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- 1 Title page
- 2
- 3 Title

4 The acute effects of cannabidiol on the neural correlates of reward anticipation and feedback

- 5 in healthy volunteers
- 6

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72 The authors declare no conflicts of interest.

73 Abstract

74 Background

Cannabidiol (CBD) has potential therapeutic benefits for people with psychiatric disorders
characterised by reward function impairment. There is existing evidence that CBD may
influence some aspects of reward processing. However, it is unknown whether CBD acutely
affects brain function underpinning reward anticipation and feedback. *Hypotheses*We predicted that CBD would augment brain activity associated with reward anticipation and

82 feedback.

83

84 *Methods*

We administered a single 600mg oral dose of CBD and matched placebo to 23 healthy participants in a double-blind, placebo-controlled, repeated-measures design. We employed the monetary incentive delay (MID) task during functional magnetic resonance imaging (fMRI) to assay the neural correlates of reward anticipation and feedback. We conducted whole brain analyses and region-of-interest (ROI) analyses in pre-specified reward-related brain regions.

91

92 *Results*

93 The MID task elicited expected brain activity during reward anticipation and feedback, 94 including in the insula, caudate, nucleus accumbens, anterior cingulate, and orbitofrontal 95 cortex. However, across the whole brain, we did not find any evidence that CBD altered 96 reward-related brain activity. Moreover, our Bayesian analyses showed that activity in our

97	ROIs was similar following CBD and placebo. Additionally, our behavioural measures of
98	motivation for reward did not show a significant difference between CBD and placebo.
99	

100 Discussion

- 101 CBD did not acutely affect the neural correlates of reward anticipation and feedback in
- 102 healthy participants. Future research should explore the effects of CBD on different
- 103 components of reward processing, employ different doses and administration regimens, and
- 104 test its reward-related effects in people with psychiatric disorders.

106 Introduction

107 Reward processing refers to the neural, psychological and behavioural processes that underpin 108 the seeking and consumption of rewards (Berridge et al., 2009). The human brain reward 109 system is made up of key regions such as the ventral tegmental area (VTA), ventral and dorsal 110 striatum, anterior cingulate cortex, the orbitofrontal cortex, ventral pallidum, amygdala, insula, 111 thalamus and parahippocampal regions (Haber and Knutson, 2010; Knutson and Greer, 2008). 112 Fronto-striatal loops pass reward-related information from the prefrontal cortex to subcortical 113 regions and back again, such that organisms can orient attention to, be motivated for, and 114 consume rewards (Haber and Knutson, 2010).

115

116 Reward processing is perturbed in a variety of psychiatric disorders, including depression 117 (Eshel and Roiser, 2010; Knutson, Wimmer, et al., 2008; Whitton et al., 2015), addiction 118 (Balodis and Potenza, 2015; Goldstein and Volkow, 2011) and schizophrenia (Gold et al., 119 2008; Juckel et al., 2006; Strauss et al., 2013). Dysfunctional reward processing therefore 120 represents an important transdiagnostic neurocognitive mechanism which may contribute to 121 the emergence of various psychiatric disorders (Husain and Roiser, 2018; Insel, 2010; Whitton 122 et al., 2015). Hence, the reward circuit is a potential target for novel psychiatric drug treatments. Successful manipulation of the reward system could lead to the amelioration of impaired 123 124 reward learning, motivation and pleasure, observed across various clinical diagnoses.

125

The endocannabinoid system plays an important role in modulation of the brain's reward processes (Bloomfield et al., 2016; Parsons and Hurd, 2015; Solinas et al., 2009). CB1 receptors are expressed at a moderate level at the origin of the mesolimbic dopamine pathway, the VTA, and at a higher level at the terminal region, the nucleus accumbens (NAcc) (Curran et al., 2016; Solinas et al., 2009).

131

132 Cannabidiol (CBD) is the second most abundant cannabinoid in the cannabis plant (Upton et 133 al., 2014; Pertwee, 2008) and at typical doses CBD is non-intoxicating (Haney et al., 2016; 134 Hindocha et al., 2015; Lawn et al., 2016; Martin-Santos et al., 2012). CBD has therapeutic 135 potential in a variety of psychiatric disorders (Freeman et al., 2019; Khan et al., 2020). 136 Preclinical research has demonstrated that CBD administration can affect reward-related 137 behaviours, particularly reducing drug-seeking behaviour (Hay et al., 2018; Katsidoni et al., 138 2013; Parker et al., 2004; Ren et al., 2009; Schier et al., 2014; Viudez-Martínez et al., 2018). 139 Speculatively, CBD could ameliorate addictive behaviour by enhancing the sensitivity of the 140 reward system to natural rewards, such that pharmacological rewards are less desired. The 141 effects of CBD on the mesolimbic dopamine system are, however, equivocal (Renard et al., 142 2017).

