



Increased ventral striatal BOLD activity during non-drug reward anticipation in cannabis users

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ABSTRACT

Despite an increased understanding of the pharmacology and long-term cognitive effects of cannabis in humans, there has been no research to date examining its chronic effects upon reward processing in the brain. Motivational theories regarding long-term drug use posit contrasting predictions with respect to how drug users are likely to process non-drug incentives. The reward deficiency syndrome (RDS) of addiction posits that there are deficits in dopamine (DA) motivational circuitry for non-drug rewards, such that only drugs of abuse are capable of normalizing DA in the ventral striatum (VS). Alternatively, the opponent process theory (OPT) holds that in individuals prone to drug use, there exists some form of mesolimbic hyperactivity, in which there is a bias towards reward-centred behaviour concomitant with impulsivity. The current study examined BOLD responses during reward and loss anticipation and their outcome deliveries in 14 chronic cannabis users and 14 drug-naïve controls during a monetary incentive delay (MID) task. Despite no significant behavioural differences between the two groups, cannabis users had significantly more right VS BOLD activity during reward anticipation. Correlation analyses demonstrated that this right VS BOLD response was significantly correlated with life-time use and reported life-time cannabis joints consumed. No correlations between cannabis abstinence and BOLD responses were observed. We also observed a number of group differences following outcome deliveries, most notably hypoactivity in the left insula cortex in response to loss and loss avoidance outcome notifications in the cannabis group. These results may suggest hypersensitivity during instrumental response anticipation for non-drug rewards and a hyposensitivity to loss outcomes in chronic cannabis users; the implications of which are discussed with respect to the potentially sensitizing effects of cannabis for other rewards.

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Introduction

The long-term use of cannabis has been linked to deficits in learning, memory and executive functioning in humans (Nestor et al., 2008; Grant et al., 2003; Solowij et al., 2002; Bolla et al., 2002). Critically it has been proposed that compromised cognitive processing in chronic cannabis users may promote continued drug consumption, which may be exacerbated following drug withdrawal (Goldstein and Volkow, 2002; Volkow et al., 2002). Subjective reports concerning the effects of cannabis suggest that its reinforcing properties, like other drugs of abuse, are related to its effects on the ventral striatal (VS) “reward circuitry” (Huestis et al., 2007; D’Souza et al., 2008; Hunault et al., 2008), where inhaled cannabis has been shown to induce dopamine (DA) release (Bossong et al., 2009). Despite an increased understanding of the cognitive and psychopharmacological effects of

cannabis, little is known regarding its chronic effects upon reward processing in the human brain.

Despite significant evidence of VS responses to drug-associated stimuli in substance dependence (London et al., 1999; Grusser et al., 2004; Sinha and Li, 2007), there is little evidence that drug users are abnormal in their responses to stimuli that predict non-drug rewards. Motivational theories regarding drug use make contrasting predictions with respect to how drug users may differentially recruit the VS in response to cues that signal non-drug incentives (Bjork et al., 2008). The reward deficiency syndrome (RDS) and the allostatic hypotheses (AH), for example, view addiction as a deficit in DA motivational circuitry for non-drug rewards, such that only drugs of abuse are able to normalize DA at the VS (Blum et al., 2000; Koob et al., 2004). The incentive salience hypothesis (ISH) attributes compulsive drug use to alterations in striatal functioning, in which drug cues reputedly acquire increased incentive-motivational value (Berridge and Robinson, 1998; Robinson and Berridge, 2001).

Substance-dependent patients have also been shown to exhibit both impulsive and reward-centred choice behaviour under laboratory conditions, with cocaine and alcohol dependence associated with

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an increased preference for small immediate over larger delayed rewards (Bickel and Marsch, 2001; Bechara et al., 2001; Heil et al., 2006; Bjork et al., 2004). This may suggest that in individuals who are both prone to, and engage in chronic drug use, there exists some combination of both mesolimbic reward hyperactivity and hypoactive frontocortical punishment-avoidance circuitry (Solomon and Corbit, 1973; Bechara et al., 2005; Bickel et al., 2007). Therefore, drugs such as cannabis, capable of engaging the VS, as described above, may induce a heightened and indiscriminate response to cues which signal all forms of potential reward, an effect which is related to the magnitude of life-time drug use.

Given that cannabis increases DA in the VS (Bossong et al., 2009) and that the anticipation of non-drug incentives reliably activates (Schultz et al., 1997; Knutson et al., 2001; O'Doherty 2004) and increases DA release in this region (Schott et al., 2008) the current investigation hypothesised that 1) chronic cannabis use would be associated with altered VS functioning during reward anticipation, together with differential neural activity in limbic and paralimbic regions during outcome delivery and 2) that VS, limbic and paralimbic activity would be related to life-time cannabis exposure, but not abstinence.

Material and methods

Participants

Fourteen cannabis users and 14 controls were recruited from the general public and academic institutions around Dublin. A semi-structured interview was conducted to screen participants for past or present history of psychiatric or neurological illness. Information pertaining to any form of treatment (counselling, psychological, psychiatric), past or present, was carefully detailed, with any potential participant describing any major life-time psychiatric event or brain injury (e.g., head trauma resulting in a loss of consciousness, seizure or stroke) considered ineligible for the study. Also, any reports of familial psychiatric history (i.e. sibling, parent, grand parent) meant that people were additionally considered ineligible to participate. All participants completed inventories for mood (Beck Depression Inventory – BDI II) and drug use (questionnaire taken from the Addiction Severity index Lite-CF; see [Questionnaires](#) section) to screen for depression and past or concurrent abuse of other substances (e.g., alcohol, amphetamines, benzodiazepines, cocaine, MDMA, hallucinogens and opiates) during a subsequent in-person screening session (Beck et al., 1996; McLellan et al., 1992). Information concerning alcohol, nicotine and cannabis use in each participant was indexed in number of years (life-time) and occasions of recent use (last 30 days). Other drug use information on each participant was indexed by the total number of separate occasions (life-time) and the total number of recent separate occasions (last 30 days).

Cannabis group participants were required to have regularly consumed cannabis (5–7 days/week) for the previous 2 years in order to be eligible for the study. Participants in the cannabis group were additionally required to have smoked a minimum of 500 joints in their life-time, in order to exclude potential participants with relatively low cannabis use. All cannabis users were required to provide a positive urine sample for Δ^9 -tetrahydrocannabinol (Δ^9 THC), with additional screening for methadone, benzodiazepines, cocaine, amphetamines, opiates, barbiturates and tricyclic antidepressants (Cozart[®] RapiScan, UK) taking place. Control group participants were also tested for Δ^9 THC and the above adulterants. While the identification and quantification of cannabis metabolites in urine may have proved advantageous as a potential predictor of brain functioning, past studies have shown that estimates of recent use, life-time use and age of onset of use, are reliable predictors of behavioural performance and BOLD activity in cannabis users (Solowij et al., 2002; Bolla et al., 2002, 2005; Block and Ghoneim, 1993; Pope et al., 2003;

Pope and Yurgelun-Todd, 1996; Chang et al., 2006). Therefore, urinalyses were conducted merely to verify the presence of Δ^9 THC in cannabis participants only and the absence of all other drug metabolites in both control and cannabis participants. Following confirmation of eligibility in both cannabis users and controls, people were invited to participate in the study.

