The impact of regular ecstasy use on memory function

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ABSTRACT

Aim To assess memory impairment in a group of regular users of ecstasy compared with a group of regular users of cannabis, after accounting for possible confounding factors such as other drug use, premorbid intelligence and psychopathology.

Method Comparative and regression analysis was used to determine the presence or absence of a difference in memory function between 40 regular ecstasy users and 37 regular users of cannabis, who were interviewed at the National Drug and Alcohol Research Centre in Sydney, Australia. Regression analysis was used to find associations between life-time exposure to ecstasy use and memory performance. Memory function was assessed using an age-standardized memory test. Other scales were used to assess premorbid intelligence, physical and psychological health, drug withdrawal and other drug use.

Results Initial comparative analysis showed a trend towards a significantly poorer performance by the regular ecstasy-using group on the ‘auditory immediate memory’ and ‘auditory delayed memory’ indices. When regression analysis was performed an estimate of verbal intelligence was found to be the most predictive of most memory indices including ‘auditory immediate memory’ and ‘auditory delayed memory’. Life-time exposure to ecstasy was not predictive of the memory indices. The current frequency of cannabis use was found to have some predictive effect for immediate and delayed visual memory.

Conclusion This study does not show memory impairment in a group of ecstasy users relative to cannabis using controls. The previously reported association of life-time exposure to ecstasy and memory was not found. The findings may indicate a confounding role of cannabis use, as has been recently reported.

KEYWORDS 3,4-methylenedioxymethamphetamine, cognitive functioning, ecstasy, MDMA, memory.

INTRODUCTION

Although the link between 3,4-methylenedioxymethamphetamine (MDMA) (ecstasy) and serotonergic neurotoxicity is quite clear in animal models, there is less evidence for neurotoxic reactions in humans. A recent paper in the _Lancet_ expressed concern about the risk of neurotoxicity associated with ecstasy use (Boot, McGregor & Hall 2000). Dose-dependent morphological changes found in cultured human serotonergic JAR cells treated with MDMA (Simantov & Tauber 1997) indicate that MDMA has neurotoxic potential in humans. However, discrepancies between dosing regimes in animals and typical patterns of drug use by recreational human users of ecstasy create uncertainties about the effects of MDMA in vivo. Studies of the effects of ecstasy use in humans have focused on three areas. First, indirect measures of serotonergic function in humans, such as decreased CSF levels of 5-hydroxyindoleacetic acid, in recreational ecstasy users compared with ‘ecstasy-naive controls (Peroutka 1987; Ricaurte et al. 1990; McCann et al. 1994; Bolla, McCann & Ricaurte 1998; McCann et al. 1999) have been used to suggest that MDMA-induced serotonergic neurotoxicity may occur in humans. The indirect nature
of these studies limits the conclusions that can be drawn from their findings, but the data are consistent with a serotonergic deficit.

More recently, researchers have used brain imaging to assess the effect of long-term ecstasy use (McCann et al. 1998; Chang et al. 1999; Obrocki et al. 1999; Semple et al. 1999; Chang et al. 2000). Only one of the brain imaging studies suggested that ecstasy use causes neurotoxicity (McCann et al. 1998). The method used by this study, directly assessing serotonergic function using radioligands selective for serotonin transport molecules, was used by another research group which drew a different conclusion (McCann et al. 1998; Semple et al. 1999). Their suggestion of a reversible change in the serotonergic neurones or a downregulation of their action has been supported by other studies (Chang et al. 1999; Chang et al. 2000).

Finally, researchers have sought to characterize the effects that putative serotonergic dysfunction has on the functioning of the individual. A particular focus for this research is to assess the cognitive functioning of abstinent ecstasy users. Compared to users of other drugs and drug-naive controls, ecstasy users have displayed a varied pattern of deficits. These studies are outlined below.

A recent paper has commented on the possible confounding effects of cannabis use in previous research of this nature. Croft et al. (2001) studied drug-naive subjects, cannabis-using subjects and ecstasy- and cannabis-using subjects on a battery of neuropsychological tests. There was no difference between the three groups on an estimate of verbal intelligence; however, the combined group of drug users performed more poorly than the control group on measures of visual recognition, non-spatial associative learning, working memory, verbal memory and manual dexterity. The cannabis-using group and the ecstasy and cannabis group did not differ on any of the tests. In covariate analysis, all deficits apart from speed of processing were shown to be more closely related to cannabis use than ecstasy consumption. Although intelligence was not adjusted for in covariate analysis, these results suggest cannabis as a possible confound.

