Effects of Opioid Dependence on Visuospatial Memory and Its Associations With Depression and Anxiety

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Introduction: The cognitive impact of opioid dependence is rarely measured systematically in everyday clinical practice even though both patients and clinicians accept that cognitive symptoms often occur in the opioid-dependent population. There are only a few publications which utilized computerized neuropsychological tests to assess possible impairments of visuospatial memory in opioid-dependent individuals either receiving opioid replacement therapy (ORT) or during subsequent short-term abstinence and the effects of anxiety and depression.

Methods: We assessed a cohort of 102 participants, comprising i) a stable opioid-dependent group receiving methadone maintenance treatment (MMT) (n = 22), ii) a stable opioid-dependent group receiving buprenorphine (BMT) (n = 20), iii) a current abstinent but previously opioid-dependent group (ABS) (n = 8), and iv) a control group who have never been dependent on opioids. The Cambridge Neuropsychological Automated Test Battery (CANTAB) neuropsychological tasks undertaken by participants included: Delayed Matching to Sample (DMS), Pattern Recognition Memory (PRM), Spatial Recognition Memory (SRM), and Paired Associate Learning (PAL) tasks. Three clinical measures were used to assess the severity of anxiety and depressive illness: Hospital Anxiety Scale-Hospital Anxiety Depression (HADA)-(HADD), Beck Depression Inventory (BDI), and Inventory of Depressive Symptomatology (self-report) (ISD-SR).

Results: The methadone- and buprenorphine-treated groups showed significant impairments (p < 0.001) in visuospatial memory tasks but not the abstinent group. Impairments in visuospatial memory strongly correlated with higher mood and anxiety symptom severity scores (p < 0.001).

Discussion: These results are broadly consistent with previous studies. Uniquely, though, here we report a strong relationship between visuospatial memory and depression and anxiety scores, which might suggest common illness mechanisms.

Keywords: memory, opioid dependence, heroin, methadone, buprenorphine, depression, anxiety
INTRODUCTION

Substance misuse is a chronic condition often characterized by remissions and relapses (1). Individuals with a history of long-term opioid dependence may demonstrate cognitive impairments, primarily within the executive functioning domains (2–8).

These impairments have been linked to grey matter reductions in the prefrontal cortex, anterior mid-cingulate cortex, and basal ganglia (9), brain regions thought responsible for the regulation of cravings, pain, and emotional experience. In addition, other studies have reported how opioids affect memory, learning, and emotional disturbances (2, 3, 10, 11). Depression has long been associated with widespread cognitive deficits (12) which tend to worsen over a life span (13).

Specific memory tasks have shown to be sensitive and useful in detecting brain dysfunction in the temporal and amygdalo-hippocampal regions (14), which are consistently reported as functionally abnormal in mood disorders and sensation-seeking behaviors (15–17).

Importantly, these brain regions are also relevant to the neurobiology of substance misuse (18) with similar symptoms such as mood, anhedonia, and anxiety associated with drug dependence (19). These symptoms may represent a risk factor for the development of dependence and also may constitute a specific factor by which dependence is maintained, as well as strongly associated with major depressive disorder (MDD). However, depressive and anxiety symptoms have rarely been investigated in opioid dependence within a clinical environment.

Previous studies showed impairments in episodic memory (20), visual memory, verbal memory, information processing, problem solving (21), and spatial, tactile, and verbal memory (2) in heroin-, morphine-, and methadone-dependent participants. Curran and colleagues showed that a single dose of methadone could negatively impact on episodic memory in opiate users (20).

Previously, we have shown that visuospatial memory was impaired in chronic heroin and methadone-dependent participants, those maintained on methadone as part of opioid replacement therapy (ORT), or patients prescribed opioids for chronic pain (10). However, to our knowledge, there are no previous studies reporting the impact of opioid dependence on memory during short-term abstinence from opioids.

Here, we tested the following hypotheses:
(i) Visuospatial memory impairments are associated with current opioid exposure. Conversely, we therefore predicted that abstinence would be associated with no significant impairments.
(ii) Cognitive impairments would correlate with mood and anxiety ratings. Specifically, we predicted that participants with higher depression and anxiety symptoms would have greater visuospatial memory impairments.

METHODS

Study approval was granted by the East of Scotland Research Ethics Committee (REC reference number: 06/S1401/32) and written informed consent obtained from all participants. National Health Service (NHS) Scotland Research Governance approval was provided by the NHS Fife Research and Development Department.