Cannabis users qualifying to participate reported, on average: 6.1 years (range = 2.5–17) of life-time cannabis use; the consumption of 7258 life-time cannabis joints (range = 700–34,403); 20 day use in the last 30 days (range = 6–30); the consumption of 64 cannabis joints in the last 30 days (range = 15–140) and 108 h of cannabis abstinence prior to testing (range = 12–504). We deliberately recruited cannabis users at varying stages of abstinence in order to test whether BOLD responses in the VS during reward processing would be associated with the subacute effects of, and withdrawal from, cannabis – our hypothesis being that life-time exposure, rather than recent use, would determine neural activity. All participants were right-handed as confirmed by the Edinburgh Handedness Inventory (Oldfield, 1971). All participants' structural brain scans were examined and cleared of structural abnormalities by a registered radiologist. All research participants provided informed consent and were financially compensated.

Monetary incentive delay task (MID)

We used a “monetary incentive delay task” (MID), which was based on that originally employed by Knutson (Knutson et al., 2001). The version used in the current study, however, differed from that originally developed on two levels. First, we did not use differential magnitudes of financial reward and punishment. Second, in the original version of the task, the anticipatory cue and trial outcome periods were time-invariantly yoked. To overcome any possible cross-contamination between anticipatory cue and outcome-related activation, we used extended temporal jittering between cue periods and target responses, and between outcome notifications and the commencement of the next trial. The temporal jittering enabled the separation of the BOLD signal related to instrumental response anticipation from outcome deliveries using deconvolution analyses. This was confirmed when designing the task by using the image analysis software to screen for multicollinearity between the anticipation and outcome regressors.

A condition of the approval we received from our institutional review board was the requirement that all participants were paid the same amount of money upon concluding the experiment (40 Euro money voucher for participation). During the practice session, prior to scanning, however, participants were informed that they would receive the total amount of money won during the experiment. Therefore, participants believed their winnings would be contingent upon their MID performance. All participants practiced the task prior to scanning, clearly demonstrating the ability to discriminate between the significance of the three “cues” used to indicate trial type (see below for cue description). While being scanned participants performed the MID task, during which they anticipated potential monetary gain, loss or no potential monetary outcome. During each trial, participants viewed one of three coloured squares (cue) that indicated the potential to win fifty cents (green square), lose fifty cents (red square) or experience no financial outcome (blue square) following their response to an upcoming visual target (see Fig. 1). Each cue was presented for a variable duration (2–8 s), after which participants made a button press response upon the presentation of a visual target (star located within a circle). Participants received feedback (1500 ms) following their response to the visual target, after which there was an end fixation period (2–8 s) before the commencement of the next trial. Responses to the visual target falling within (“hits”) or outside (“misses”) a 400 ms response deadline received feedback appropriate for that particular trial (see Fig. 1). We

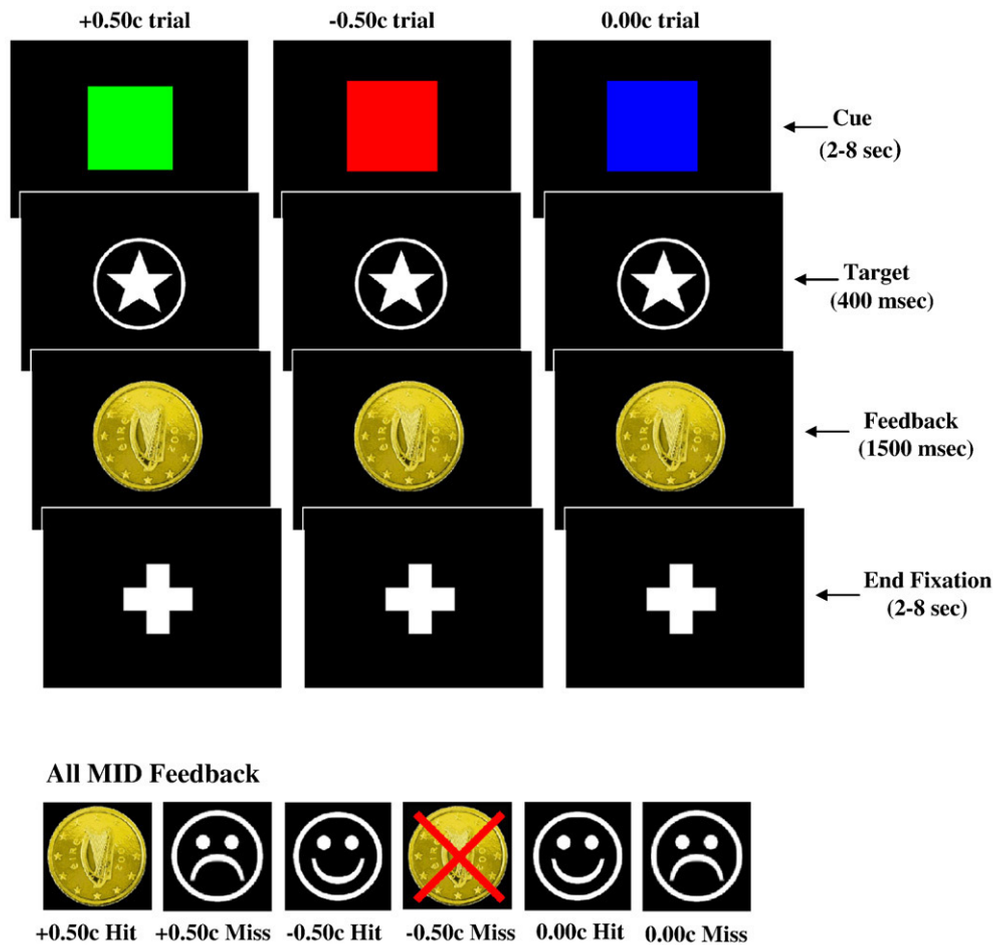


Fig. 1. Monetary incentive delay (MID) task structure. Participants were cued (2–8 s) regarding a potential financial “win”, “loss” or “no-outcome” using one of the three coloured squares. Participants were then required to respond to a target stimulus following cue presentation. The target stimulus was presented for 400 ms, during which participants were required to respond, as quickly as possible, with a button press on a hand-held key pad. Following this, participants received feedback regarding their response (1500 ms), before the presentation of an end fixation period (2–8 s), during which they saw a centrally located crosshair.

chose this 400 ms time frame in order to yield accuracy levels at around 50%, which would serve to maintain the participant's interest in the MID task. Therefore, participants had 400 ms to respond to the visual target in order to be successful on a “win”, “loss” or “no-outcome” trial. There were a total of 27 trials in each epoch (nine “win”, nine “loss” and nine “no-outcome”), with each trial lasting between 6 and 18 s. The MID was composed of three epochs, with each epoch lasting 340 s. The order of cue presentation within each epoch was randomised across the three epochs. Dependent measures derived from the data included percentage accuracy and reaction time for “win”, “loss” and “no-outcome” conditions. The task was programmed and run using E-Prime version 1.1 (Psychology Software Tools, Pittsburgh, USA).

Questionnaires

The Beck Depression Inventory II and National Adult Reading Test (NART) were administered to all participants prior to scanning (Beck et al., 1996; Nelson and O'Connell, 1978). Information concerning alcohol and drug use (see Table 1) was obtained from all participants using a questionnaire taken from the Addiction Severity index Lite-CF (McLellan et al., 1992). Prior to scanning, cannabis users also provided information concerning withdrawal and cannabis craving (Brower et al., 1988; Heishman et al., 2001). Information regarding cannabis withdrawal, modified from a cocaine withdrawal checklist, was obtained using a thirty two-item checklist, on which cannabis group

participants were required to rate, on a scale of 0 (none) to 3 (severe), symptoms they had experienced in the previous 24 h. The Marijuana Craving Questionnaire is made up of 12 statements, which the participant rates according to a seven-point Likert-type scale from “strongly disagree” to “strongly agree”. Responses to the questionnaire are then divided into four specific constructs made up of *compulsivity* (inability to control cannabis use), *emotionality* (the use of cannabis in anticipation of relief from withdrawal or negative mood), *expectancy* (the anticipation of positive outcomes from smoking cannabis) and *purposefulness* (the intention and plan to use cannabis for positive outcomes), as they relate to cannabis use.

fMRI acquisition

All scanning was conducted on a Philips Intera Achieva 3.0 Tesla MR system (Best, The Netherlands) equipped with a mirror that reflected the visual display, which was projected onto a panel placed behind the participants' head outside the magnet. The mirror was mounted on the head coil in the participants' line of vision.