143

Human research has shown that CBD can acutely alter neural, behavioural and psychological 144 145 processes relating to reward, including effort sensitivity (Lawn et al., 2016), attentional bias to 146 drug pictures (Hindocha et al., 2018; Morgan et al., 2010), drug consumption (Freeman et al., 147 in press; Morgan et al., 2013), neural response to music reward (Freeman et al., 2018) and levels of stress-induced social anxiety (Bergamaschi et al., 2011; Zuardi et al., 1993), without 148 149 producing reinforcing or unpleasant side-effects (Haney et al., 2016). However, it is not known 150 if CBD specifically acts on the human brain's reward circuitry, or acts by another mechanism. 151 Furthermore, if CBD does act on the reward system, its effects on reward anticipation and 152 reward feedback have not been parsed.

153

154 The monetary incentive delay (MID) task is a well-validated functional magnetic resonance 155 imaging (fMRI) task which, through its structure, allows for investigation of the neural 156 correlates of reward anticipation and reward feedback (Balodis and Potenza, 2015; Knutson et 157 al., 2001). Meta-analyses of MID task results show reward anticipation and feedback recruit 158 overlapping and distinct regions (Knutson and Greer, 2008; Oldham et al., 2018). Both 159 processes activate striatal regions, while reward anticipation activates the thalamus and insula, 160 and reward feedback preferentially activates prefrontal cortex areas. Importantly, neural 161 activity during reward anticipation in the ventral striatum correlates with dopamine release in 162 the same region (Schott et al., 2008), demonstrating the task engages the mesolimbic dopamine 163 system.

164

165 CBD seemingly has opposite effects to the primary intoxicating cannabinoid found in cannabis, 166 delta-9-tetrahydracannabinol (THC), on both brain and behavioural outcomes (Bhattacharyya 167 et al., 2010; Bloomfield et al., 2016; Englund et al., 2013). CBD enhanced striatal activation 168 during a verbal memory task, while THC dampened striatal activity (Bhattacharyya et al., 169 2010). In the MID task, acute THC administration has been shown to attenuate the widespread 170 neural response to reward feedback (van Hell et al., 2012) and attenuate the neural response in 171 the nucleus accumbens during reward anticipation in people with nicotine dependence (Jansma 172 et al., 2013). Therefore, one might expect CBD to do the opposite: augment neural response to reward anticipation and feedback. Furthermore, a pro-reward function action could underlie 173 174 CBD's putative anti-addiction, anti-depressant and anxiolytic effects.

175

In summary, the endocannabinoid system plays an important role in the brain's reward circuitry and both preclinical and human research has demonstrated that CBD can modulate rewardrelated behaviours. However, previous human studies have tended to investigate CBD's impact alongside THC. Moreover, they have focused on psychiatric symptom-based measures, rather than precise components of reward processing, such as anticipatory and consummatory reward

181 processes which are indexed by the well-validated MID task. No study has examined the 182 specific, isolated effect of CBD on the human brain during reward processing. Based on its 183 opposing effects to THC and its ostensibly therapeutic effects in disorders characterised by 184 reward dysfunction, we predicted that CBD would augment the neural response to reward 185 anticipation and feedback.

187 Methods

188 *Design and participants*

The study used a double-blind, randomized, placebo-controlled, repeated-measures design to compare the effects of oral CBD 600mg with matched placebo (PBO). Drug order was balanced and randomised. Drug order was completely concealed from participants and concealed from experimenters until data collection, entry, and analysis had been completed.

193

We tested 28 healthy participants. Four participants did not complete both sessions, so they were excluded. Furthermore, one participant did not complete the MID task correctly, so they were excluded. That left 23 participants in our analysis.

197

198 Participants were recruited through public advertisement. Inclusion criteria were: (1) age 18-199 70 years; (2) right-handed; (3) fluent in English. Exclusion criteria were: (1) positive urine 200 screen for recreational drug use (Alere Toxicology UC-10A; amphetamines, barbiturates, 201 benzodiazepines, cocaine, methamphetamine, morphine, methadone, phencyclidine, tricyclic 202 antidepressants, THC), (2) recent (within the past six months) use of any psychotropic 203 (recreational or medical) drug, including cannabis, (3) positive breath test for alcohol, (4) 204 carbon monoxide \geq 5 parts per million (ppm), (5) problematic alcohol use, as defined by a score 205 \geq 8 on the Alcohol Use Disorder Identification Test (AUDIT) (Saunders et al., 1993), (6) more 206 than ten lifetime uses of cannabis or CBD, (7) more than five lifetime uses of any other 207 recreational drug, (8) nicotine dependent, as defined by a score greater than three on 208 Fagerstrom Test for Nicotine Dependence (Heatherton et al., 1991), (9) current or past mental 209 or physical health issues or learning impairments, based on an adapted version of the Diagnostic and Statistical Manual of Mental Disorders-IV (DSM-IV) Structured Clinical 210 211 Interview (SCID) (Gibbon and Spitzer, 1997), (10) positive reading on urine pregnancy test,

