Associations Between Cocaine, Amphetamine or Psychedelic Use and Psychotic Symptoms in a Community Sample

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**Objective:** To investigate whether there is an association between use of cocaine, amphetamines, or psychedelics and psychotic symptoms.

**Method:** Cumulated data from a prospective, longitudinal community study of 2588 adolescents and young adults in Munich, Germany were used. Substance use was assessed at baseline, 4-year and 10-year follow-up using the Munich Composite International Diagnostic Interview; psychotic symptoms were assessed at 4-year and 10-year follow-up. Multinominal logistic regression analyses, adjusted for sociodemographic factors, common mental disorders, other substance use, and childhood adversity (adjusted odds ratios, AOR), revealed associations between cocaine, amphetamine or psychedelic use and psychotic symptoms.

**Results:** Lifetime experience of psychotic symptoms was associated with lifetime use of cocaine (AOR 1.94; 95%CI 1.10-3.45), amphetamines (AOR 1.69; 95%CI 0.98-2.93), psychedelics (AOR 2.37; 95%CI 1.20-4.66) and all three substances (AOR 1.95; 95%CI 1.19-3.18).

**Conclusion:** Associations between psychotic symptoms and use of cocaine, amphetamines, and/or psychedelics in adolescents and young adults call for further studies to elucidate risk factors and developmental pathways.

**Keywords:** Substance-Related Disorders, Cocaine, Amphetamine, Psychedelics, Psychoses
Significant Outcomes

• After adjusting for potential confounding factors, use of cocaine, amphetamines or psychedelics was associated with psychotic symptoms.

• Among a subgroup of persons without a mood or anxiety disorder, analysis revealed an association between use of cocaine, amphetamines or psychedelics and psychotic symptoms.

Limitations

• The low positive predictive value of the M-CIDI does not allow for analysis of psychotic disorders; analysis is limited to psychotic symptoms.

• The relatively low prevalence of psychotic symptoms among this community sample limited the ability to analyse for a temporal relationship between substance use and psychotic symptoms.

• Self-report of lifetime substance use and psychotic symptoms may lead to recall errors or erroneous information.
1. Introduction

Although the association of drug use with mood, anxiety and personality disorders has been well documented (1-3), there has been an increased interest in the risk of psychosis among those using illegal drugs. Several clinical and community-based studies have demonstrated an association between cannabis use and psychotic symptoms (4-11). Non-cannabis drug use has been examined among persons with established psychotic disorders (12,13) and psychotic symptoms (14-17). Some of these studies demonstrate an association between use of stimulants and psychotic symptoms (12,15-17). In investigating the temporal relationship between substance (e.g. cannabis, amphetamine, cocaine, psychedelics) use and onset of psychotic symptoms, several studies report that in the majority of study participants, substance use was initiated prior to experiencing psychotic symptoms (17-20). However, there are also findings referring to predominantly secondary onset of substance use after psychotic symptoms (21).

In the present study, we aim to extend this area of inquiry by examining the association between use of three non-cannabinoid illegal drugs – cocaine, amphetamines and psychedelics – and psychotic symptoms. We chose these specific illegal drugs because each of them has neurobiological underpinnings to psychosis. For example, it has long been known that cocaine and amphetamines increase levels of dopamine by powerfully blocking the dopamine transporter (22-24). While psychedelics are primarily serotonin-2A receptor agonists, lysergic acid diethylamide (LSD) in particular is known to have a high affinity for dopamine receptors (25,26). It is not surprising, then, that substance intoxication with these substances includes psychotic symptoms such as paranoia and hallucinations.
Although elevated levels of psychotic symptoms among persons using cocaine or amphetamines have been reported in general population samples (14,16), clinical samples (12,13) and among prison inmates (15), the strength of the association has varied considerably.

There are a number of limitations of studies on non-cannabinoid illegal drug use and psychotic symptoms. First, studies using clinical or forensic samples are limited by selection bias (12,13,15). Second, many studies have not adjusted for confounding variables such as other mental disorders, drug-induced psychosis, and use of cannabis or other drugs, such as opiates, sedatives, or PCP (13,14,16,17).