A total of 102 participants were opportunistically enrolled in this study with four groups: (i) a stable opioid-dependent group receiving methadone maintenance treatment (MMT) (n = 22), (ii) a stable opioid-dependent group receiving buprenorphine (BMT) (n = 20), (iii) a current abstinent but previously opioid-dependent group (ABS) (n = 8), and (iv) controls, with no history of illicit heroin, methadone, or buprenorphine use (n = 52). Patients had a diagnosis of Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition (DSM-IV), Opioid Dependence and a history of poly-substance misuse with heroin as the primary “drug of choice” preceding initiation of MMT.

An extensive detailed screening was assessed by two clinicians (A.B. or F.D.), which included sociodemographic information collection and a semi-structured interview to obtain detailed previous histories of drug and alcohol use and current opioid dependence status (Table 1 and Supplementary Table 1). Clinical histories and diagnoses were obtained using the structured Mini International Neuropsychiatric Interview (MINI Plus v 5.0) (22) together with a detailed review of individual clinical care records. The latter included recording the dose of methadone and buprenorphine that each participant received at the time of testing. A morphine equivalent calculation was performed in accordance to a previous publication by Vieweg et al. (23). Each methadone dose was multiplied by 20, and each buprenorphine dose was multiplied by 12 (23). Ongoing abstinence from illicit drug use was also objectively confirmed just prior to scanning with a urine drug test (24) using automated enzyme-mediated immunoassay to classify any detected drug (25). The Clinical Opioid Withdrawal Scale (COWS) was used to quantify the level of opioid withdrawal if present (26). Previous care records from Addiction Services, psychiatric notes, and general practitioners’ records confirmed the absence of hepatitis B and C and HIV. Other exclusion criteria included: past or current histories of psychotic disorders; post-traumatic stress disorder (PTSD); antisocial and borderline personality disorders; neurological and neurodevelopmental disorders; significant head injury; confirmed history of non-fatal overdose episodes; and co-occurring benzodiazepine, stimulant, and/or alcohol dependence.

Current and premorbid intelligence was estimated using the Wechsler Abbreviated Scale of Intelligence (WASI) and National Adult Reading Test (NART) (27, 28).

Visuospatial Memory Tasks

The Cambridge Neuropsychological Automated Test Battery (CANTAB, www.camcog.com) comprises a series of computerized memory tasks (29). As previously reported, the following tasks have shown specificity to detect impairments in visual memory performance [Delayed Matching to Sample (DMS), Pattern Recognition Memory (PRM), Spatial Recognition
TABLE 1 | Demographic, clinical, and substance use history data.

|                        | MMT (N = 22) | BMT (N = 20) | ABS (N = 8) | HC (N = 52) | Statistics |
|------------------------|-------------|--------------|-------------|-------------|------------|
| Number                 | 22          | 20           | 8           | 51          |            |
| Age in years           | 33.6        | 37.4         | 37.6        | 28.0        | P < 0.001  |
| NART                   | 114.3 (5.2) | 98.0 (13.5)  | 106.4 (15.6)| 117.5 (6)  | P < 0.001  |
| HADA                   | 6.0 (4.3)   | 4.8 (2.7)    | 4.0 (2.3)   | 3.5 (3.4)   | P = 0.04   |
| HADD                   | 4.4 (3.5)   | 4.4 (2.9)    | 8.0 (1.5)   | 1.2 (2.3)   | P < 0.001  |
| BDI                    | 12.4 (10)   | 9.9 (6.3)    | 9.0 (1.8)   | 3.7 (5.2)   | P = 0.02   |
| IDS-SR                 | 17.8 (12)   | 12.6 (6.6)   | 14.0 (3.2)  | 7.9 (7.3)   | P < 0.001  |
| Fagerstrom (total score)| 3.4 (2.3)   | 3.9 (2.3)    | 3.5 (2.8)   | ns          |            |
| OD (methadone or buprenorphine in mg) | 73.4 (60.8) | 11.0 (6.7) | –          | –          | P < 0.001  |
| Daily intake expressed as morphine equivalent dose in mg | 1,835.5 (1,277) | 888.0 (533) | –          | –          | P < 0.001  |
| Age when first used heroin in years | 20.2 (4.4) | 21.7 (5.4) | 20.0 (4.7) | –          | ns        |
| Age when dependent on opioids in years | 20.2 (4.4) | 23.6 (5.9) | 22.9 (8.5) | –          | ns        |
| Age when injecting opioids in years | 21.8 (4.2) | 24.8 (6) | 22.7 (6.9) | –          | ns        |
| Years of opioid use    | 12.9 (4.4)  | 13.4 (6.7)   | 13.4 (7.6)  | –          | ns        |
| Age when first used benzodiazepine in years | 17.2 (5.8) | 21.7 (7.7) | 15.6 (6.6) | P < 0.04   |
| Days of benzodiazepine use in the last 30 days | –          | –          | –          | –          | –          |
| Age when first used cocaine in years | 17.3 (1)   | 21.9 (6.6)  | 18.3 (4.2)  | –          | ns        |
| Days of cocaine use in last 30 days | –          | –          | –          | –          | –          |
| Age when first used cannabis in years | 13.3 (3.8) | 15.8 (5.3) | 13.1 (1.2) | –          | ns        |
| Days of cannabis use in last 30 days | –          | –          | –          | –          | –          |
| Age when first used alcohol in years | 10.5 (7.9) | 15.1 (3) | 13.0 (1.9) | –          | 0.04      |
| Days of alcohol use in last 30 days | –          | –          | –          | –          | –          |
| Duration abstinence (days) | –          | –          | 102.2 (61.3)| –          | –          |

