive symptoms. Results

Participants

Of the 95 participants included in the study, 9 controls and 6 patients had fewer than 24 valid reports and were excluded from the analyses. Drop-out was not associated with group (OR = 0.94, 95% CI 0.32–2.73, P = 0.91) or cannabis use during the ESM week (β = 1.01, 95% CI 0.88–1.16, P = 0.90). The final study sample consisted of 80 participants (42 patients and 38 controls; Tables 1 and 2). The OPCRIT diagnoses according to RDC were: schizophrenia (n = 10), schizoaffective disorder (n = 28), and unspecified functional psychosis (n = 4). Groups differed significantly in age, but not in gender (Table 1), or in lifetime or current frequency of cannabis use or lifetime use of other drugs (Table 3).

Overall symptom levels

Overall, patients reported significantly lower levels of positive affect (β = −0.56, 95% CI −1.03 to −0.09, P = 0.019) and higher levels of negative affect than controls (β = 0.41, 95% CI 0.07 to 0.75, P = 0.020). Patients and controls differed significantly in the intensity of hallucinations they reported (β = 0.26, 95% CI 0.17 to 0.70, P = 0.024). Frequency of cannabis use during the ESM week was significantly higher in patients than in controls (OR = 1.17, 95% CI 1.12 to 2.77, P = 0.015).

Self-medication effects

Neither positive affectprevious nor negative affectprevious predicted cannabis use reported at the following beep (OR = 1.08, 95% CI 0.99–1.16, P = 0.09 and OR = 0.97, 95% CI 0.88–1.08, P = 0.64 respectively). Similarly, no association was found between delusionsprevious and subsequent cannabis use (OR = 0.99, 95% CI 0.890–1.06, P = 0.97) or hallucinationsprevious and cannabis use (OR = 0.96, 95% CI 0.83–1.11, P = 0.59).

Cannabis effects on mood in patients v. controls

Cannabis use was associated with subsequent increases in positive affect (β = 0.21, 95% CI 0.14 to 0.29, P < 0.001). Overall, cannabis use had no effect on negative affect (β = −0.04, 95% CI −0.09 to 0.01, P = 0.12). The cannabis use × group interaction, however, was significant for negative affect. Thus, patients were more sensitive to the mood-enhancing effects of cannabis than controls (i.e. large and significant decreases in negative affect were observed after cannabis use in patients, but not in controls; cannabis × group interaction: χ2 = 6.43, d.f. = 1, P = 0.011; Table 4). No such interaction effect was observed for changes in positive affect after cannabis use (χ2 = 0.98, d.f. = 1, P = 0.32).
Cannabis effects on psychosis in patients v. controls

Cannabis use was significantly associated with subsequent increases of hallucinations (β = 0.05, 95% CI 0.01–0.08, P = 0.015). No main effect of cannabis use was found for subsequent delusion intensity (β = −0.02, 95% CI −0.07 to 0.04, P = 0.58). Patients and controls differed in their sensitivity to the hallucinogenic effects of cannabis (cannabis × group interaction χ² = 3.66, d.f. = 1, P = 0.056; Table 4). Patients reported significant increases in hallucinations (β = 0.08, 95% CI 0.03–0.13, P = 0.002), whereas the association between cannabis use and hallucinations was small and non-significant in controls (β = 0.01, 95% CI −0.04 to 0.06, P = 0.75). Further post-hoc analyses for the separate hallucination items showed that the cannabis effects in patients were particularly associated with auditory hallucinations (‘hearing voices’; β = 0.11, 95% CI 0.04–0.17, P = 0.003; Table 4) but that they were less clearly associated with visual hallucinations. No interaction effects were found for delusions (χ² = 1.17, d.f. = 1, P = 0.28; Table 4).

Temporal dynamics of cannabis effects

A follow-up post-hoc analysis was conducted, assessing the duration of cannabis effects on mood and hallucinations in the patient group only, by entering both cannabis use and cannabis useprevious simultaneously in the same model. This suggested that increases in positive affect were observed in the short term (β = 0.31, 95% CI 0.19–0.43, P < 0.001 for cannabis use) rather than the long term (β = 0.07, 95% CI −0.056 to 0.19, P = 0.28 for cannabis useprevious). For hallucinations, however, when both cannabis use and cannabis useprevious were entered simultaneously in the same model, increases in hallucinatory experiences were observed only in the long term (β = 0.09, 95% CI 0.01–0.17, P = 0.030 for cannabis useprevious) and not in the short term (β = 0.06, 95% CI −0.01 to 0.14, P = 0.11 for cannabis use). This suggests that the association between cannabis and hallucinatory experiences was most prominent after a longer period of time compared with its shorter-term mood-enhancing effects.