Using data from a prospective-longitudinal community study of adolescents and young adults, we examine associations between cocaine, amphetamine or psychedelic use and psychotic symptoms. Since psychotic symptoms can occur during the intoxication phase of these non-cannabinoid drugs, we were specifically interested in examining whether these drugs were associated with psychotic symptoms that were not limited to the intoxication periods. We predict that cocaine, amphetamine or psychedelic use will be associated with psychotic symptoms, independent of sociodemographic factors, adverse childhood experiences, and other drug (cannabis, opiates, sedatives, PCP) use, which are factors known to be associated with development of psychotic symptoms (27,28). Due to methodological limitations, we were unable to examine temporal relationships between substance use and psychotic symptoms in this study.

_Aims of the study_

To examine the association between use of cocaine, amphetamine and psychedelics and psychotic symptoms in a large, representative community sample of adolescents and young adults followed over approximately 10 years.
2. Methods

2.1 Sample and overall design

Data were collected as part of the Early Developmental Stages of Psychopathology (EDSP) study. The EDSP is a prospective-longitudinal study on prevalence, incidence, risk factors and course of substance use disorders and mental disorders in a representative community sample of adolescents and young adults (29,30). The baseline investigation was conducted in 1995 with the total sample of 14- to 24-year-olds (N=3,021) and a response rate of 71%. The first follow-up study was conducted only among the younger cohort in 1996/1997 with a response rate of 88% among 14-17 year olds. The second follow-up (“4-year follow-up”) was carried out in 1998/1999 with a response rate of 84% among all 14- to 24-year-olds of the baseline investigation. The third follow-up (“10-year follow-up”) was also conducted among all participants in 2004/2005 and reached a response rate of 73%.

The sample was randomly drawn from government registries in Munich, Germany. All participants provided informed consent after complete description of the study. Because the study was designed as a longitudinal panel with emphasis on the early developmental stages of psychopathology and substance use disorders, 14-15-year-olds were sampled at twice the probability of 16-21-year-olds, and four times the probability of 22-24-year-olds. The community sample has been demonstrated to be representative of the general population (29).

2.2 Diagnostic Assessment

Diagnostic assessments in all waves were based on the computer-assisted Munich-Composite International Diagnostic Interview (M-CIDI/DIA-X, 31) which allows for the assessment of symptoms, syndromes, and diagnoses of 48 mental disorders according to the
DSM-IV criteria (32) and for collection of data on onset, duration, severity, and psychosocial impairment. Diagnostic findings were obtained by using the M-CIDI/DSM-IV algorithms. At baseline, the lifetime version of the M-CIDI was used. At each follow-up, the interval version was applied. In all assessments, the M-CIDI was supplemented by a respondents’ booklet that included scales and questionnaires for assessing psychological constructs relevant to the study (30). Interviews were conducted by trained clinical interviewers (mainly psychologists in postgraduate training) and closely monitored by the staff members and clinical supervisors during weekly supervision sessions. Correct interview administration was also verified by phone-calls to participants. Follow-up interviews were conducted using the same procedures (29,30). Test-retest reliability and validity, which were fair to good (kappa 0.56-0.81 for DSM-IV diagnostic categories), have been reported in detail elsewhere (33,34).