A study by Bhattachary (2001) assessed 26 regular users of ecstasy, 18 novice users, 16 abstinent users and 20 ecstasy-naive subjects. The researcher found that all of the ecstasy-exposed groups scored significantly lower scores than the drug-naive controls on a test of immediate verbal memory, while the regular ecstasy users and abstinent group differed significantly from the controls on a measure of delayed verbal memory. Regular users and abstinent users also performed significantly worse on a measure of verbal fluency. Life-time consumption of ecstasy and days since last ecstasy use were significant predictors of immediate verbal memory in a regression including cannabis use and IQ. Life-time consumption of ecstasy was also significantly correlated with scores of delayed verbal memory and verbal fluency. Although attempts were made to assess the effect of cannabis use on the results, differing drug use among the groups may confound the analysis.

Gouzoulis-Mayfrank et al. (2000) assessed 28 regular ecstasy users, 28 drug-naive controls and 28 subjects matched for cannabis use with the ecstasy-using group. The ecstasy users performed significantly more poorly on measures of verbal learning and immediate visual memory. The ecstasy users had significantly worse attention scores than the cannabis group, which may have interfered with their performance on the memory assessment. Morgan (1999) found a significant impairment of verbal memory in 25 ecstasy users compared with 22 polydrug using controls, but failed to find significant differences in working memory (Morgan 1998).

Verkes et al. (2001) studied men who were moderate (Parrott et al. 1998) and heavy (Parrott et al. 1998) users of ecstasy and ecstasy-naive (Wareing, Fisk & Murphy 2000). They found that the ecstasy users scored significantly more poorly on tests of reaction time, memory span, word recognition and figure recognition. However, intelligence was not assessed and there was no attempt to account for differences in drug use history. Rodgers (2000) also failed to account for intelligence when finding a significant impairment in delayed memory between a group of regular ecstasy users and a group of regular cannabis users.

Wareing et al. (2000) found various impairments of central executive functioning and information processing speed when controls (Chang et al. 1999) were compared with current (Chang et al. 1999) and abstinent (Chang et al. 1999) users of ecstasy. Intelligence and other drug use were not accounted for in the analyses, and differences in anxiety and arousal were found to diminish group differences in one of the subtests. Assessment of verbal fluency and visual memory did not yield group differences. McCann et al. (1999) similarly found impairments in working memory tasks when 22 ecstasy users were compared with 23 ecstasy-naive controls. There was an inadequate assessment of other drug use and scores for memory tests were not adjusted for intelligence. Similarly, Parrott et al. (1998) did not note other drug use or intelligence of a small group of regular (Chang et al. 1999) and novice (Chang et al. 1999) ecstasy users and 10 drug-naive controls. Thus it is difficult to interpret the findings.

Bolla et al. (1998) found impaired immediate verbal memory and delayed visual memory after controlling for verbal intelligence in a group of ecstasy users, but again failed to adequately account for other drug use.

In previously published research on the cognitive functioning of abstinent ecstasy users, other factors that
may influence the subjects’ performance have often been overlooked. A failure to control adequately for factors such as polydrug use and premorbid intellectual ability make it difficult to interpret the findings of some of these studies. This study was designed to assess the memory of regular ecstasy users, while accounting for factors other than ecstasy use, particularly the concomitant use of cannabis and intelligence.

**METHOD**

**Study design**

An observational study of subjects exposed to ecstasy and to other drugs, using regression analysis, was employed to address the relationship between exposure to ecstasy and memory function.

**Participants**

Eighty-eight subjects participated in the study. Three subjects did not complete the testing, thus their results have not been included. One subject gave a drug history in the interview that differed from the one given during the initial screening, and as a result of this new information the subject was not suitable for inclusion. The fully tested sample of ecstasy users consisted of 47 subjects, all with a history of regular use over 6 months or longer with a minimum frequency of once a month. Subjects were recruited in Sydney through advertisements in local music magazines, as well as flyers posted at the University of New South Wales. The comparison group consisted of 37 regular cannabis users with no history of regular ecstasy use. Cannabis users were considered the best comparison group, because a majority of subjects in the ecstasy-using group were regular users of cannabis around the time of testing (see Table 1), and all subjects in the ecstasy-using group were regularly using some other illicit drug in the months preceding the test. A small amount of ecstasy use was allowed in the comparison group (life-time exposure of up to five tablets), as a greater range of total use of ecstasy would give the best possible sample for regression analysis. It was also considered that robust group differences in memory function would not be masked by a small amount of ecstasy consumption, particularly if the effects on memory were due to neurotoxicity.