Each scanning sequence began with a reference scan to resolve sensitivity variations. A parallel sensitivity encoding (SENSE) approach with a reduction factor of 2 was utilised for all T1-weighted image acquisitions (Pruessmann et al., 1999). 180 high-resolution T1-weighted anatomic MPRAGE axial images (FOV 230 mm, thickness 0.9 mm, voxel size 0.9×0.9×0.9) were then acquired (total duration

Table 1
Mean and SEM for control and cannabis groups on demographic and drug use history.

	Control (n = 14)	Cannabis (n = 14)
Age	23.1 ± 1.2	22.1 ± 1.2
Years of education	16.1 ± 0.4	17.1 ± 0.6
Verbal intelligence score (NART)	123.0 ± 0.8	123.9 ± 0.8
Beck Depression Inventory II Score	7.2 ± 2.1	8.1 ± 2.1
Females/males	3/11	2/12
Years of alcohol use	6.9 ± 1.2	6.1 ± 1.2
Alcohol use in last month (no. of days)	7.8 ± 1.4	8.3 ± 1.8
Alcohol use age onset (years)	16.2 ± 0.5	15.6 ± 0.5
Years of nicotine use	7.9 ± 1.7	5.2 ± 1.3
Cigarettes/day	10.0 ± 2.4	10.0 ± 2.8
Nicotine use in last month (no. of days)	15.0 ± 4.2	12.9 ± 4.1
Number of packs in last month	11.8 ± 3.6	12.0 ± 3.8
Nicotine use age onset (years)	16.2 ± 0.6	16.4 ± 1.0
Amphetamine use (no. of times)	3.3 ± 1.2	3.0 ± 1.8
Amphetamine use in last month (no. of times)	0.0 ± 0.0	0.0 ± 0.0
Cocaine use (no. of times)	4.8 ± 0.3	6.1 ± 1.2
Cocaine use in last month (no. of times)	0.0 ± 0.0	0.0 ± 0.0
MDMA use (no. of times)	3.6 ± 0.9	4.5 ± 0.6
MDMA use in last month (no. of times)	0.0 ± 0.0	0.0 ± 0.0
Hallucinogenic use (no. of times)	2.0 ± 0.4	3.2 ± 0.4
Hallucinogenic use in last month (no. of times)	0.0 ± 0.0	0.0 ± 0.0
Cannabis use (years)	0.0 ± 0.0	6.1 ± 1.1
Life-time joints (number)	3.0 ± 0.6	7258.6 ± 2512.8
Heavy cannabis use (years)	0.0 ± 0.0	4.5 ± 1.1
Days of use in last month (number)	0.0 ± 0.0	20.1 ± 2.5
Joints in last month (number)	0.0 ± 0.0	64.8 ± 10.7
Cannabis use age onset (years)	17.0 ± 0.3	16.1 ± 0.4
Cannabis abstinence (h)		108.0 ± 39.7
Cannabis withdrawal score (out of 32)		11.6 ± 2.1
Cannabis craving scores (each item out of 21)		
Compulsivity		5.9 ± 0.8
Emotionality		7.1 ± 1.2
Expectancy		10.6 ± 1.0
Purposefulness		12.1 ± 1.5

325 s), to allow subsequent activation localization and spatial normalization.

Functional data were acquired using a T2* weighted echo-planar imaging sequence collecting 32 non-contiguous (10% gap) 3.5 mm axial slices covering the entire brain (TE = 35 ms, TR = 2000 ms, FOV 224 mm, 64 × 64 mm matrix size in Fourier space). The functional scans had a total duration of 340 s per run.

Data processing and analyses

All analyses were conducted using AFNI software (Cox, 1996). Following image reconstruction, the three 3-D time series (epochs 1, 2 and 3) were concatenated and motion-corrected using 3-D volume registration (least-squares alignment of three translational and three rotational parameters). No participant's head moved >2 mm in any direction during the functional acquisitions. Activation outside the brain was removed using edge detection techniques. A single deconvolution analysis calculated activation for the cue and outcome periods.

Cue analyses

A regression analysis estimated activation for the “win”, “loss” and “no-outcome” cue periods. The three cue-period regressors were convolved with a standard haemodynamic response to accommodate the lag time of the blood oxygen level-dependent (BOLD) response. Multiple regression analyses calculated activation as a percentage change relative to the baseline, which consisted of the fixation period at the end of each trial.

The percentage change activation maps were re-sampled to 1 mm³ resolution, warped into standard Talairach space (Talairach and Tournoux, 1988) and spatially blurred with a 3-mm isotropic rms Gaussian kernel. Group activation maps for each condition of the task (“win”, “loss” and “no-outcome” cues) were determined with one-

sample *t*-tests against the null hypothesis of zero activation change (i.e. no change relative to the between-trial fixation periods). Using a voxelwise statistical threshold ($t = 3.4$, $p \leq 0.005$), a series of Monte Carlo simulations were then conducted (1000 iterations) to determine the frequency of clusters of significant voxels produced purely by chance. From this frequency distribution, we then selected the cluster size (276 μ l given our parameters) that occurred less than 5% of the time by chance, which, when combined with the voxelwise statistical threshold, resulted in a clusterwise threshold of $p \leq 0.05$ (corrected).

The thresholded group *t*-test maps for the “win”, “loss” and “no-outcome” cue periods were combined within each group to form group OR maps. Following this, group OR maps were combined to form an overall OR map. Thus, the overall OR map included those significant voxels from the “win”, “loss” or “no-outcome” cue periods in either the control or cannabis groups. This final map yielded functionally-defined regions of interest for between-group and between-condition comparisons. The approach taken here, wherein we determine activated areas in which to conduct cluster-level between-group comparisons, was chosen because the cluster-level analysis tends to be more statistically robust than between-group voxelwise comparisons.

Outcome analyses

We observed very large areas of significant activation during all outcome periods in both groups, which included the anterior cingulate (ACC), insula [BA13], dorsolateral prefrontal cortex (dlPFC) and nucleus accumbens (NAcc). Due to the volume of these activations, we opted for the whole brain, between groups voxelwise analysis. Analyses were conducted using the same voxelwise threshold ($p < 0.005$) and cluster criterion (276 μ l) used for the cue analyses as described above.

All between-group analyses of mean cluster activation for the cue and outcome periods were initially conducted using univariate analyses of variance. Significant group × condition interactions were decomposed using independent *t*-tests, given an *a priori* interest in BOLD differences between the two groups during specific conditions (i.e. “loss” and “win”).

Results

Demographics and drug use

Table 1 shows the group demographic and drug use history for both samples. The groups did not significantly differ on age, years of education, verbal intelligence, BDI scores or indices of alcohol, nicotine and other drug use. Cannabis use data (no. of life-time joints) were found to be significantly skewed in the cannabis sample, and therefore, log transformed (log₁₀) for further analyses.

