

1 **Adolescent THC impacts on mPFC dopamine-mediated cognitive processes in male and female rats**

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11

12 **Abstract**

13 *Rationale:* Adolescent cannabis use is linked to later-life changes in cognition, learning, and memory. Rodent experimental
14 studies suggest Δ^9 -tetrahydrocannabinol (THC) influences development of circuits underlying these processes, especially
15 in the prefrontal cortex, which matures during adolescence.

16 *Objective:* We determined how 14 daily THC injections (5mg/kg) during adolescence persistently impacts medial prefrontal
17 cortex (mPFC) dopamine-dependent cognition.

18 *Methods:* In adult Long Evans rats treated as adolescents with THC (AdoTHC), we quantify performance on two mPFC
19 dopamine-dependent reward-based tasks—strategy set shifting and probabilistic discounting. We also determined how
20 acute dopamine augmentation with amphetamine (0, 0.25, 0.5 mg/kg), or specific chemogenetic stimulation of ventral
21 tegmental area (VTA) dopamine neurons and their projections to mPFC impacts probabilistic discounting.

22 *Results:* AdoTHC sex-dependently impacts acquisition of cue-guided instrumental reward seeking, but has minimal effects
23 on set-shifting or probabilistic discounting in either sex. When we challenged dopamine circuits acutely with amphetamine
24 during probabilistic discounting, we found reduced discounting of improbable reward options, with AdoTHC rats being more
25 sensitive to these effects than controls. In contrast, neither acute chemogenetic stimulation of VTA dopamine neurons nor
26 pathway-specific chemogenetic stimulation of their projection to mPFC impacted probabilistic discounting in control rats,
27 although stimulation of this cortical dopamine projection slightly disrupted choices in AdoTHC rats.

28 *Conclusions:* These studies confirm a marked specificity in the cognitive processes impacted by AdoTHC exposure. They
29 also suggest that some persistent AdoTHC effects may alter amphetamine-induced cognitive changes in a manner
30 independent of VTA dopamine neurons or their projections to mPFC.

31
32 Key words: THC, cognition, dopamine, ventral tegmental area, medial prefrontal cortex, chemogenetics

34 Introduction

35 Cannabis is one of the most widely used drugs among adolescents, and its availability is increasing around the world.
36 Human studies show that early exposure to cannabis, and especially its main psychoactive constituent Δ^9 -
37 tetrahydrocannabinol (THC), is associated with later-life cognitive impairments, and increased risk for psychiatric disorders
38 including schizophrenia and addiction (Curran et al. 2016; Ehrenreich et al. 1999; Jenni et al. 2017; Malone et al. 2010;
39 Murray et al. 2022; Rubino and Parolaro 2016; Schneider 2008; Volkow et al. 2016). However, in humans it is difficult to
40 dissect whether THC exposure causes these associations, or whether early cannabis use and long-term deficits both result
41 instead from other underlying comorbidities or risk factors. Rodent models are thus essential for establishing casual effects
42 of THC on the developing adolescent brain.

43 Neurodevelopmental disruptions persisting long after adolescent cannabis use are plausible because adolescence is a
44 dynamic critical period for structural and functional brain remodeling, especially in late-developing structures like the
45 prefrontal cortex (PFC) (Andersen 2003; Casey et al. 2000). Some of this age-dependent plasticity seems to involve the
46 endocannabinoid system, with dynamic changes in cannabinoid receptors (CBRs) and endocannabinoids (ECB) occurring
47 across adolescence (Bara et al. 2021; Ellgren et al. 2008; Heng et al. 2011; Lee et al. 2016; Simone et al. 2022). Might
48 THC, which also acts via CBRs, disrupt this age-dependent ECB signaling system and thus leave long-lasting
49 consequences on the brain? If so, the adolescent-developing PFC (Peters et al. 2022; Scheyer et al. 2023; Spear 2000),
50 and its dopaminergic inputs from ventral tegmental area (VTA), which are actively innervating during this period (Hoops and
51 Flores 2017; Manitt et al. 2011; Reynolds et al. 2018), are a likely candidate for cognition-relevant neurodevelopmental
52 insults caused by adolescent THC (Molla and Tseng 2020; Renard et al. 2017a; Renard et al. 2017b).