All participants were screened in a telephone interview. Exclusion criteria were: (a) reported regular use of any drug other than alcohol or cannabis, which was more frequent than their ecstasy use; (b) use of prescribed psychotropic medication at the time of testing; (c) current or previous diagnosis of an organic brain disorder; (d) a history of head trauma, unconsciousness, fits, convulsions, epileptic seizures or attention deficit hyperactivity disorder; (e) a hearing or visual impairment that prevented completion of the tests; (f) past or current diagnosis with or treatment for a psychiatric illness (except depression) and/or drug or alcohol abuse; (g) current diagnosis with depression or receipt of treatment for depression; or (h) a first language other than English.

**Procedure**

Individuals who met the criteria for participation in the ecstasy-using group or the comparison group were required to abstain from drugs and alcohol for at least 24 hours prior to testing. Those who agreed to this requirement then attended the National Drug and Alcohol Research Centre for testing. All participants signed informed consent forms prior to participation in the study. Testing lasted approximately 2 hours.

A battery of neuropsychological tests, consisting of the Wechsler Memory Scale–III (WMS-III) (Wechsler 1997a), the ‘Vocabulary’ subtest of the Wechsler Adult

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**Table 1** Means (standard deviations) for self-reported drug consumption.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Ecstasy n = 40</th>
<th>Control n = 37</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>39</td>
<td>35</td>
</tr>
<tr>
<td>Average per month (units)</td>
<td>71.4 (91.0)</td>
<td>74.9 (87.7)</td>
</tr>
<tr>
<td>Cannabis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>25</td>
<td>37</td>
</tr>
<tr>
<td>Average per month (joints)</td>
<td>67.9 (285.6)</td>
<td>62.6 (119.2)</td>
</tr>
<tr>
<td>Amphetamine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (g)</td>
<td>0.5 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Cocaine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (g)</td>
<td>0.2 (0.5)</td>
<td>–</td>
</tr>
<tr>
<td>Heroin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (‘hits’)</td>
<td>0.3 (1.2)</td>
<td>–</td>
</tr>
<tr>
<td>LSD</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (‘tabs’)</td>
<td>0.2 (0.6)</td>
<td>–</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (tablets)</td>
<td>0.1 (0.4)</td>
<td>–</td>
</tr>
<tr>
<td>Inhalants</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current regular use</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Average per month (‘hits’)</td>
<td>5.9 (22.4)</td>
<td>–</td>
</tr>
</tbody>
</table>

*a* indicates significant difference at the 0.001 probability level; *b* indicates significant difference at the 0.05 probability level; *c* only amyl nitrate and nitrous oxide used by subjects; *d* indicates number of subjects.
Intelligence Scale–III (WAIS-III) (Wechsler 1997a) and the Kaufman Brief Intelligence Test (K-BIT) (Kaufman & Kaufman 1990) was administered to the subjects. Scores on the WMS-III are calculated to produce eight memory indices: (1) ‘auditory immediate memory’; (2) ‘visual immediate memory’; (3) ‘immediate memory’, a composite of (1) and (2); (4) ‘auditory delayed memory’, which assesses delayed recall of information presented in (1); (5) ‘visual delayed memory’, which assesses delayed recall of information presented in (2); (6) ‘auditory recognition delayed memory’, which assesses the ability of the subject to recognize pairs of words read aloud by the examiner, which constituted part of (1) and (4); (7) ‘general memory’, a composite of (4), (5) and (6); and (8) ‘working memory’.

Other measures were given. The Symptom Check List 90–Revised (SCL-90-R) (Derogatis 1994) was used as a self-report measure of general psychopathology and emotional distress. The physical and psychological health and wellbeing of the subjects was assessed using the SF-36 questionnaire (Ware & Sherbourne 1992).