Discussion

Cannabis use in daily life was associated with subsequent increases in hallucinatory experiences, in particular auditory hallucinations. Patients with a psychotic disorder were more sensitive to the hallucinogenic effects of cannabis than healthy controls. Overall, cannabis enhanced mood, with patients being more sensitive to the positive effects of cannabis on negative affect (i.e. stronger decreases in negative affect after cannabis use). In addition, the data suggest that the positive effects of cannabis on mood are acute, whereas its association with psychotic experiences is sub-acute. Neither negative affect nor hallucinations nor delusions predicted cannabis use, arguing against self-medication effects in daily life. In addition, the effect on subsequent symptom level remained both large and significant after adjustment of symptom level at the previous assessment (i.e. symptoms preceding the use of cannabis).

The current momentary assessment data confirm epidemiological and experimental findings, showing that patients with a psychotic disorder are more sensitive to the psychosis-inducing effects of cannabis than healthy controls. Furthermore, the current findings extend this to the real world of everyday life. The effect sizes were small but the cumulative effect may be considerable.
as exposure to cannabis and subsequent fluctuations in symptoms were observed several times a day over several days. The method of multiple assessments in daily life furthermore has the advantage of avoiding retrospective assessment of symptom states. These are most likely to be distorted by cannabis in the patient group, as increased sensitivity to cannabis operates not only at the symptom level, but also at the level of memory function.20 A crucial point in the interpretation of the results is the validity of the psychosis measures, especially because the results seem to indicate that hallucinations may be a more sensitive phenotype than delusions to study the acute effects of cannabis in daily life. Previous research has shown that patients can distinguish between hearing real voices and verbal hallucinations in self-report questionnaire studies. As such, the interpretation of the results is the validity of the psychosis measures, especially because the results seem to indicate that hallucinations may be a more sensitive phenotype than delusions to study the acute effects of cannabis in daily life.

Table 4 Effects of cannabis use on subsequent symptom levels, patients vs. controls

<table>
<thead>
<tr>
<th>Group</th>
<th>Cannabis −, mean (s.d.)</th>
<th>Cannabis +, mean (s.d.)</th>
<th>Cannabis effect sizea</th>
<th>Group x cannabisb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive affect</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td>4.99 (1.14)</td>
<td>5.12 (1.17)</td>
<td>$\beta = 0.18$, 95% CI 0.07 to 0.28, $P = 0.01$</td>
<td>$\chi^2 = 0.98$, d.f. = 1, $P = 0.32$</td>
</tr>
<tr>
<td>Patients</td>
<td>4.30 (1.26)</td>
<td>4.46 (1.32)</td>
<td>$\beta = 0.24$, 95% CI 0.15 to 0.35, $P &lt; 0.001$</td>
<td></td>
</tr>
<tr>
<td>Negative affect</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td>1.30 (0.65)</td>
<td>1.29 (0.70)</td>
<td>$\beta = 0.03$, 95% CI −0.05 to 0.10, $P = 0.47$</td>
<td>$\chi^2 = 6.43$, d.f. = 1, $P = 0.011$</td>
</tr>
<tr>
<td>Patients</td>
<td>1.96 (1.16)</td>
<td>1.78 (0.96)</td>
<td>$\beta = 0.10$, 95% CI −0.17 to −0.03, $P = 0.0043$</td>
<td></td>
</tr>
<tr>
<td>Delusions</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td>1.87 (0.82)</td>
<td>1.87 (0.89)</td>
<td>$\beta = 0.02$, 95% CI −0.06 to 0.10, $P = 0.70$</td>
<td>$\chi^2 = 1.11$, d.f. = 1, $P = 0.28$</td>
</tr>
<tr>
<td>Patients</td>
<td>2.47 (1.25)</td>
<td>2.45 (1.26)</td>
<td>$\beta = 0.05$, 95% CI −0.12 to 0.03, $P = 0.25$</td>
<td></td>
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<tr>
<td>Hallucinations</td>
<td></td>
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</tr>
<tr>
<td>Controls</td>
<td>1.00 (0.07)</td>
<td>1.00 (0.13)</td>
<td>$\beta = 0.01$, 95% CI −0.04 to 0.06, $P = 0.75$</td>
<td>$\chi^2 = 3.66$, d.f. = 1, $P = 0.056$</td>
</tr>
<tr>
<td>Patients</td>
<td>1.38 (0.88)</td>
<td>1.40 (0.95)</td>
<td>$\beta = 0.08$, 95% CI 0.03 to 0.13, $P = 0.002$</td>
<td></td>
</tr>
<tr>
<td>Auditory hallucinations</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td>1.00 (0.08)</td>
<td>1.01 (0.25)</td>
<td>$\beta = 0.01$, 95% CI −0.06 to 0.08, $P = 0.72$</td>
<td>$\chi^2 = 3.36$, d.f. = 1, $P = 0.067$</td>
</tr>
<tr>
<td>Patients</td>
<td>1.40 (1.03)</td>
<td>1.50 (1.21)</td>
<td>$\beta = 0.11$, 95% CI 0.04 to 0.17, $P = 0.003$</td>
<td></td>
</tr>
</tbody>
</table>