(11) breast-feeding, (12) known allergies or aversions to CBD, microcrystalline cellulose,
gelatine or lactose, (13) colour blindness, (14) MRI contraindications, (15) current use of
psychiatric medications.

215

Participants were reimbursed £10/hour for their time. This study was approved by the UCL
ethics committee (Project Number: 3325/002), and all participants provided written informed
consent.

219

220 <u>Assessments</u>

221 The Monetary Incentive Delay (MID) task (Knutson et al., 2000) (Figure 1)

The MID task is a well-validated task that allows measurement of neural activity during reward anticipation and reward feedback using functional magnetic resonance imaging (fMRI). We used an adapted version of the original (Knutson et al., 2000).

225

226 In our version of the task, a cue (a square) is first presented for 500ms, which signals whether 227 the trial is a win trial (if the square is orange) or a neutral trial (if the square is blue). On a win 228 trial, the participant has the opportunity to win 30p if they respond to a subsequent target in 229 time. On a neutral trial, the participant cannot win or lose any money, but they are asked to 230 respond to the subsequent target as quickly as they can anyway. Following the cue, there is a 231 blank screen, the anticipation phase, for 2-4s in which the participant waits for the target. 232 Subsequently, the target (a white square) is presented and the participant must respond to it as 233 quickly as they can by pressing a button with their thumb on their right hand. Initially, 234 participants must respond to the target within 300ms in order to get a 'hit'. However, following 235 a successful 'hit', the next trial's target must be responded to within a time that is 16.67ms 236 shorter than the previous trial in order to get another 'hit'. Following a 'miss', the next trial's 237 target must be responded to within a time that is 16.67ms longer than the previous trial in order 238 to get a 'hit'. This is to calibrate the participant's performance to 'hit' roughly 50% of the time. 239 Following the target, feedback is presented for roughly 1000ms (although this changes on a 240 trial-by-trail basis along with changes in target duration). If it is a 'win' trial and the participant 241 gets a 'hit', then the participant wins 30p and is told 'Hit. You win 30p'. If it is a 'win' trial 242 and the participant gets a 'miss', then the participant does not win money and is told 'Miss'. If 243 it is a 'neutral' trial and the participant gets a 'hit', then the participant does not win money and 244 is told 'Hit'. If it is a 'neutral' trial and the participant gets a 'miss', then the participant does 245 not win money and is told 'Miss'. The current total won is always displayed on the feedback 246 screen. Following the feedback, there is an inter-trial interval (ITI) between 1.2 and 9.2s when 247 a blank screen is shown.

248

There are 48 trials in total, of which 24 are neutral trials in which no money can be earned and 24 are win trials in which money can be earned. The order of win trials was fixed, so that win trials did not appear consecutively. Each win trial provides the opportunity to win 30p; this amount does not vary, as in some previous MID task versions (Knutson et al., 2008). There are also no loss trials. The task lasts for 12 minutes.

254

The MID task produces measures of brain activity associated with reward anticipation and reward feedback. It also produces behavioural measures of mean reaction time to respond to the target on successful 'win' and 'neutral' trials and the proportion of 'hits' on 'win' and 'neutral' trials.

259

260 [Insert Figure 1]

262	Demographics	1
202	Demographics	

263 We recorded participants' age, sex, weight and BMI.

- 264
- 265 Beck Depression Inventory (BDI) (Beck et al. 1996)
- 266 A self-reported scale of depression severity which consists of 21 items. This measured the
- 267 participants' depressive symptomatology over the preceding two weeks to the first study visit.
- 268 Higher scores reflect a higher severity of depression.
- 269
- 270 Alcohol Use Disorder Identification Test (AUDIT) (Saunders et al., 1993)
- A self-reported scale which screens for problematic alcohol use and consists of 10 items. Scores
- range from 0 to 40, with higher scores reflecting more severe problematic alcohol use. A score
- 273 of 8 or more is considered hazardous.
- 274
- 275 Fagerstrom Test for Nicotine Dependence (FTND) (Heatherton et al., 1991)
- 276 A self-reported scale of nicotine dependence consisting of six items. Total scores range from 0
- to 10, with higher scores reflecting higher nicotine dependence.
- 278
- 279 Wechsler Test for Adult Reading (WTAR) (Ginsberg et al., 2003)
- A test of reading ability which is a proxy of verbal intelligence. It includes 50 words that must
- 281 be read aloud and pronounced correctly.
- 282
- 283 Plasma CBD levels
- 284 Blood samples were collected using EDTA vacutainers and centrifuged immediately. Plasma
- samples were stored at -80°C prior to analysis. CBD concentrations were determined using gas
- chromatography mass spectroscopy (GC/MS) with a lower limit of quantification of 0.5mg/ml.