MID performance

Fig. 2a shows the MID accuracy (% “hits”) for the three conditions in the control and cannabis groups. A two (group) by three (condition) univariate analysis of variance showed that there was a significant effect of condition ($F_{2, 78} = 6.5$, $p < 0.01$), no effect of group ($F_{1, 78} = 0.02$, $p = 0.9$) and no condition × group interaction ($F_{2, 78} = 0.5$, $p = 0.6$). Pairwise comparisons confirmed a significant difference in accuracy between the “no-outcome” and “loss” ($p < 0.05$), “no-outcome” and “win” ($p < 0.01$), but not the “loss” and “win” conditions ($p = 1.0$). Fig. 2b demonstrates MID reaction time (ms) on “hit” trials on the three conditions for both groups. Here, there was a significant effect of condition ($F_{2, 78} = 5.8$, $p < 0.01$), no effect of group ($F_{1, 78} = 3.1$, $p = 0.08$) and no condition × group interaction ($F_{2, 78} = 0.2$, $p = 0.8$). Pairwise comparisons indicated a significant difference in reaction time between the “no-outcome” and

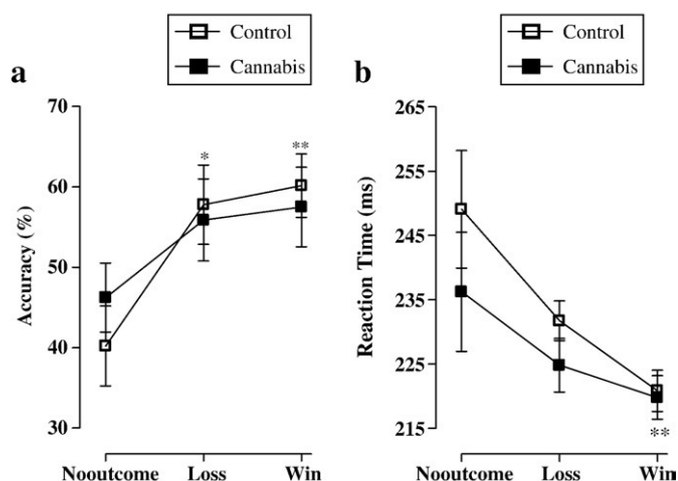


Fig. 2. (a) Mean percentage accuracy (* $p < 0.05$ “loss” versus “no-outcome”, ** $p < 0.01$ “win” versus “no-outcome”) and (b) mean reaction time (ms) (** $p < 0.01$ “win” versus “no-outcome”) in the controls and cannabis users on the MID task (means and standard errors). Data were analyzed using two (group) \times three (cue) univariate analyses.

“win” ($p < 0.01$), but not between the “no-outcome” and “loss” ($p = 0.06$) conditions.

fMRI

Cue analyses

Fig. 3 demonstrates the pattern of activation in both the cannabis and control groups during the “win” cue period of the MID task. Table 2 lists the areas of significant activity during the “win”, “loss” and “no-outcome” cue presentation periods in both the control and cannabis groups. All data across conditions and between groups in each brain region were found to be normally distributed.

Regions of significant activity for both groups were observed in both left and right hemispheres and included the cerebellum, cingulate gyrus, ventral striatum, medial frontal gyrus, fusiform gyrus and cuneus. Two (group) \times three (cue) univariate analyses demonstrated significant BOLD activity differences between the two groups in a number of areas. In the right VS there was no effect of cue ($F_{2, 78} = 2.3, p = 0.1$), a significant effect of group ($F_{1, 78} = 6.0, p < 0.05$), in which the cannabis group had significantly more activity than controls, and a significant cue \times group interaction ($F_{2, 78} = 4.5, p < 0.05$). *Post hoc* tests indicated greater BOLD activity in the cannabis group during the “win” cue ($p < 0.05$) period only (see Figs. 4a and b). We also observed a significant positive correlation between BOLD activity in this region during the “win” cue period and the number of reported life-time cannabis joints consumed ($r = 0.6, p < 0.05$) (see Fig. 4c).

Due to the large size of the observed right VS cluster (2008 μ l, encompassing the caudate, lentiform nucleus and putamen), we additionally perform a *post hoc* analysis using a higher voxelwise threshold ($t = 4.2, p \leq 0.001$) to determine the exact signal location for the observed group BOLD difference during the “win” cue period. This produced two smaller clusters. The first cluster (114 μ l) was located in the right ventral putamen ($x = 22, y = 3, z = -5$) and showed a significant effect of cue ($F_{2, 78} = 3.9, p < 0.05$), a significant effect of group ($F_{1, 78} = 4.6, p < 0.05, \text{Cannabis} > \text{Control}$) but no cue \times group interaction ($F_{2, 78} = 0.4, p = 0.7$). Pairwise comparisons for cue indicated that there was significantly greater BOLD activity during the “loss” cue compared to the “no-outcome” period ($p < 0.05$). Within group *post hoc* testing showed that in the cannabis group there was significantly more BOLD activity during the “loss” ($p < 0.05$) and “win” ($p < 0.05$) cue periods compared to the “no-outcome” cue period (see Figs. 5a and b). The second cluster (66 μ l), located in the right

putamen ($x = 19, y = 10, z = -1$), showed no effect of cue ($F_{2, 78} = 0.4, p = 0.7$), no effect of group ($F_{1, 78} = 1.2, p = 0.3$), but a significant cue \times group interaction ($F_{2, 78} = 3.6, p < 0.05$). Between group *post hoc* tests showed that the cannabis group had significantly greater BOLD activity than the control group ($p < 0.05$) during the “win” cue period (see Figs. 5a and c). We additionally observed a significant positive correlation between BOLD activity in this putamen cluster during the “win” cue period and the number of reported life-time cannabis joints consumed ($r = 0.7, p < 0.01$) (see Fig. 5d).

The right cerebellar (declive of vermis) BOLD response showed no effect of cue ($F_{2, 78} = 0.5, p = 0.6$), a significant effect of group ($F_{1, 78} = 5.7, p < 0.05$), wherein the cannabis group had significantly more BOLD activity than controls, and a significant cue \times group interaction ($F_{2, 78} = 3.9, p < 0.05$). The interaction was driven by significantly greater BOLD activity during the “loss” ($p < 0.05$) and “win” ($p < 0.05$) cue periods in the cannabis group relative to the controls (see Figs. 6a and b). We additionally observed a significant positive correlation between BOLD activity in this region during the “win” cue period and the number of reported life-time cannabis joints consumed ($r = 0.8, p < 0.001$) (see Fig. 6c).

BOLD activity in the right medial frontal gyrus [BA6] showed no effect of cue ($F_{2, 78} = 0.4, p = 0.7$), a significant effect of group ($F_{1, 78} = 6.2, p < 0.05, \text{Control} > \text{Cannabis}$) but no significant cue \times group interaction ($F_{2, 78} = 0.8, p = 0.5$). In the left fusiform gyrus [BA37], there was a significant effect of cue ($F_{2, 78} = 4.4, p < 0.05$), a significant effect of group ($F_{1, 78} = 6.8, p < 0.05, \text{Control} > \text{Cannabis}$) and a significant cue \times group interaction ($F_{2, 78} = 3.9, p < 0.05$). Pairwise comparisons for cue showed that there was greater BOLD activity during “win” compared to the “loss” ($p < 0.05$) and “no-outcome” ($p < 0.05$) periods. Further *post hoc* tests of the interaction revealed that it was only the control group that had significantly more BOLD activity during the “win” cue compared to the “loss” ($p < 0.05$) and “no-outcome” ($p < 0.05$) cue periods. Moreover, the control group had significantly more activity than cannabis users ($p < 0.05$) during the “win” cue period. Finally, for the right cuneus [BA18], there was no effect of cue ($F_{2, 78} = 1.0, p = 0.4$), a significant effect of group ($F_{1, 78} = 4.4, p < 0.05, \text{Cannabis} > \text{Control}$) but no cue \times group interaction ($F_{2, 78} = 2.1, p = 0.1$). For clusters in the left MFG and cingulate gyrus, we failed to observe any significant findings involving condition, group or any interaction between these two factors.