a. Regression coefficient indicates change in symptom score associated with no cannabis use vs. cannabis use, analyses adjusted for age, gender, alcohol use, overall level of cannabis use during the experience sampling method week and symptom level at the previous beep.
b. Chi-squared (d.f. = 1) test for the interaction term, adjusted for age, gender, alcohol use, overall level of cannabis use during the experience sampling method week and symptom level at the previous beep.

Biological plausibility

A reciprocal interaction between the endocannabinoid and dopamine system may explain the psychotogenic effects of cannabis in individuals with increased liability to psychosis as the patient group.27 In the central nervous system, THC (A-9-tetrahydrocannabinol), the psychoactive component of cannabis (binds to cannabinoid 1 (CB1) receptors, the primary receptor site for endocannabinoids. Endocannabinoids act as retrograde signals at central nervous system synapses, by activating presynaptic CB1 receptors.26,27 Activation of CB1 receptors inhibits presynaptic neurotransmitter release. The endocannabinoid system thus plays an important role in the modulation of other neurotransmitters (e.g. gamma-aminobutyric acid and glutamate) and may thereby indirectly influence dopamine firing as well.28 Exogenous cannabinoids such as THC, however, disrupt these subtle, fine-tuning effects of endocannabinoids.29 Animal research has shown that THC evokes burst-firing in the ventral tegmental area and thereby increases dopamine concentrations in striatal

Increased addiction potential in patients

In agreement with two longitudinal studies on this topic,8,9 the present data did not support direct mechanisms of self-medication, since cannabis use was not predicted by previous changes in symptom level or mood. However, cannabis did improve mood, particularly in patients. The combination of differential sensitivity in patients to the acute rewarding effects and the sub-acute negative influences of cannabis on psychotic symptoms (despite the fact that the majority of patients were using antipsychotic medication) may be helpful in explaining the model of cannabis use in patients with psychosis, as proposed by Spencer et al.8 According to this model, use of cannabis is driven by expectations that individuals may have about the (acute) effects of cannabis. The sub-acute negative psychotinic effects may then be experienced as evidence that more use is necessary to bring about the anticipated rewarding effects. Hallucinatory experiences are strongly associated with negative affect, fuelling further use in order to experience acute improvement in mood. Patients may be more sensitive to these mechanisms as they (a) have overall higher levels of negative affect and (b) are more sensitive to the mood- and psychosis-enhancing effects of cannabis. The motivation to enhance affect may consequently direct the individual to future use and cannabis dependence, despite the long-term negative impact cannabis may have on functional outcome.24 The finding that cannabis use was not predicted by changes in mood at the previous beep, could be explained by the fact that in the context of ongoing high levels of negative affect, patients may delay use until they find an appropriate time to smoke (with mood fluctuations occurring further back in the chain of events).
The cannabis effects as presented in this study might thus be attributable to the effects of THC on dopaminergic neurotransmission. Striatal dopamine is thought to play a crucial role in attributing salience to stimuli in the environment. As a consequence, a hyperdopaminergic state may facilitate psychotic experiences by enabling false attribution of significance to ambiguous stimuli. Auditory hallucinations may then be most prominent after exposure to THC as they occur in the moment, whereas delusions may be secondary interpretations of aberrant perceptions. Recently, the acute effects of THC on striatal dopamine were investigated for the first time in human healthy volunteers, using experimental positron emission tomography paradigms. Bossong et al found THC-induced striatal dopamine release, whereas Stokes and colleagues did not find such effects on dopamine transmission. These divergent findings may be due to the fact that individuals differ in their sensitivity to THC, with individuals expressing high psychosis liability, such as patients with schizophrenia, being more vulnerable than healthy controls. Epidemiological work has now identified several factors that co-participate with THC in causing psychosis, as pre-existing psychotic symptoms, exposure to childhood trauma, and a functional polymorphism in the catechol-O-methyltransferase gene have been shown to moderate the effects of cannabis on psychosis outcome. The results of the current study further support the idea that gene–environment interactions underlie the cannabis–psychosis association by showing that individuals at increased risk for psychosis (such as patients) are more sensitive to both the psychosis-inducing and mood-enhancing effects of THC.