287

288 Drug Administration

289 Participants were administered a single dose of 600mg oral CBD (pure synthetic (-)-CBD, STI 290 Pharmaceuticals, Essex, England) or matched placebo (lactose powder) in identical, opaque 291 capsules on each testing session. The CBD was formulated in 50mg capsules. Participants 292 swallowed all 12 capsules at their own pace under invigilation of the experimenter. 600mg was 293 chosen as it produces an increase in plasma concentrations after acute administration 294 (Babalonis et al., 2017; Englund et al., 2013), is well tolerated in humans (Grotenhermen et al., 295 2017), produces a significant anxiolytic effect (Bergamaschiet al., 2011), produces opposing 296 effects to THC on the striatum as assessed by fMRI (Bhattacharyya et al., 2010), and elicits 297 anti-psychotic like effects in combination with THC (Bhattacharyya et al., 2015).

298

299 <u>Procedure</u>

300 Participants completed a screening on the telephone during which initial eligibility criteria 301 (drug use, FTND, AUDIT, MRI contraindications, allergies, medical information, and 302 handedness) were assessed and basic participant details were recorded. Participants that 303 appeared eligible on the phone were invited to attend experimental sessions. Participants were 304 asked to fast from midnight the day before both sessions, and refrain from smoking tobacco 305 and consuming alcohol for 24 hours before the start of the sessions. Upon arrival, participants 306 underwent urine tests to verify they were not pregnant (if female) and they had not recently 307 taken recreational drugs. They also completed breath tests for alcohol and carbon monoxide.

308

309 Eligible participants then completed two seven-hour experimental sessions, when they received
310 CBD or PBO on the first session, and the other drug condition on the second session.
311 Experimental sessions were separated by a minimum seven-day wash-out period (>4 times the

elimination half-life) to minimize carryover effects of CBD (Consroe et al., 1991). The BDIand WTAR were completed immediately after drug administration on the second session.

314 Previous research suggests that CBD reaches the peak level of plasma concentration after 315 approximately 2.5 hours (Babalonis et al., 2017). Therefore, 2.5 hours after drug 316 administration, participants underwent MRI scanning for 1.5 hours to complete the MID task, 317 as well as other tasks and scans, which will be reported elsewhere. Participants' blood samples 318 were taken straight after the scan finished, which was approximately 4 hours and 15 minutes 319 after drug administration. After a standardised lunch provided by the experimenter, participants 320 completed a series of questionnaires and computer tasks, results of which will be reported elsewhere. 321

322

323 <u>Power calculation</u>

A power calculation was conducted using G*Power (version 3.1.9.2). This showed that a sample size of 20 would have 81% power to detect a significant (p<0.05, two-tailed) difference between CBD and placebo (PBO) with a moderate or greater effect size of d=0.5. This effect size was based on the previous finding of the difference in the attentional bias toward cigarette cues between 800mg CBD vs. placebo in nicotine-dependent users (Hindocha et al., 2018). We then recruited extra participants to account for expected participant dropout and exclusions.

330

331 MRI data acquisition

MRI data was collected using a 3-Tesla Siemens Verio MRI Scanner at the Robert Steiner MR unit at Hammersmith Hospital, London. Functional imaging used a multiband (acceleration factor = 2) gradient-echo T2*-weighted echo-planar imaging (EPI) sequence with 42 slices per volume (TR = 2400ms; TE = 30ms; in-plane matrix = 64 x 64; 3mm isotropic voxels; flip angle = 62° ; bandwidth = 1594 Hz/pixel; 304 volumes; a slice thickness of 3mm; field of view = 192mm x 192mm). The phase encoding direction was from anterior to posterior. Echo spacing was 0.71ms. There were 3 dummy scans at the beginning of the scan, which were not included in in our dataset. For structural acquisition, a T1-weighted structural volume was acquired for all participants using a Magnetisation Prepared Rapid Gradient Echo (MPRAGE) scan (TR = 2300ms; TE = 2.28ms, TI= 900ms, flip angle = 9°, field of view= 256mm, image matrix = 256 with 1-mm isotropic voxels; bandwidth = 200 Hz/pixel).