A thorough drug history was taken from all participants. Life-time history of ecstasy, amphetamine, cannabis and alcohol use was obtained if they had been used regularly. Regular use was defined as at least once a month for a period of 6 months. Patterns of drug and alcohol use were noted using a modified version of Skinner’s Life-time Drinking History (Skinner 1979). Use of cocaine, opiates, benzodiazepines, barbiturates, hallucinogens (LSD and Psilocybin mushrooms) and inhalants was also assessed, taking into account current and heavy regular use.

Current drug dependence was assessed using the Severity of Dependence Scale (SDS) (Gossop et al. 1995) for amphetamine, cannabis, alcohol, ecstasy and cocaine. Participants were also required to complete a self-report questionnaire, which assessed physical and psychological symptoms of withdrawal. This questionnaire was based upon a report on amphetamine dependence (Topp, Mattick & Lovibond 1995) as well as the DSM-IV description of withdrawal from amphetamine or similarly acting sympathomimetic drugs (American Psychiatric Association 1994). Subjects were asked to rate how much they were suffering from a particular symptom at the time of testing using a linear scale numbered 0–4.

During the interview, subjects were required to provide a urine sample that was later analysed for the presence of psychoactive drugs. Urine samples were screened for psychoactive medications as well as illicit psychoactive drugs including amphetamine, MDMA, cocaine and its metabolites, cannabis metabolites and opioids. A positive screen for any substance excluding cannabis, or the self-report of cannabis or alcohol use within 24 hours of the test, were exclusion criteria.

Statistical analysis

Statistical analyses were performed using SPSS for Windows (release 9.0). Group differences in demographic variables, drug use measures, cognitive performance indices and other variables were assessed using independent-sample t-tests. Each of the eight WMS-III memory indices was then included as the dependent variable in eight separate, fully planned multiple linear regression analyses. Age, gender, vocabulary subtest scores, the transformed variable of life-time exposure to ecstasy, the SDS for amphetamines, and the current frequencies of cannabis and cocaine use were the independent variables used.

The regression analyses were used to assess the relationship between life-time exposure to ecstasy and memory. Life-time exposure to ecstasy was the selected ecstasy use parameter for two reasons. First, there was a range of consumption from very low to very high consumption. Secondly, neurotoxic substances such as solvents have been shown to affect the central nervous system in a dose-related fashion (Bleeker et al. 1991). Thus, life-time exposure to ecstasy and cognitive function was of most interest. Severity of dependence on amphetamine is related to cognitive function, while other measures of amphetamine use are not (McKetin & Mattick 1997), thus this measure was used for the regression models. The current frequency of regular use of cannabis and cocaine were thought to indicate most effectively the extent of the pattern of use, with higher frequencies of use more likely to impart a negative influence on memory function. The current frequency of use relates to the frequency of use in the month prior to testing.

RESULTS

Seven subjects in the ecstasy using group were excluded after the urine screen. Six subjects tested positive for psychoactive substances other than cannabis metabolites. One subject failed to give a urine sample and was thus excluded. The final group consisted of 77 subjects: 40 subjects were in the ecstasy-using group and 37 were in the comparison group. Subjects in the two groups were not significantly different in terms of their age, education level and gender ratio. The ecstasy-using group did perform significantly worse on the WAIS-III ‘vocabulary’ subtest \( (P = 0.001) \), although there was no difference between the two groups in intellectual ability as measured by the K-BIT. The means and standard deviations for these variables are presented in Table 2.
Patterns of drug use in the sample

Patterns of ecstasy use are shown in Table 3. All participants in the ecstasy group had a history of regular ecstasy use as described in the method section. Participants in the ecstasy group reported having taken an average of 258 ecstasy tablets; however, it must be noted that one individual reported a lifetime consumption of 3583 tablets. Without this case, the average was 89 ecstasy tablets. The mean duration of use of ecstasy in the sample was 46 months. The average number of ecstasy tablets taken by the comparison group was 1.6.

The members of the ecstasy group were more likely to use other drugs than those in the comparison group. Current regular use of alcohol, cannabis, amphetamine, cocaine, heroin, lysergic acid diethylamide (LSD), benzodiazepines and inhalants are presented in Table 1. Psilocybin mushrooms and barbiturates were not used regularly by any subjects. Comparison subjects reported only current regular cannabis and alcohol consumption, with other drugs not being used regularly, if at all. Apart from cocaine and amphetamine use, other drugs were not used by sufficient numbers of participants to generate a group difference.