2.2.1 Assessment of substance use and substance use disorders

The methods and previous results on substance use disorders of the study have been published in greater detail elsewhere (1,35-43). Briefly, cocaine, amphetamines, and psychedelics and other illegal substance use (frequency and quantity), as well as DSM-IV abuse and dependence were assessed in Section L (drugs) of the M-CIDI/DIA-X. The section starts with screening questions on prescription drug use followed by questions on use of illegal substances. In the case of an affirmative response on the screening questions, a list of specific substances together with their “street-names” is presented. The list includes cocaine, amphetamines, and psychedelics, and five other classes of illegal substances (e.g. opioids, cannabis) as well as categories for “other” illegal substances and polysubstance use. Following this probe for the type of substance(s) used, questions are posed regarding the frequency and
quantity. If a respondent reports using any illegal substance 5 or more times, questions for each substance of these substances are posed in order to assess symptoms of DSM-IV abuse and dependence. The section is skipped for respondents who would not answer openly to questions about illegal substance use. Inter-rater reliability of the CIDI substance use, abuse and dependence sections is in the acceptable range (kappa 0.55-0.83 for DSM-IV diagnoses of abuse or dependence of a substance) and good agreement was found between clinician-assigned DSM-IV substance use diagnoses and those assigned according to the M-CIDI DSM-IV algorithms (kappa 0.83-0.86 for DSM-IV diagnoses) (33,35). In this study we refer to the lifetime use of cocaine, amphetamines and psychedelics five or more times.

2.2.2 Assessment of psychotic symptoms

Psychotic symptoms were assessed in the Psychosis section (G) of the M-CIDI/DIA-X at 4-year follow-up and 10-year follow-up. At the 4-year follow-up, prior lifetime history of psychotic symptoms was assessed, whereas at 10-year follow-up, symptoms during the interval period were assessed. A positive rating on any of the 15 core psychosis items of the M-CIDI (delusions, hallucinations) was regarded as a presence of a psychotic symptom. Participants were questioned to ensure that none of the positive responses were due to direct effects of medication, drug or alcohol use. The M-CIDI has been demonstrated to have a very low positive predictive value (0.226), a low specificity (0.6), a good negative predictive value (0.973) and sensitivity (0.875) for any specific psychotic disorder, such as schizophrenia (33). For this reason, psychotic symptoms, and not disorders, were assessed. This methodology has been utilized previously in epidemiologic work that has focused on examining the link between cannabis and psychosis (9).
2.3 Statistical analysis

Data were weighted to consider different sampling probabilities as well as systematic non-response at baseline; numbers (N) are reported unweighted. The Stata 10.1 software package was used to calculate proportions and standard errors as well as robust confidence intervals for weighted data (44). Cumulated data from baseline and all follow-up assessments are reported in this study. Since the follow-up CIDI questions refer to the time since the last interview, aggregating the information from the different waves reflects the cumulative lifetime status up to the last completed assessment at 10-year follow-up (either 4-year or 10-year follow-up; the cohort aged 14-17 years at baseline was also assessed at 18 months).

3021 participants completed the baseline assessment. Of those, 2719 completed 4-year or 10-year follow-up, the interviews in which psychotic symptoms were assessed. 131 participants refused to respond openly to the questions about illegal substance use on at least one assessment and were excluded leaving a sample of 2588 participants for our analyses.

To examine associations between the three illegal drug groups and psychotic symptoms, multinomial logistic regression analyses were used, with psychotic symptoms as the outcome and use of cocaine, amphetamines and psychedelics as independent variables, where those using a specific drug five or more times were compared to those using that substance zero to four times. First we report odds ratios (OR) and 95% confidence intervals (95% CI) from unadjusted logistic regression analyses. Second, we report adjusted odds ratios (AOR). Analyses were adjusted in two stages, first, for age, sex, social class, and urbanicity (Adjusted 1), and subsequently further adjusted for alcohol use disorder, nicotine dependence, cannabis use, other drug use disorder, any mood or anxiety disorder, and childhood adversity (Adjusted 2).
Childhood adversity included any qualifying trauma before the age of ten, death of a father or mother before the age of ten, separation or divorce of parents before the age of ten, or not growing up with biological parents for most of the time (all assessed at the baseline interview). Variables such as drug use disorders (2), mood or anxiety disorders (45,46), urbanicity (28) and childhood adversity (27) have been adjusted for in the analysis because they have been demonstrated to be associated with psychotic symptoms.
3. Results