343

344 *Functional magnetic resonance imaging (fMRI) data analyses*

Image pre-processing and analysis were performed using FSL's fMRI Expert Analysis Tool
(FEAT) (FMRIB Software Library v6.0, Analysis Group, FMRIB, Oxford, UK) (Jenkinson et

al., 2012). Data were pre-processed before being subject to first and second-level analyses.

348

349 Pre-processing

350 FSL's brain extraction tool (BET) was used to strip the brain from the skull. FMRIB Automated 351 Segmentation Tool was used to separate out grey matter, white matter, and cerebrospinal fluid. 352 Functional images were realigned to the middle volume using FSL's MCFLIRT procedure, in 353 order to correct for head motion. Subsequently, the functional images were co-registered to the 354 individual participant's structural image and normalised to the MNI-152 (Montreal 355 Neurological Institute) template using FEAT's non-linear transformation procedure with a 356 10mm warp resolution. An isotropic 6mm full-width at half-maximum Guassian kernel (i.e. 357 twice the voxel size) was then applied to spatially smooth images. A high-pass filter (100s cut-358 off) was applied to remove low-frequency noise. Images were visually inspected to ensure that 359 the pre-processing had worked correctly.

T₁-weighted structural images were also skull-stripped with FSL's BET and normalised to the
 MNI-152 template.

363

364 *First level analyses*

365 Timestamps and durations for each event (cue, anticipate, target, feedback, inter-trial-interval) in the MID task were extracted from the task output files using scripts written in Matlab 366 367 (Mathworks Inc., United States). A general linear model was created with the following 368 explanatory variables (i.e. regressors): (1) reward anticipation (i.e. anticipate-win), (2) no 369 reward anticipation (i.e. anticipate-neutral), (3) reward feedback on a successful win trial (i.e. 370 feedback-win-hit), (4) no reward feedback on an unsuccessful win trial (i.e. feedback-win-371 miss), (5) no reward feedback on a successful neutral trial (i.e. feedback-neutral-hit), (6) no 372 reward feedback on an unsuccessful neutral trial (i.e. feedback-neutral-miss). Each event was 373 modelled with a boxcar function with the event's duration convolved with the canonical haemodynamic response function, using the gamma function. Extended motion parameters and 374 375 temporal derivatives were included as additional regressors-of-no-interest.

376

377 These contrasts were then calculated:

378 (1) 'reward anticipation': anticipate-win > anticipate-neutral.

- 379 (2) 'reward feedback': feedback-win-hit > feedback-neutral-hit.
- 380

381 Second level analyses

382 Whole brain analysis

383 The second-level fMRI data analysis was also performed with FSL's FEAT pipeline (Jenkinson

et al., 2012), using a random effects analysis with FMRIB's Local Analysis of Mixed Effects

(FLAME). We analysed the two contrasts specified above at the second level. We used clusterwise correction, with a cluster-defining threshold of z=2.3 and an alpha value of 0.05. We conducted one-sample t-tests for both contrasts, collapsing across both drug conditions, to investigate the overall effect of the task (reward anticipation and reward feedback) on brain activity. Secondly, we conducted paired t-tests for both contrasts to investigate the differences, in both directions, between CBD and PBO.

391

392 Region of interest (ROI) analyses

393 ROIs were pre-specified based on a meta-analysis of MID fMRI results for significantly 394 activated regions for reward anticipation and feedback (Knutson and Greer, 2008). There were 395 eight ROIs for anticipation and seven ROIs for feedback, as shown in Table 1. The Talairach 396 coordinates from Knutson and Greer (2008) were converted to MNI coordinates using the 397 mni2tal MATLAB function created by the University of Cambridge Medical Research Council 398 Cognition and Brain Sciences Unit (http://imaging.mrc-399 cbu.cam.ac.uk/imaging/MniTalairach). We used these coordinates as the centres for our spherical ROIs, with radii of 5mm. The ROIs were created using FSLeyes and fslmaths 400 401 functions. We then extracted average unstandardized beta values (with arbitrary units) from 402 these regions for the two contrasts described above.

403

We then ran one-sample t-tests (against a score of zero) to test whether the task elicited the expected anticipation and feedback activation in the hypothesised regions. Subsequently, we ran paired t-tests for an effect of drug (CBD vs. PBO) on the activation in these anticipation and feedback ROIs. We reduced the alpha value to 0.006 to account for the multiple tests (i.e. ROIs) within each contrast.

We examined the extracted beta values for normality by visually inspecting histograms of the data, checking for kurtosis and skewness values >1, using Kolmogrov-Smirnov tests and looking for outliers as shown by SPSS's box and whisker plots. Across all regions, for both CBD and PBO and for both reward anticipation and feedback the data were normally distributed, so data were left unchanged.