Values are mean (SD); MMT, methadone maintenance treatment group; BMT, buprenorphine maintenance treatment group; ABS, abstinent group; HC, healthy control group; N, total number; HADA, Hospital Anxiety Scale; HADD, Hospital Anxiety Depression; BDI, Beck Depression Inventory; IDS-SR, Inventory of Depressive Symptomatology (self-report); NART, National Adult Reading Test; significance * = p = 0.05, *** = p < 0.001; ns, non-significant; mg, milligrams; OD, opioid dose (methadone or buprenorphine).

Memory (SRM), and Paired Associate Learning (PAL)] and spatial memory performance [Spatial Span Task (SSP) and Spatial Working Memory (SWM)] (10).

Depression and Anxiety Rating Scales

Three clinical measures were used to assess the severity of anxiety and depressive illness: the Hospital Anxiety and Depression Scale (HADS) (30), Beck Depression Inventory (BDI) (23), and Inventory of Depressive Symptomatology (IDS): clinician (IDS-C) and self-report (IDS-SR) (31).

HADS is commonly used to determine depression and anxiety. It is a 14-item scale with 7 items that relate to depression (HADD) and 7 items to anxiety (HADA) (30). BDI and IDS are self-report inventories, and they have been mostly used to assess depression and anhedonia (32, 33). BDI demonstrated high internal consistency, with an alpha coefficient of 0.82 (34). Similarly, IDS demonstrated strong internal consistency, with an alpha coefficient of 0.88 (35).

Statistical Analysis

Data meeting assumptions of normality and homogeneity of variance were analyzed using analysis of variance (ANOVA) (36). All other data were compared using Mann–Whitney test. Preliminary analysis of all the experimental and control groups separately indicated that the samples did not come from normally distributed populations with the same standard deviation. We used a post hoc Bonferroni correction in order to control for family-wise error for unplanned tests. Mann–Whitney U tests established that NART, age, and smoking history needed to be used as covariates for hypothesis testing.

A general linear model was performed with “groups” as a factor and “visuospatial memory task performances” as dependent variables using analysis of covariance (ANCOVA). To explore the potential contribution of the impact of depression and anxiety scores on memory task performance, we added an additional correlational analysis within the ANCOVA.

Data were analyzed using the Statistical Package for the Social Science (SPSS) version 24 (SPSS Inc.) in Windows 10 on a PC computer. P values < 0.05 were considered significant.

RESULTS

Demographic Characteristics

Demographic and clinical characteristics are presented in Table 1. Participants and controls were matched on the basis of gender (all males). The MMT, BMT, and ABS groups were older
than the healthy controls (HCs) \(p < 0.001\). The HC group had higher estimated premorbid IQ \(p < 0.001\) according to the NART than the BMT and ABS groups. The mean morphine equivalent daily dose for the MMT group was significantly higher than the BMT \(p < 0.001\). Urine analyses confirmed complete absence of recent heroin, amphetamine, benzodiazepine, and cocaine prior to neuropsychological testing. The MMT group reported they first drank alcohol and consumed benzodiazepine approximately 4.5 years prior to the BMT cohort \(p < 0.04\).

### Visual Memory

#### Performance on DMS

There was a significant effect of group on the percentage of correct responses for DMS \(F(4, 78) = 7.5, p < 0.001\). *Post hoc* Bonferroni comparisons showed that participants from the MMT and BMT groups made significantly more errors than the ABS and HC groups \(p = 0.03\) and \(p < 0.001\), respectively. There was a significant effect of group on the percentage of correct responses for DMS \(F(4, 78) = 7.4, p < 0.001\). *Post hoc* Bonferroni comparisons showed that participants from the MMT and BMT groups made significantly more errors than the ABS and HC groups \(p = 0.02\) and \(p < .001\), respectively.