Physical and psychological wellbeing of subjects

The two groups were similar in terms of scores for psychological distress as determined by the Global Severity Index of the SCL-90-R and also the psychological and social subsets of the SF-36, namely ‘vitality’, ‘social functioning’, ‘emotional role’ and ‘mental health’. There were differences however, in the physical subsets of the SF-36, with ecstasy users giving significantly worse accounts of their ‘physical functioning’ ($P < 0.05$), ‘physical role’ ($P < 0.01$), ‘bodily pain’ ($P < 0.01$) and ‘general health’ ($P < 0.05$). Table 2 gives means and standard deviations for these scales. Scores for the withdrawal symptoms questionnaire were not significantly different.

Analysis of cognitive functioning

Scores for all of the WMS-III indices were not significantly different between the groups (see Table 4). There was however, a trend towards a significant difference in the ‘auditory immediate memory’ index ($P < 0.07$) and in the ‘auditory delayed memory’ index ($P < 0.08$), but the difference in verbal ability may explain these results (see later).

Twenty-four of the subjects kept for analysis had a positive screen for cannabis metabolites in their urine. The presence of THC in urine could be considered a possible confounding factor in the analysis of memory to follow. Both the regular ecstasy-using group and the comparison group had 12 subjects each who tested positive to cannabis metabolites, and the performance on the memory tests of those with positive urine screens compared with those with clean urine screens was not significantly different. The influence of these positive tests thus appears minimal.

For the linear regressions, the life-time exposure to ecstasy variable was transformed using the equation $\log(x + 1)$ to reduce skewness. This transformed variable was not found to have any significant effect in any of the regression models. The regression model including ‘auditory immediate memory’ as the dependent variable (adjusted $R^2 = 0.243, F_{7,59} = 4.481, P = 0.001$) showed that ‘vocabulary’ was the only significant predictor ($t = 8.860, P < 0.001, \beta = 0.724$). The model using ‘visual immediate memory’ as the dependent variable (adjusted $R^2 = 0.075, F_{5,59} = 1.883, P = 0.086$) was not significant; however, it did show a role of the current fre-
DISCUSSION

In this study, planned analyses were aimed to replicate the findings of previously published studies with the addition of important predictor variables (such as ‘vocabulary’). Initial trends towards significant group differences were most probably explainable by differences in verbal intelligence. The study failed to replicate the memory deficits seen in previous research, and the effect of lifetime exposure to ecstasy on memory function. However, regression analysis showing trends towards significance for the effect of the current frequency of cannabis use is in keeping with the findings of Croft et al. (2001) who found that cannabis use was more instrumental than ecstasy use in generating poorer memory performance. The finding that dependence on amphetamines is related to poorer working memory scores is consistent with previously published data (McKetin & Mattick 1997).

Memory deficits reported in other research were not found here. The disparity in the results of this study and other published research may be explained by differences in design and subject characteristics. Differences in the numbers of ecstasy tablets taken between groups could explain some of the variation of results of this and similar studies. Larger doses of ecstasy may be required for deficits to develop, and continuing exposure to ecstasy in this sample may generate detectable impairments. Secondly, the choice of comparison group for this study differed from previous studies. The inclusion of novice ecstasy users (having used five doses or less) could have hidden group differences. Only a small number of doses of ecstasy may be required to cause a deficit in memory, and thus the comparison group may be impaired in this way. This argument would not be useful to support an effect of neurotoxicity on memory performance since increasing dose would be expected to cause increasing neurotoxic damage, and thus diminished functional capacity. Robust evidence for memory impairment, if present, should still be seen between the two groups in this study. Finally, the average time since the last use of ecstasy was 18 days, with a range of 3–180 days. Residual effects of the drug may have influenced memory performance. However, it would be expected that more recent ecstasy use would impart a negative influence on memory function, and thus would accentuate group differences. Therefore, it may not be considered to be an important confound in this instance.

In summary, ecstasy use in this study was not associated with detectable memory deficits. Evidence for the possible roles of differences in verbal intelligence and cannabis use in generating relative memory impairments is provided by this study.

REFERENCES