415

416 [Insert Table 1]

417

418 In order to gain further support for either the null or alternative hypothesis for the effects of 419 CBD on brain activity during reward anticipation and feedback, we also calculated scaled 420 Jeffreys-Zellner-Siow (JZS) Bayes online calculator factors using an 421 (http://pcl.missouri.edu/bayesfactor) (Buckingham et al., 2016; Lawn et al., 2018). We used a scaled-information prior of r = 1, which is the default value recommended (Rouder et al., 2009). 422 423 For this analysis, a Bayes factor of >3 provides support for the null hypothesis (i.e. no 424 difference in activation between CBD and placebo).

425

We conducted Pearson correlations between participant CBD plasma levels and their extracted beta values for each anticipate and feedback ROI, when they were on the CBD condition. We reduced the alpha value to 0.006 to account for multiple tests (i.e. ROIs) within each contrast.

429

430 <u>Behavioural analyses</u>

We conducted a Wilcoxon signed-rank test on the plasma CBD levels for CBD compared withPBO.

- 434 We conducted 2x2 repeated-measures analyses of variance (ANOVAs) for reaction time (RT)
- 435 and the proportion of hits, with within-subjects factors of drug (CBD, PBO) and trial-type (win,
- 436 neutral).

- 437 **Results**
- 438 *Demographics*
- 439 Of the 23 participants included in the analysis, there were 12 women and 11 men, with mean
- 440 age 23.74 years (SD=4.2, range: 19-36). Participants' depression (BDI mean=2.2, SD=4.9,
- 441 range: 0 to 11) and problematic alcohol use (AUDIT mean=2.2, SD=2.8, range: 0-7) levels
- 442 were low. Participants had a mean WTAR raw score of 40.5 (SD=4.9, range: 33-49) and a
- 443 mean BMI of 22.4 kg/m² (SD=3.5, range: 17.6-35.4).
- 444
- 445 <u>Plasma CBD levels</u>
- Plasma CBD levels were higher on CBD (median=6.01ng/ml, interquartile range=4.89) than
 PBO (median=0, interquartile range=0) (Z=3.296, p=0.001).
- 448

449 <u>MID behavioural results</u>

For RT, there were main effects of drug ($F_{1, 22}=6.286$, p=0.020) and trial-type ($F_{1, 22}=15.841$, p=0.001), but there was not a significant interaction. Participants were faster to respond on win trials (mean=0.241s, SD=0.023) compared to neutral trials (mean=0.247s, SD=0.024). Participants were faster, overall, to respond under PBO (mean=0.241s, SD=0.024) compared to CBD (mean=0.247s, SD=0.024).

455

For proportion hit, there was a main effect of trial-type ($F_{1, 22}$ =43.776, p<0.001), but no main effect of drug or interaction. Participants were more likely to hit on a win trial (mean=0.612, SD=0.079) compared to a neutral trial (mean=0.437, SD=0.072).

459

461 <u>MID fMRI results</u>

Movement did not exceed 3mm (our voxel size) in any direction for any of the participants.
Mean and maximum movements were: x: mean=0.15mm (SD=0.50mm), max=0.50mm; y:
mean=0.19mm (SD=0.12), max=0.50mm; z: mean=0.34mm (SD=0.32mm), max=2.00.
Therefore we did not exclude any participants for excess movement.

466

467 Whole brain analyses

468 *Effects of task (Table 2, Figure 2, Figure 3)*

For the reward anticipation contrast, there was activation in three clusters, with peak activations in the insula bilaterally and the right paracingulate gyrus (Table 2). The right and left insula clusters extended into the right and left frontal operculum cortex, inferior frontal gyrus and orbitofrontal cortex. The paracingulate gyrus extended into the anterior cingulate gyrus, supplementary motor cortex and superior frontal gyrus (Figure 2).

474

475 For the reward feedback contrast, there was very widespread activation in two large clusters: 476 one more posterior and one more anterior (Table 2; Figure 3). The posterior had a peak 477 activation in the left occipital fusiform gyrus and extended into the bilateral cerebellum, intracalcarine gyrus, lingual gyrus, precuneus, inferior and middle temporal cortex, anterior 478 479 and posterior lateral occipital gyrus, postcentral gyrus, posterior supramarginal gyrus, and 480 hippocampus, amongst others. The anterior cluster had a peak activation in the left precentral 481 gyrus and extended into the bilateral anterior cingulate cortex, paracingulate gyrus, superior 482 and middle frontal gyrus, frontal pole, precentral gyrus, frontal medial cortex, and frontal 483 operculum, amongst others. Activity was also observed in bilateral caudate, accumbens, 484 thalamus and pallidum.