MID outcome analyses

Table 3 documents the results of the whole brain, between groups voxelwise analysis during the 6 outcome delivery periods. There were a number of group differences during these periods. Particularly interesting was the reduced left insula ($x = -34, y = 8, z = 7$) BOLD activity during the loss outcome delivery periods and similarly in this region ($x = -38, y = -1, z = -4$) during the loss avoidance (i.e. save 50 cents) notification periods in the cannabis group. Cannabis users demonstrated greater BOLD activity than controls during the neutral lose and neutral win outcome notification periods in a number of regions, including the caudate nucleus, cingulate, inferior frontal and parahippocampal gyri.

Drug-use correlations

Table 4 indicates that there were a number of significant positive correlations between cannabis use demographics and BOLD activity during the “win” cue period of the MID task. There were no correlations between alcohol use or other reported drug use measures and BOLD activity during the “win” cue period of the MID task. We did observe a significant negative correlation between reported cannabis withdrawal and fusiform gyrus BOLD during the “win” cue period (see Table 4). We additionally observed a significant negative correlation between superior parietal lobule BOLD activity during loss outcome notifications and life-time cannabis use ($r = -0.6, p < 0.05$) and the

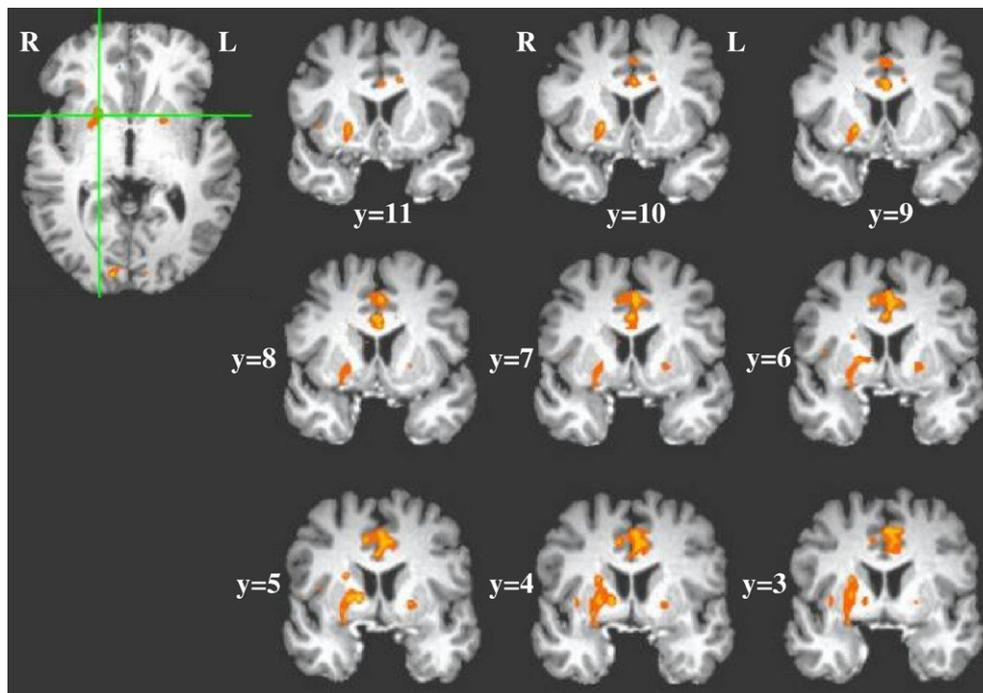


Fig. 3. Activation t -test maps (combined) for the cannabis and control groups ($p \leq 0.005$) showing coronal sections during "win" cue anticipation across the whole brain.

number of reported life-time cannabis joints consumed ($r = -0.6$, $p = 0.001$). In contrast, there were no significant correlations between self-reported use of cannabis and BOLD activity during either the "loss" or "no-outcome" cue periods. Nor were there any correlations between cue BOLD activity and MID behavioural performance (i.e. accuracy and reaction time) in either the cannabis or control groups. There were no correlations between cannabis craving and withdrawal measures and MID behavioural performance. Finally, there were no other correlations between cannabis withdrawal and cue BOLD activity or between craving and cue or outcome BOLD activity.

Discussion

The current investigation examined neural activity in chronic cannabis users during the processing of cues predicting non-drug rewards and non-drug losses. The current study, using a cohort of chronic cannabis users, who were demographically well matched to a control group, demonstrated an increased BOLD response in the right ventral striatum (VS) for cues predicting non-drug rewards. The different BOLD responses observed during reward cues occurred in the absence of any behavioural group effect on the MID, enabling us to discount performance-related neural effects from confounding these group comparisons. The observed elevated VS response to cues predictive of non-drug rewards is in contrast to that previously

observed in alcoholism and dual alcohol and cocaine dependence, suggesting that the relationship between chronic cannabis use and VS activity may be qualitatively different from that of other drugs during abstinence (Bjork et al., 2008; Wrase et al., 2007).

Animal research suggests that cues for primary reinforcers can activate DA neurons in the ventral tegmental area (VTA) and elicit DA release in the VS (Schultz et al., 1997). Human imaging studies have also demonstrated that cues for non-drug incentives reliably activate the VS BOLD response for goal-objects (Knutson et al., 2001; O'Doherty, 2004), and that reward anticipation increases DA release in this region (Schott et al., 2008). DA D_2 receptor down-regulation and reductions in pre-synaptic DA release in some drug-using groups (Heinz et al., 2004; Martinez et al., 2005; Volkow et al., 1997), it is argued, may increase the threshold required for non-drug reinforcers to activate the VS (Martin-Solch et al., 2001), thereby inducing a reward deficiency syndrome (Blum et al., 2000). The availability of striatal DA D_2 receptors, however, has not been shown to significantly differ between cannabis users and drug-naïve controls (Sevy et al., 2008), potentially ruling out a dopaminergic reward deficiency hypothesis. Given that the VTA contains a moderately high density of cannabinoid (CB_1) receptors (Herkenham et al., 1991; Tsou et al., 1998), and evidence that cannabinoids increase midbrain DA neuron activity in animals (French, 1997; Chen et al., 1990) and humans (Bossong et al., 2009), chronic cannabis use may alter reward

Table 2
Brain activations elicited by win, loss and no-outcome cue periods.

Structure	BA	HS	Vol (μ l)	x	y	z	Post hoc unpaired t -tests and p values	Direction of significance
Medial frontal gyrus	6	L	668	-5	-13	62		
Medial frontal gyrus	6	R	394	10	-9	57		Ctrl>THC
Fusiform gyrus	37	L	468	-35	-46	-10	2.7, 0.02	Ctrl>THC [W]
Cingulate gyrus		L	2220	-2	4	43		
Cuneus	18	R	276	2	-88	14		THC>Ctrl
Ventral striatum		R	2008	20	8	-4	2.8, 0.01	THC>Ctrl [W]
Declive of vermis	18	R	16,868	1	-77	-19	2.5, 0.02; 2.1, 0.05	THC>Ctrl [L] and [W]

Positive values for x, y and z Talairach coordinates denote, respectively, locations that are left, posterior and inferior relative to the anterior commissure. Table abbreviations indicate: L = left; R = right; Ctrl = controls; THC = cannabis; W = "win"; L = "loss"; NO = "no-outcome"; FG = frontal gyrus. The t and p values indicate group differences in BOLD activity using *post hoc* two-sample t -tests upon the observation of a cue \times group interaction.