More details are reported in Table 2 and Figure 1.

#### Performance on PRM, SRM, and PAL

There was a significant effect of group on the percentage of correct responses for the PRM task \(F(4, 60) = 9.3, p < 0.001\) and on the mean correct latency for the SRM task \(F(4, 60) = 6.4, p < 0.001\). Similarly, there was a significant effect of group on the total adjusted errors on the PAL task \(F(4, 75) = 6.1, p < 0.001\) and on PAL first trial memory \(F(4, 75) = 5.7, p < 0.001\) (see Figure 2).

### Spatial Memory

#### Performance on SSP and SWM

There was a significant effect of group on the SSP task (span length) \(F(4, 75) = 10.5, p < 0.001\). The BMT and ABS groups (a) made significantly more errors (between errors) \(F(4, 75) = 5, p < 0.003\) and (b) presented with a poorer strategy on the SWM task \(F(4, 75) = 9.8, p < 0.001\).

### Depression and Anxiety and Visuospatial Memory Performance

Higher HADA anxiety, BDI, and IDS-SR depression scores were significantly correlated with PAL (total error adjusted \(r (66) = 0.3, p = 0.01\), \(r (66) = 0.25, p = 0.04\), \(r (64) = 0.3, p = 0.005\), respectively). Similarly, higher HADA, BDI, and IDS-SR scores were significantly associated with PAL (first trial memory score) \(r (66) = 0.3, p = 0.007\), \(r (66) = 0.28, p = 0.02\), \(r (64) = 0.4, p = 0.001\), respectively). DMS (% correct) significantly correlated with BDI \(r (66) = 0.3, p = 0.01\) (see Table 3).

### Table 2 | Summary of neuropsychological findings for visual and spatial memory.

| Memory and learning measures | MMT (N = 22) | BMT (N = 20) | ABS (N = 8) | HC (N = 52) | Statistics |
|------------------------------|--------------|--------------|-------------|-------------|------------|
| **Visual Memory**            |              |              |             |             |            |
| DMS                           | 84.5 (11.6)  | 80.0 (15)    | 92.8 (2.1)  | 92.5 (5.9)  | \(p < 0.001\), MMT, BMT < ABS, HC*** |
| % correct                     | 80.2 (14.8)  | 75.6 (18.5)  | 91.6 (3.9)  | 90.7 (7.6)  | \(p < 0.001\), MMT, BMT < ABS, HC*** |
| DMS                           | 83.8 (10.1)  | 80.1 (11.7)  | 90.2 (0.9)  | 93.2 (4.3)  | \(p < 0.001\), MMT, BMT < ABS, HC*** |
| % correct (all delays)        | 1,997 (377)  | 2,743 (1,138)| 2,150 (454)| 1,882 (555)| \(p = 0.001\), BMT > HC*** |
| PRM                           | 125.7 (101)  | 29.9 (34.6)  | 11.0 (9)    | 57.0 (90)   | \(p = 0.001\), MMT, BMT > ABS, HC*** |
| % correct                     | 8.5 (0.8)    | 17.9 (4.5)   | 19.7 (3)    | 16.4 (9)    | \(p = 0.001\), MMT < HC, ABS*** |
| **Spatial Memory**            |              |              |             |             |            |
| SWM                           | 8.8 (15.9)   | 33.4 (21.4)  | 22.7 (16.2) | 16.6 (21.9) | \(p = 0.003\), BMT > MMT, HC*** |
| between errors                | 13.1 (14.9)  | 32.9 (6.9)   | 31.7 (6)    | 21.3 (13.4) | \(p = 0.001\), MMT < BMT, ABS*** |

* Values are mean (SD); significance *** = \(p < 0.001\); DMS, Delayed Matching to Sample; PRM, Pattern Recognition Memory; SRM, Spatial Recognition Memory; PAL, Paired Associate Learning; SWM, Spatial Working Memory; N, total number.
DISCUSSION

In this clinically well-characterized study, we have demonstrated that memory for visually presented patterns and spatial locations was impaired in individuals on ORT. This is consistent with previous studies utilizing computerized CANTAB assessment with individuals on ORT and HCs. These studies revealed that individuals on ORT exhibited impairments in comparison to controls on the PRM task (2, 3) and on the PAL task (21). In a recent meta-analysis by Baldacchino and colleagues (37), short-term memory impairments were not present in the abstinence cohorts. This is consistent with our present results, as cognitive impairments were not present in the abstinent group for both visual and spatial memory tasks.