Table 1 provides the cumulative lifetime incidence estimates of cocaine, amphetamine or psychedelic use in the EDSP sample until the 10-year follow-up. Additionally, it shows the distribution of sociodemographic factors and the cumulative incidence of substance use and specified mental disorders. Of the 2588 participants included in our analysis (Table 1), 1319 (number unweighted; percentage weighted: 49.5%) were male, and the mean age at last assessment was 27.1 years. The cumulative incidence of lifetime cocaine use (five times or more) at 10-year follow-up was 5.5%, 6.3% for amphetamines, and 3.2% for psychedelics. The cumulative lifetime incidence for any non-cannabinoid illegal substance (combined group of cocaine, amphetamines or psychedelics) use was 8.5%.

At the last follow-up, the cumulative lifetime incidence of experiencing two or more psychotic symptoms was 218 (8.3%) for the total sample (N = 2588). Among persons having used a substance five or more times, 32 (22.5%) experienced two or more psychotic symptoms for cocaine use, 37 (21.3%) for amphetamine use, 24 (28.2%) for psychedelics, and 49 (20.6%) for any non-cannabinoid illegal substance (combined group of cocaine, amphetamines or psychedelics) (Table 2). The cumulative lifetime incidence of two or more psychotic symptoms did not differ between males (121, 8.9%, 95% CI 7.35-10.73) and females (97, 7.7%, 95% CI 6.27-9.58).

As shown in Table 3, a strong association was found between lifetime use of a substance and two or more psychotic symptoms in comparison to those without any psychotic symptoms (ORs ranging between 3.54-5.12). Although adjusting for covariates decreased the size of the associations, the lifetime use of cocaine, psychedelics and use of any non-cannabinoid illegal substance (combined group of cocaine, amphetamine, psychedelics) remained significantly
associated with lifetime occurrence of two or more psychotic symptoms (AORs between 1.94-2.37). The association between amphetamine use and two or more psychotic symptoms did not remain significant after adjusting for covariates. After adjusting for covariates, the associations between lifetime substance use and one psychotic symptom were not significant for any of the substances examined.

In order to investigate whether the association between psychotic symptoms and substance use was independent of having a mood or anxiety disorder, we additionally performed subgroup analyses with data from all participants who had not met criteria for a mood or anxiety disorder during their lifetime (N = 1224). A strong association was found between lifetime use of a substance five or more times and two or more psychotic symptoms in comparison to those without psychotic symptoms (ORs ranging from 4.28-8.23, Table 4). However, after adjusting for covariates, the association between lifetime substance use and two or more psychotic symptoms was significant only for psychedelics (OR 3.56, 95% CI 1.20-10.61). Within this subgroup, none of the associations between lifetime substance use and one psychotic symptom remained significant after adjusting for covariates.
4. Discussion

The main finding of the present study was that in adolescents and young adults, use of cocaine, amphetamine and/or psychedelics, was associated with psychotic symptoms. These associations were significant when adjusted for variables known to increase risk for psychosis, and most associations also remained after adjustment for use of alcohol, nicotine, and cannabis. Our results extend the previous work (12,15-17) by showing an association between both psychedelics and stimulants (cocaine and amphetamine), and psychotic symptoms among a community-based population while excluding substance-induced psychosis. Among the subgroup of people without a mood or anxiety disorder, the strength of associations between use of a substance and psychotic symptoms was similar for cocaine, amphetamine, and the combined group of substances, however, psychedelics showed a slightly stronger association that remained significant after adjusting for covariates.