**Limitations**

Several limitations need to be taken into account. First, reports on cannabis use and symptoms were based on self-report and were not confirmed by, for example, urine analysis. However, as consumption of cannabis is legal in The Netherlands and 32–51% of the assessments were cannabis moments, underreporting of drug use is unlikely. Experience sampling method adherence research has shown that individuals generally adhere very well with ESM procedures as carried out in the current investigation. Second, the validity of the reports may be challenged by negative effects of cannabis on cognition. The number of drop-outs due to invalid reports, however, was similar to that reported in a recent ESM adherence study. Third, patients were on average 10 years older than controls, thus duration of cannabis use may have been higher in patients than in controls. Adjustment for age did not change the results significantly. However, it may be useful to match on age and duration of cannabis use in future ESM studies. Fourth, it was found recently that patients with a first episode of psychosis show a preference for higher potency cannabis compared with healthy controls. In the current study, however, type and potency of cannabis was not controlled for in the analyses. It is unlikely, however, that this explains the stronger effects of cannabis on psychosis in the patient group. This is because the percentage of THC of the cannabis available in The Netherlands is correlated with its price and on average patients paid lower prices than controls for the cannabis they used during this study (β = −1.40, 95% CI: −2.85 to 0.05, P = 0.058). Nevertheless, in future ESM studies it would be interesting to include information on type and potency of cannabis (e.g. by means of hair samples) especially given the differential effects of THC and cannabidiol on mood and psychotic symptoms. Fifth, participants were regular cannabis users, which raises the question of whether the results would generalise to less frequent use of cannabis. D’Souza et al found frequent users to be blunted to the acute psychotomimetic effects of Δ9-THC. Epidemiological work however, has also shown that the long-term effects of cannabis may increase the risk for psychosis in a dose–response fashion, a finding that may be suggestive of a sensitisation process. More research is needed to investigate the moderating effect of duration of previous exposure on the acute psychosis-inducing effects of Δ9-THC.

**Clinical implications**

In patients, the clinical goal may be to intervene as early as possible to limit progression of the illness to more severe states associated with comorbidity of psychotic illness and cannabis misuse. Cannabis use is frequently a concern in treatment, yet it is often insufficiently discussed between patient and mental health provider. Since the consumption of cannabis and use-related activities play an important role in the everyday social lives of patients, it is of major importance to get better insight into the mechanisms and patterns of use. Some patients who were participating in this study reported that completion of the ESM booklets and the feedback they received afterwards on their patterns of use and related symptoms may have already changed their conceptions about cannabis use to some extent. This suggests that the data presented here may be of great clinical use, being applicable at the individual level as well. A combination of motivational interviewing and cognitive–behavioural therapy (in which not only the positive but also the affective symptoms are addressed) has been proposed to be most effective for patients with psychosis and comorbid substance misuse. The experience sampling data described here validate this idea by emphasising the need to recognise individual differences in sensitivity to cannabis and addiction potential resulting from differences in (genetic) liability to psychosis.
Psychosis reactivity to cannabis use in daily life: an experience sampling study
Cécile Henquet, Jim van Os, Rebecca Kuepper, Philippe Delespaul, Maurice Smits, Joost à Campo and Inez Myin-Germeys
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