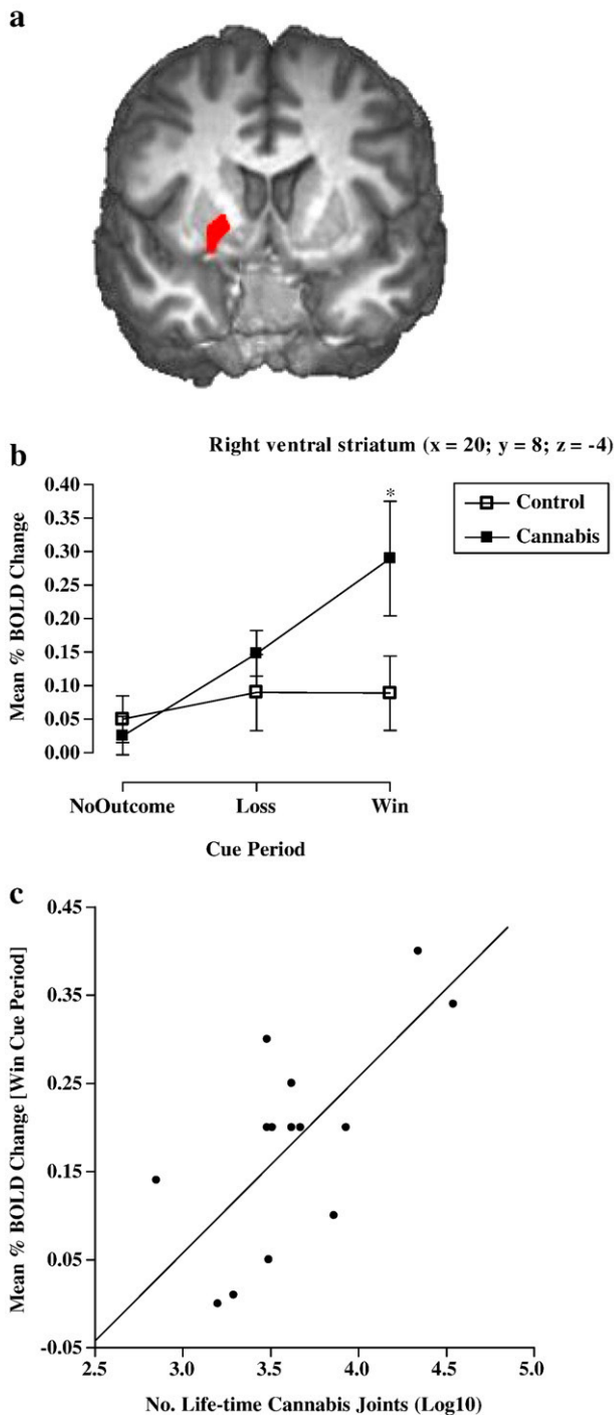


Fig. 4. (a) Mean brain activity in the right VS ($x = 20, y = 8, z = -4$), (b) graph showing cannabis and control groups significantly differed in right VS BOLD activity during “win” cue periods ($*p < 0.05$ independent t -tests) and (c) correlation between right VS BOLD activity during the “win” cue period and the number of reported life-time cannabis joints smoked ($r = 0.6, p < 0.05$).

processing through sensitizing mesolimbic circuits (under the assumption that the observed effects resulted from, rather than preceded, the cannabis use).

Past research has shown that estimates of life-time cannabis use and life-time “dose” (i.e. cannabis joints) are reliable predictors of behavioural performance and BOLD activity in cannabis users (Solowij et al., 2002; Bolla et al., 2002; Block and Ghoneim, 1993; Bolla et al., 2005; Pope et al., 2003; Pope and Yurgelun-Todd, 1996; Chang et al., 2006). Correlation analyses revealed that a participant’s reported

years of cannabis use and the number of life-time cannabis joints smoked independently predicted their VS BOLD response during cues for non-drug rewards. Importantly, this VS correlation was observed in the absence of any correlations between craving indices or other drug use measures and BOLD activity in this area, potentially ruling out these factors as contributors to the VS response.

We do not believe that the present results were influenced by cannabis abstinence at the time of testing the cannabis-using group, as we found no significant associations between hours of abstinence and task performance or BOLD responses. Cannabis users did demonstrate a reduced BOLD response in the left fusiform gyrus [BA37], which was significantly negatively correlated with reported cannabis withdrawal. The data, however, do not suggest that the VS findings reflect cannabis withdrawal, as there were no associations between withdrawal scores and BOLD activity in the VS. Furthermore, animal research suggests that there is a decline in mesolimbic DA activity during cannabis withdrawal (Diana et al., 1998), which might predict reductions in the VS BOLD response, rather than increases, as observed herein.

The incentive-sensitization theory of addiction proposes that sensitized mesolimbic neural circuits function to attribute incentive salience to reward-related stimuli, allowing reward cues to trigger excessive “wanting” for the reward (Berridge and Robinson, 1998). In drug addiction, however, the focus of sensitized “wanting” is believed to be primarily towards drug cues and drug rewards, rather than natural rewards (Robinson and Berridge, 2001). Despite this assertion, sensitization has been shown to enhance the pursuit of natural rewards in animals, where pre-treatment with amphetamine, cocaine and morphine has been observed to significantly increase cue-elicited approach behaviour for food, water and sexual contact (Mitchell and Stewart, 1990; Fiorino and Phillips, 1999; Harmer and Phillips, 1999; Taylor and Horger, 1999; Wyvell and Berridge, 2001). This may suggest that chronic pre-exposure to cannabis in humans might sensitize mesolimbic neural circuits, an effect which is manifested by cue-triggered VS responses during the pursuit of non-drug rewards. Therefore, if these results are indeed indicative of sensitization within the VS for non-drug rewards, they may also have significant implications for the future use and misuse of other drugs, as well as other forms of behaviour.

Subjective reports concerning the effects of cannabis suggest that its reinforcing properties are related to its effects on the brain’s “reward circuitry” (Huestis et al., 2007; D’Souza et al., 2008; Hunault et al., 2008; Bossong et al., 2009). There is evidence that Δ^9 THC exposure in animals affects the developmental plasticity of the reward system (Singh et al., 2006), and that the consumption of cannabis can predict a significantly higher risk for the subsequent use of other more dangerous illicit substances in humans (Fergusson and Horwood, 2000; Lessem et al., 2006). Cannabis users have also been shown to demonstrate more sexual risk (Bon et al., 2001; Castilla et al., 1999; Wingood and DiClemente, 1998; Poulin and Graham, 2001) and pathological gambling behaviour (Kausch, 2003; de Carvalho et al., 2005; Petry and Tawfik, 2001; Toneatto and Brennan, 2002), perhaps indicative of an inability to balance the immediate pursuit of rewards against the long-term negative consequences of actions. Importantly, laboratory-based evidence suggests that cannabis users have a greater level of impulsivity and an increased sensitivity for small, but immediate, rewards (Whitlow et al., 2004; Simons and Arens, 2007), consistent with the notion of mesolimbic reward hyperactivity, together with reductions in frontocortical punishment-avoidance circuitry (Solomon and Corbit, 1973, 1974; Bechara, 2005; Bickel et al., 2007). Therefore, one hypothesis, arising from the current findings, is that chronic cannabis use in humans may induce a VS hypersensitivity to other rewards (e.g., money, sex), thus increasing the likelihood of future reward seeking, risk-taking behaviour, and potentially, the pursuit of more deleterious and illicit drugs of abuse. Despite this evidence, there is the possibility that these VS differences