We previously reported that cognitive processes particularly associated with the prefrontal cortex are disrupted during chronic opioid use but not during abstinence (9). Our results could be explained by frontal lobe dysfunction (9, 38–40), which can potentially cause impairments on tasks requiring optimal memory function with patients receiving ORT. In addition, the identified impairments within the opioid-dependent groups on ORT point to specific correlations with depression and anxiety, particularly with tasks sensitive to the anatomical location of the medial temporal lobe.

This is consistent with numerous studies in healthy volunteers identifying the medial temporal lobe, such as the hippocampus and amygdala, as the area where memory-sensitive tasks are encoded (41, 42). Of specific interest, the medial temporal lobe regions have been reported 1) as structurally abnormal in depressive disorder (16) and 2) as one of the main putative candidate regions for both the development and the maintenance of dependence (18) and depression (43).

Regarding possible limitations of the present study, we recruited only males, so these findings shouldn’t be generalized to females (44). Drug use and clinical histories were collected based upon self-report, and no blood, hair, or saliva samples were available to confirm the accuracy of the information given; however, our study did acquire urine drug screen analysis to confirm the absence of recent illicit drug use prior to every session. Additionally, the present study recruited well-matched subjects with regard to their previous drug history in the experimental groups and excluded regular and dependent users of most psychoactive substances, such as alcohol and benzodiazepines, as they have been shown to profoundly impact neuropsychological performance (18). We couldn’t control the effect of nicotine, which may have influenced our results due to its known neuropsychological effects on visual and spatial memory (45). The buprenorphine group had a significant lower morphine equivalent dose than the methadone group, which may impact our findings; however, no statistically significant correlations were present. Larger studies with long-term abstinence are required to fully validate the observed reversibility and possible extinction of these impairments.

Clinical Relevance

Patients’ questions about the effects of opioid dependence on memory and its impact during abstinence cannot comprehensively be answered, due to a current lack of research in this area (10). More data are required on the consequences of opioid dependence on memory in order to evaluate the acceptability of differential treatments, such as methadone and buprenorphine, and perhaps maximize abstinence periods (46). Previous studies have indicated the importance of detecting memory impairments using highly structured and extensive neuropsychological batteries. This is further highlighted in the present study, indicating that opioid-dependent individuals have memory loss in both visual and spatial domains. Early identification of memory impairments associated with opioid dependence could improve the current standard clinical method of assessment. Elucidating the cognitive and neural mechanisms responsible for the formation and maintenance of opioid-related associative dependence has the

![FIGURE 1](A) Delayed Matching to Sample (DMS) task (% correct) box plots: the stable opioid-dependent group receiving methadone maintenance treatment (MMT) and that receiving buprenorphine (BMT) made significantly more errors than the abstinent but previously opioid-dependent group (ABS) and healthy controls (HCs) (p < 0.001) groups. (B) Pattern Recognition Memory (PRM) task (% correct) box plots: the MMT and BMT made significantly more errors than the ABS and HC (p < 0.001) groups.
potential for opening up new therapeutic trajectories during both the prevention and/or reversal of the significant effects on memory and learning, which may be a vulnerability for development and maintenance of opioid dependence. Notably, our results highlight the possibility that opioid-dependent individuals may benefit from focused treatments for depression and anxiety symptoms during ORT.

In particular, understanding the underlying neurocognitive and brain substrates linked to a dual close relationship between comorbid substance misuse and mood states may (a) reveal potential new interventions for the treatment of protracted opioid dependence and/or relapse (18) and (b) provide the required biomarkers to create predictive algorithms to detect early dependence and abstinence (6, 7).

**CONCLUSION**

In summary, our results found that opioid-dependent participants exhibited visuospatial memory impairments closely associated with depression and anxiety scores. These impairments were not present in short-term abstinence, suggesting reversible impairments. Further studies need to explore the effect that mood plays in cognitive impairments observed in this and other dependent populations (e.g. nicotine and alcohol). Indeed, identifying and characterizing the visuospatial memory abilities and their potential mechanisms of action may be of crucial importance in identifying potential common mechanisms controlling the switch from the non-dependent to substance-dependent states and ultimately achieving abstinence in the opioid-dependent population.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study are available on request to the corresponding author.

**ETHICS STATEMENT**

Study approval was granted by the East of Scotland Research Ethics Committee (REC reference number: 06/S1401/32) and written informed consent obtained from all participants. National Health Service (NHS) Scotland Research Governance approval was provided by the NHS Fife Research and Development Department.

**AUTHOR CONTRIBUTIONS**

ST wrote the first draft of the manuscript with AB’s input and created the figures and tables. FD and JS provided revisions to versions of the draft manuscript. ST formatted the manuscript for publication.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpsyt.2019.00743/full#supplementary-material
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