486	[Insert Table 2]
487	[Insert Figure 2]
488	[Insert Figure 3]
489	
490	Effects of the drug
491	No significant clusters were found for CBD>PBO or PBO>CBD for either reward anticipation
492	or feedback.
493	
494	ROI analyses
495	Effects of task (Table 3)
496	For reward anticipation, only the right insula was significantly activated (t_{22} =3.87, p=0.001)
497	during reward anticipation.
498	
499	For reward feedback, the left $(t_{22}=3.31, p=0.003)$ and right $(t_{22}=3.38, p=0.003)$
500	parahippocampal gyri, right caudate (t_{22} =3.46, p=0.002) and left nucleus accumbens (t_{22} =4.02,
501	p=0.001) were significantly activated during reward feedback.
502	
503	[Insert Table 3]
504	
505	Effects of drug (Table 4)
506	CBD did not differ from PBO in all of the ROIs during reward anticipation (ps>0.1).
507	Furthermore, all but one of the ROIs had a Bayes factor>3, in favour of there being no

508 difference between drug conditions.

510 CBD did not differ from PBO in all of the ROIs during reward feedback (ps>0.3). Furthermore,

all the ROIs had Bayes factors>3, in favour of there being no difference between drugconditions.

- 513
- 514 [Insert Table 4]
- 515

516 <u>Correlations</u>

- 517 There were no significant correlations between plasma CBD levels and activation in any of the
- 518 ROIs during anticipation or feedback.
- 519

521 Discussion

522 We hypothesised that brain activity would be greater during reward anticipation and feedback 523 following 600mg of oral CBD compared to PBO. However, this was not the case. We found 524 no evidence that CBD affects the brain's response to reward anticipation or feedback. 525 Furthermore, in pre-specified reward-related brain regions (Knutson and Greer, 2008), using 526 Bayesian analyses, we found support for there being no difference in neural activity between 527 CBD and PBO. Overall, we found no support for CBD affecting the neural correlates of reward 528 anticipation and feedback or behavioural measures of motivation for reward in healthy 529 volunteers.

530

531 Across both drug conditions, in the whole brain, our MID task elicited reward anticipation 532 activation in the bilateral insula and paracingulate gyrus, extending into inferior frontal gyri 533 and orbitofrontal cortex. In our ROI analysis, the right insula was significantly activated during 534 reward anticipation. Reward feedback elicited extensive activity across anterior and posterior 535 parts of the brain, including a range of reward-related brain regions. In our ROI analysis, the 536 right caudate, left nucleus accumbens and bilateral parahippocampal gyri were activated during 537 reward feedback. These analyses demonstrate that anticipation and feedback of reward produced activity in several expected brain regions. Further support that the task functioned 538 539 adequately is that both reaction time and hit rate were significantly affected by trial type, such 540 that participants were faster and more likely to successfully hit the target on win trials compared 541 to neutral trials. Importantly, our plasma results demonstrate that the 600mg oral dose of CBD 542 was absorbed.

543

In terms of behavioural outcomes, CBD led to longer reaction times compared to PBO overall.
However, there was no interaction between drug and trial-type; CBD did not reduce reaction

times more for win trials than it did for neutral trials. Hence CBD did not affect our behavioural measure of motivation for reward; it simply increased reaction time, in general (i.e. comparably for both trial-types). This is somewhat surprising given previous research has not found CBD to affect reaction speed in general (Belgrave et al., 1979; Fusar-Poli et al., 2009; Hindocha et al., 2018).

551

552 Despite some existing evidence that CBD can impact reward function, we found null results 553 for its effects on the neural correlates of reward anticipation and feedback. This absence of 554 impact on reward circuitry, may contribute to the lack of reinforcing and abuse potential of 555 CBD (Haney et al., 2016). To our knowledge, no previous study has examined the effects of 556 CBD alone on brain activity associated with reward processing or motivation for reward. 557 Previous studies have often investigated how inhaled CBD moderates THC's effects (Freeman 558 et al., 2018; Lawn et al., 2016), which may have contributed to the discrepancy. Moreover, 559 other studies have explored more complex components of reward function, including 560 attentional bias toward drug pictures (Hindocha et al., 2018; Morgan et al., 2010). Other 561 components of reward processing, including reward learning and subjective pleasure could also 562 still be sensitive to a 600mg dose of oral CBD. CBD's acute effects on human behaviour and subjective experience are seemingly complicated and enigmatic (Bergamaschi et al., 2011; 563 564 Fusar-Poli et al., 2009; Haney et al., 2016; Morgan et al., 2010). The same may well be true 565 with regards to CBD's impacts on reward processing.