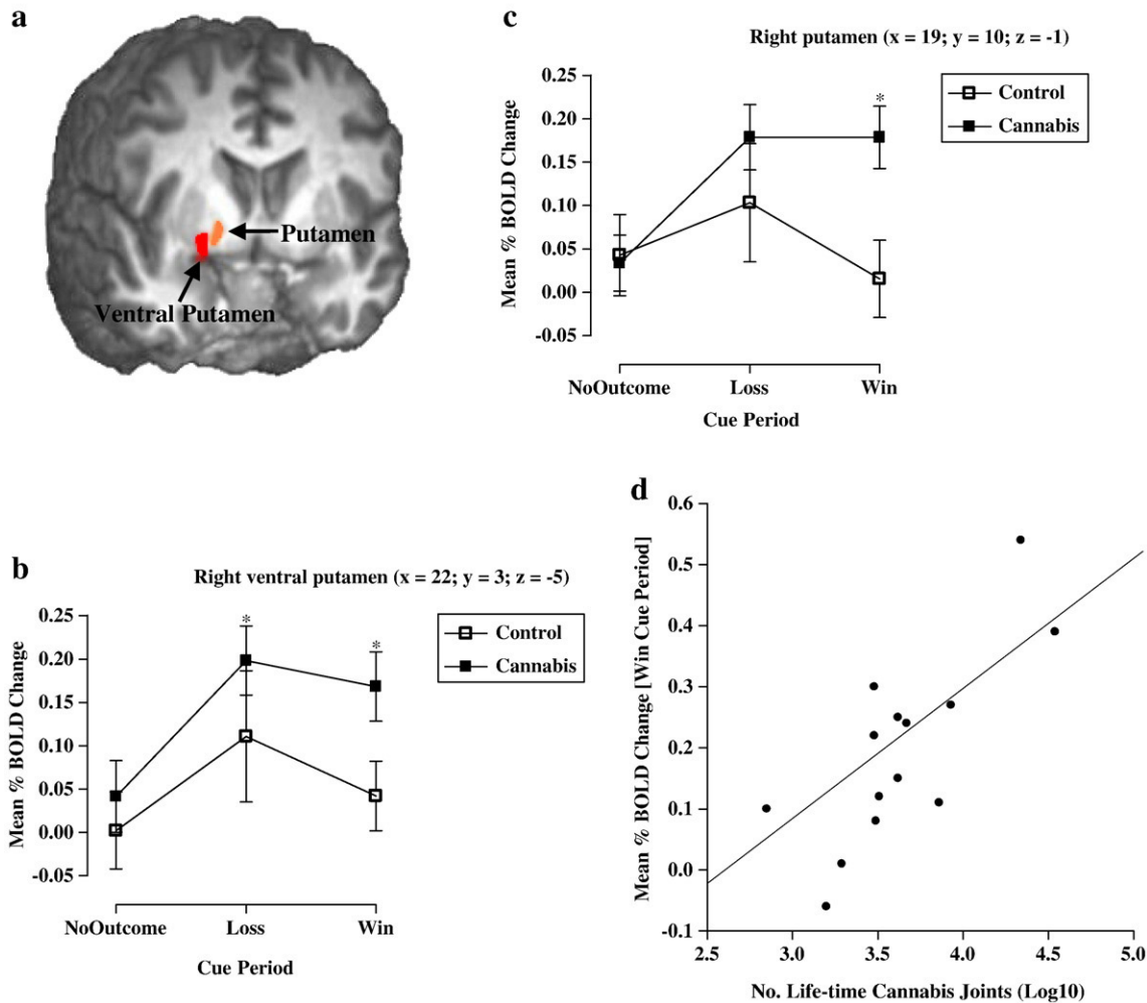


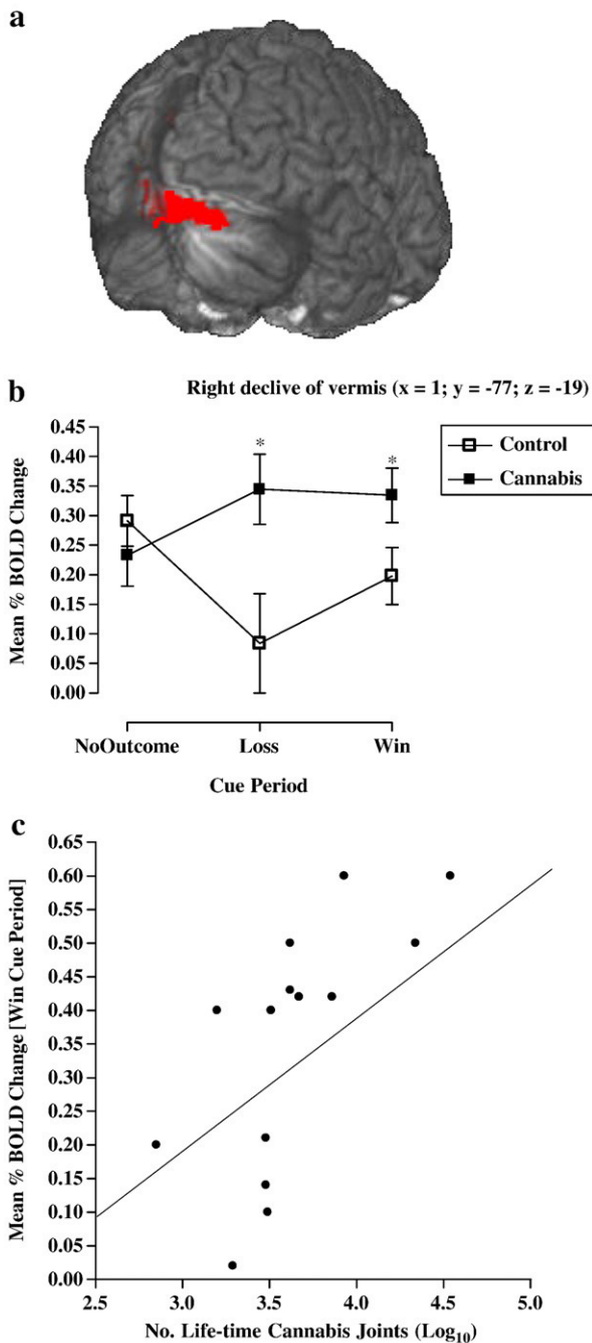
Fig. 5. Post hoc analysis showing (a) mean brain activity in the right ventral putamen ($x = 22, y = 3, z = -5$), (b) graph showing cannabis users had a greater BOLD response to “loss” and “win” cues compared with “no-outcome” cues ($*p < 0.05$ paired *t*-tests) in the right ventral putamen, (c) graph showing cannabis users had a greater BOLD response in the right putamen ($x = 19, y = 10, z = -1$) compared to controls during “win” cue presentation ($*p < 0.05$ independent *t*-tests) and (d) correlation between right putamen BOLD activity during the “win” cue period and the number of reported life-time cannabis joints smoked ($r = 0.7, p < 0.01$).

in cannabis users may have preceded cannabis use. Cannabis use may well have arisen from pre-existing differences in VS functioning. Therefore, we cannot unequivocally state that the VS differences observed herein are a direct consequence of chronic cannabis consumption and its effects on DA mesolimbic reward circuitry.

The current study also demonstrated that cannabis users had a significantly greater BOLD response in the right declive of vermis during both “loss” and “win” cue periods. There is evidence that the cerebellum plays a role in cognitive processes required for executing goal-directed behaviours and conditioned response learning (Paradiso et al., 1999; Logan and Grafton, 1995). Moreover, there is evidence for cerebellar vermis connections to DA cell body regions in the VTA, with the VTA shown to project to the cerebellum (Snider et al., 1976; Ikai et al., 1992). Vermis activity has been shown to occur during the provision of non-drug rewards or their anticipation (Rogers et al., 1999; Kunig et al., 2000; Martin-Solch et al., 2001; Knutson et al., 2001), which may explain activation patterns observed here. Furthermore, the vermis has been shown to respond to drug-related stimuli in cocaine (Volkow et al., 2003) and alcohol (Schneider et al., 2001) dependence, with increased cerebellar activity observed during cognitive tasks in alcoholics (Desmond et al., 2003) and cocaine addicts (Hester and Garavan, 2004). The present study also demonstrated that in the cannabis-using group, there were significant relationships between BOLD activity in the declive of vermis (during the “win” cue period) and cannabis use history (years of use and life-

time joints). This group difference in vermis BOLD activity, together with the observed association with cannabis use, may suggest that chronic cannabis use exaggerates cerebellar goal-directed activity in response to cues predictive of non-drug reinforcers.