4.1 Strengths and Limitations

Strengths of the study include a population-based sample not limited by selection bias associated with treatment-seeking samples, and measurement of symptoms based on a standardised interview conducted by psychologists. Although community studies like the EDSP study data have some advantages in many ways, measurement of the outcomes is limited to psychotic symptoms (30). Despite the remarkable prevalence of 8.5% for psychotic symptoms in this community survey, data could not be analysed for diagnostic outcomes such as schizophrenia, or another psychotic disorder, because the M-CIDI cannot validly assess psychotic disorders according to DSM-IV criteria (33). Being based only on psychotic symptoms the results should not be interpreted as a positive association with psychosis in the
sense of diagnostic categories, as individual symptoms are more likely to occur than symptoms meeting a full set of criteria for a diagnosis of a disorder. The low prevalence of psychotic symptoms also limited our ability to perform a prospective analysis to examine whether a temporal relationship exists between substance use and psychotic symptoms. It should be noted that the methodology used in the present study has been previously utilised and critically discussed in several other examinations on this topic with regard to cannabis disorders (6,9). Additionally, two other limitations have to be noted. First, findings are based on self-reported use patterns and symptoms, which require accurate reports of the amount of a substance as well as accurate reports on symptoms. As such, there is a possibility of recall bias, but the confidential interview method used may help to minimize recall errors and erroneous information. Our study data assessed lifetime rates of substance use in a prospective, longitudinal design, which might account for somewhat elevated rates of substance use in comparison to some other studies, however, other population-based surveys, such as the National Household Survey on Drug Abuse found comparable estimates of substance use disorders. Among 12-29 year olds, 12-month rates of alcohol abuse were 27.4% and 14.5% for men and women, respectively, and 12.1% and 6.6% for alcohol dependence for men and women, respectively (47). Second, this community sample comprises a relatively well-educated and economically stable urban population. Therefore, estimates may not be generalisable to other more demographically diverse populations, and replication of findings in other community-based samples is needed.

4.2 Clinical Relevance/Possible Implications
Several cases of adverse reactions to illegal substances, including death, have been reported, however, many individuals use them without harm, which may engender a false sense of security from adverse outcomes (48). Illegal substance use and substance use disorders occur at a high rate among the population, especially among adolescents and young adults (3,17,38,41) and the co-occurrence between illegal substance use and psychosis is more prevalent than would be predicted by chance alone (20,49). While the exact mechanisms underlying the occurrence of prolonged psychotic symptoms associated with non-cannabinoid substance use are not known, theories have been proposed. There is some data to support a theory that repeated exposure to a stimulant, such as cocaine or amphetamines, leads to changes in the central nervous system, akin to ‘kindling’, implying that stimulant use may cause a psychosis that would not have occurred in the absence of the stimulant (50). There is less research in the area of prolonged psychotic symptoms associated with psychedelic use, however, a model where use of psychedelics might be associated with accelerating or precipitating the onset of an illness that would have ultimately have developed in an individual, has been proposed (51).

Because of conflicting evidence showing that substance use may begin before or after the onset of psychotic symptoms, studies examining the biochemical interactions and possible genetic factors at play will be important in proving or disproving the etiologic theories proposed. Studies examining genetic factors have begun in the areas of cannabis-psychosis association with examination of the catechol-O-methyltransferase gene (52) and methamphetamine-psychosis association with examination of protein interacting with C kinase (PICK1) gene (53) and polymorphisms therein that may modulate clinical outcomes after exposure to those substances. Similar genetic links associated with development of psychosis in individuals using other non-cannabis substances should be investigated, in addition to further elucidating non-biologic
factors such as environmental influences. Ongoing advances in neuroimaging such as diffusion tensor imaging (54) may also prove valuable in identifying structural abnormalities in many psychiatric disorders and possibly abnormalities common to both psychotic and substance use disorders.

In conclusion, findings from this study contribute to a growing body of literature that suggests that use of non-cannabinoid drugs such as cocaine, amphetamines or psychedelics is associated with psychotic symptoms. While neither the causality nor the possible biochemical mechanisms underlying this association have yet been demonstrated, this nonetheless presents a significant public health concern and the potential to utilize public education to reduce drug use and cases of psychosis in young adults should be carefully considered. Nonetheless, there remains a further need for longitudinal data in order to clarify the temporal sequence and whether substance use is, in fact, a causal factor in experiencing psychotic symptoms.
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