566

567 Furthermore, long-term daily administration of CBD, as delivered in clinical trials (Freeman et 568 al., in press; Leweke et al., 2012; McGuire et al., 2018), could produce different effects on the 569 neural correlates of reward anticipation and feedback. We only delivered a single oral 600mg 570 dose in healthy volunteers. CBD likely has complex, variable dose-response functions on diverse psychological outcomes (Zuardi et al., 2017). Nevertheless, experimental medicine approaches, such as this one, are needed to efficiently examine the acute effects of potentially therapeutic drugs in human models of psychiatric targets, where clinical trials are costly and protracted. Future research into CBD's effects on reward processing should expand the reward components assessed and utilise different doses. It should also examine consequences of repeated, long-term administration, which may allow for CBD levels to build up in the body and have greater impacts on receptor expression and endocannabinoid levels.

578

579 The present results leave open the intriguing possibility that CBD may only exert an effect on 580 reward networks that have already been perturbed, for example in people with a drug addiction. 581 CBD administration has been shown to modulate reward-related behaviours in animals when 582 addiction is being modelled (Katsidoni et al., 2013; Parker et al., 2004; Ren et al., 2009; Schier 583 et al., 2014; Viudez-Martínez et al., 2018). Moreover, behavioural evidence from human 584 studies suggests that CBD can reduce the salience of drug-related cues in those with cannabis 585 (Morgan et al., 2010) and nicotine (Hindocha et al., 2018) dependencies, and reduce drug cue-586 induced cravings in those addicted to heroin (Hurd et al., 2019). Additionally, a four-week 587 treatment of CBD dose-dependently decreased cannabis use in a clinical trial of people with 588 cannabis use disorder (Freeman et al., in press). In all of these studies, CBD attenuated atypical 589 reward-related behaviours conferred by addiction, suggesting a restorative effect. Therefore, 590 the null findings reported in the present study could have resulted from our sample of healthy 591 volunteers. Future neuroimaging research should therefore administer CBD to participants 592 thought to have perturbed reward systems, including those with addiction.

593

594 The reward system is thought to be critically involved in the emergence and/or maintenance of 595 a variety of psychiatric disorders, including depression (Nestler and Carlezon, 2006; Whitton 596 et al., 2016), schizophrenia (Kapur et al., 2005; Whitton et al., 2016) and addiction (Berridge 597 and Robinson, 2016; Goldstein and Volkow, 2011). If it emerges that CBD does have accepted 598 therapeutic effects in these domains, further research will be needed to understand whether or 599 not the mechanism is related to reward circuitry. Moreover, an improved understanding of 600 CBD's pharmacological actions and their relative importance in treating reward-related 601 psychological symptoms will be important in the development of cannabinoid-based 602 psychiatric medicines. One possible avenue for future research would be to further understand 603 and capitalize on CBD's agonism of the serotonin-1a receptor (Russo et al., 2005), in order to 604 potentially disrupt addition and depressive symptoms.

605

606 Strengths and Limitations

607 Our study has a number of strengths. First and foremost, it was a double-blind, placebo-608 controlled experiment addressing a novel and important research question. Second, we utilised 609 a well-validated fMRI task which elicited activity in many expected brain regions and 610 appropriately affected behavioural performance. Third, CBD was absorbed into the 611 bloodstream. Fourth, we conducted Bayesian analyses to provide support for null findings.

612

613 However, there are some limitations. Despite stimulating activity in many expected brain 614 regions, the MID failed to produce anticipatory activation in the striatum, which is the region 615 most commonly found to respond in this stage of the task (Oldham et al., 2018). Thus, CBD 616 could theoretically affect striatal activity (Bhattacharyya et al., 2010) and we may have failed 617 to detect it here. Finally, although CBD was absorbed relative to placebo, our plasma levels 618 were lower than that seen in previous oral CBD studies (Haney et al., 2016; Millar et al., 2018). 619 This may have been caused by our fasting participants, as a large, high-fat meal eaten before 620 CBD administration can augment bioavailability four-fold (Taylor et al., 2018). Therefore, we cannot exclude the possibility that if greater quantities of CBD had been absorbed, we may have observed different results. We also do not know whether 600mg is the optimal dose to manipulate reward processing, especially given CBD's potentially inverted U-shaped doseresponse curve (Zuardi et al., 2017). Additionally, we did not control or account for female participants being in different stages of their menstrual cycle, which can affect psychopharmacological phenomena (Bolea-Alamanac et al., 2018).

627

628 Conclusion

To conclude, in healthy volunteers, a single, oral 600mg dose of CBD did not affect the neural
correlates of reward anticipation and feedback, or behavioural measures of motivation for
reward.

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