We additionally observed group differences with respect to outcome-related BOLD activity, most notably detecting left insula cortex hypoactivity in the cannabis group in response to loss notification deliveries. Error-related insula activity has previously been demonstrated in healthy control subjects (Hester, Foxe, Molholm, Shpaner, and Garavan, 2005; Klein et al., 2007), and is believed to play a crucial role in integrating bodily states and affective value for reward-related adaptive behaviour (Critchley et al., 2001; Craig, 2002). Anatomically, the insula is well positioned to integrate a linking of affective value with adaptations in behaviour, possibly through its bidirectional connections with regions implicated in reward and decision making, such as the orbitofrontal cortex, amygdala, anterior cingulate and VS (Reynolds and Zahm, 2005). Research also suggests that the insula and interoceptive awareness are critical to drug craving and addiction (Gray and Critchley, 2007; Naqvi, Rudrauf, Damasio, and Bechara, 2007; Paulus, 2007), whereby the insula monitors interoceptive “urges” for rewarding stimuli such as a drug of abuse. The relative insensitivity of our cannabis group in their error-related insula activity following loss outcome notifications may also be consistent with recent findings in cannabis users demonstrating an absence of insula and anterior cingulate activity during an

**Table 3**

Voxel-wise analyses ($p < 0.005$; 276 μ l cluster criterion) showing whole brain activations elicited by the MID delivery outcome periods in the control and cannabis groups.

Structure	BA	HS	Vol (μl)	x	y	z	Direction of significance
<i>Neutral lose outcome</i>							
Cingulate gyrus	32	L	771	-1	24	27	THC>Ctrl
Superior temporal gyrus	22	R	503	48	-45	13	THC>Ctrl
Precentral gyrus	6	L	468	-21	-18	63	THC>Ctrl
Cuneus	19	L	436	-20	-84	35	THC>Ctrl
Superior frontal gyrus	8	R	419	17	47	44	THC>Ctrl
Inferior frontal gyrus	45	R	418	52	24	7	THC>Ctrl
Culmen	L	392	-37	-48	-20	THC>Ctrl	
Middle frontal gyrus	8	R	355	42	27	47	THC>Ctrl
Caudate nucleus	R	333	9	8	10	THC>Ctrl	
Postcentral gyrus	3	L	317	-49	-18	47	THC>Ctrl
Middle occipital gyrus	31	L	278	-28	-75	14	THC>Ctrl
Brainstem	L	333	0	-19	-30	THC>Ctrl	
<i>Neutral win outcome</i>							
Parahippocampal gyrus	27	L	1072	-15	-39	2	THC>Ctrl
Declive of vermis	R	546	17	-82	-17	THC>Ctrl	
Uncus	24	R	321	14	-6	-21	THC>Ctrl
Parahippocampal gyrus	37	R	285	29	-46	-8	THC>Ctrl
<i>Lose 50 cent outcome</i>							
Superior parietal lobule	7	L	672	-38	-55	54	THC>Ctrl
Insula	13	L	368	-34	8	7	Ctrl>THC
Precentral gyrus	6	R	323	35	-5	34	Ctrl>THC
<i>Save 50 cent outcome</i>							
Insula	13	L	428	-38	-1	-4	Ctrl>THC
<i>Miss 50 cent outcome</i>							
Superior frontal gyrus	8	L	581	-19	47	37	Ctrl>THC
<i>Win 50 cent outcome</i>							
Paracentral lobule	31	L	278	0	-24	45	Ctrl>THC

The two groups significantly differed in a number of regions during the different delivery outcome periods. Nomenclature: Neutral lose and Neutral win following “no-outcome” trials; Lose 50 cents and Save 50 cents following “loss” trials; Miss 50 cents and Win 50 cents delivery following “win” trials. Positive values for x, y and z Talairach coordinates denote, respectively, locations that are left, posterior and inferior relative to the anterior commissure.

The findings of the current study suggest that in chronic cannabis users, there is an increased VS BOLD response to stimuli which predict potential non-drug rewards, together with a deficit in insula activity following loss and loss avoidance outcomes. Furthermore, the observed VS hyperactivity during reward anticipation was associated with the duration (in years) of cannabis use and the estimated number of life-time cannabis joints consumed. Notwithstanding the possibility that these VS differences may have preceded cannabis use and/or be due to a small control group effect, these findings may suggest a “dose–response” sensitization effect on DA incentive processing within mesolimbic circuitry. One outstanding question with respect to the current findings concerns whether these effects would generalise to cannabis users who show high levels of

Fig. 6. (a) Mean brain activity in the right declive of vermis ($x = 1, y = -77, z = -19$), (b) graph showing cannabis and control groups significantly differed in the right declive of vermis BOLD activity during “loss” and “win” cue presentation ($*p < 0.05$ independent t -tests) and (c) correlation between the right declive of vermis BOLD activity during the “win” cue period and the number of reported life-time cannabis joints smoked ($r = 0.7, p < 0.01$).

error awareness task (Hester et al., 2009), suggesting a potential deficit in error-related monitoring in this population. We also observed insula hypofunctioning in cannabis users following loss avoidance outcome trials. While reward learning is typically associated with the mesolimbic DA system (McClure et al., 2003; O’Doherty et al., 2003), which we surmise may be sensitized by the effects of cannabis for the anticipation of rewards, this result may reflect potential differences in other neurotransmitter systems of cannabis users (e.g., serotonin, noradrenaline and acetylcholine), which are known to modulate insula activation during learning (Yu and Dayan, 2003; Berman et al., 2000).

Table 4

Correlations between BOLD activity during the “win” cue period in the cannabis group, and the cannabis use and withdrawal measures. MFG = medial frontal gyrus.

Structure	Years of use	Life-time joints	Cannabis withdrawal
Left MFG/BA6	$r = 0.4, p = 0.2$	$r = 0.3, p = 0.3$	$r = -0.2, p = 0.6$
Right MFG/BA6	$r = 0.5, p < 0.05$	$r = 0.7, p < 0.01$	$r = -0.5, p = 0.07$
Fusiform gyrus/BA37	$r = 0.4, p = 0.2$	$r = 0.3, p = 0.3$	$r = -0.7, p < 0.01$
Left Cingulate gyrus	$r = -0.4, p = 0.2$	$r = 0.7, p < 0.01$	$r = -0.1, p = 0.8$
Right cuneus/BA18	$r = 0.6, p < 0.05$	$r = 0.7, p < 0.01$	$r = -0.2, p = 0.4$
Right ventral striatum	$r = 0.6, p < 0.05$	$r = 0.6, p < 0.05$	$r = -0.04, p = 0.9$
Right ventral putamen	$r = 0.3, p = 0.3$	$r = 0.4, p = 0.1$	$r = 0.02, p = 0.9$
Right putamen	$r = 0.5, p < 0.05$	$r = 0.7, p < 0.01$	$r = -0.1, p = 0.7$
Right declive of vermis	$r = 0.7, p < 0.01$	$r = 0.8, p < 0.001$	$r = -0.4, p = 0.1$

unmotivated and avolitional behaviour, which may be more characteristic of cannabis-dependence. Furthermore, future studies will be needed to examine factors which determine whether it is drug or non-drug rewards that become excessively “wanted” in chronic cannabis users, and indeed other drug-using populations.

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