53 The adult PFC is crucial for purposeful, goal-directed behaviors driven by the ability to flexibly converge our internal states,
54 like reward motivation, with outside external information about contexts, cues, and rules (Miller 2000; Miller and Cohen
55 2001; Ott and Nieder 2019). Executive functions like working memory, attention, rule shifting, and decision-making require
56 PFC-dependent cognitive control. Rodent studies show adolescent cannabinoid drug exposure can cause persistent deficits
57 in working memory (De Melo et al. 2005; O'Shea et al. 2004; Quinn et al. 2008; Schneider and Koch 2003), social cognition
58 (O'Shea et al. 2004; Renard et al. 2017a; Renard et al. 2017b; Zamberletti et al. 2014), and cognitive flexibility (Egerton et
59 al. 2005; Gomes et al. 2015; Jacobs-Brichford et al. 2019; Szkudlarek et al. 2019) that may depend upon PFC.

60 Furthermore, dopamine in PFC plays a major role in decision making, working memory, cognitive flexibility, and goal-
61 directed behaviors (Floresco 2013; Goldman-Rakic 1995; Goto et al. 2007; Seamans and Yang 2004). PFC dopamine

dysfunction has also been implicated in schizophrenia symptoms (Davis et al. 1991; Howes and Kapur 2009), and there is a clear link between recent cannabis use and onset of psychosis in humans (Andreasson et al. 1987; D'Souza et al. 2004; Hambrecht and Hafner 2000), though it is not clear that this link is causal in nature (Fergusson et al. 2005; Henquet et al. 2005; Sewell et al. 2009). Since VTA dopamine neuron axons actively infiltrate mPFC during adolescence (Hoops and Flores 2017; Manitt et al. 2011; Reynolds et al. 2018) and are thus subject to disruption by THC exposure, and since THC impacts dopamine signaling at several key nodes in reward and salience circuits (Behan et al. 2012; Corongiu et al. 2020; Ferland et al. 2023; Renard et al. 2017a; Renard et al. 2017b), it is plausible that adolescent THC might exert some of its neurodevelopmental disruptions by impacting the function of cognitive flexibility-relevant dopaminergic inputs to mPFC.

We therefore explored this possibility in a series of experiments quantifying effects of a well-characterized, translationally relevant adolescent THC exposure protocol in rats (Ruiz et al. 2021a; Ruiz et al. 2021b; Torrens et al. 2022; Torrens et al. 2020) upon adulthood mPFC dopamine-dependent cognition.

Materials and Methods

Subjects

Long Evans rats ($n = 29$ males, 25 females) were used for set shifting, probabilistic discounting and amphetamine experiments, and transgenic TH:Cre ($n = 19$ males, 9 females) and wildtype littermates ($n = 14$ males, 4 females) were used for chemogenetic experiments. Rats were pair or triple-housed in same-sex groups at weaning (postnatal day; PD21), in a temperature, humidity, and pathogen-controlled colony under a 12:12hr reverse light/dark cycle. Water was provided *ad libitum* and food was restricted to ~85% of free-feeding weight during behavioral testing, starting at ~PD70. Experiments were approved by University of California Irvine's Institutional Animal Care and Use Committee and were conducted in accordance with the NIH Guide for the Care and Use of Laboratory Animals.

Drugs

THC was provided in ethanol by the NIDA Drug Supply Program. For injection, THC was prepared fresh each day; ethanol is evaporated under N_2 , and THC is reconstituted in 5% Tween-80 in saline, with heat and sonication, to 2 ml/kg for intraperitoneal (IP) injection. D-amphetamine hemisulfate salt (amphetamine) was attained from Sigma and mixed in saline at 0.5 mg/ml for injection. Clozapine-N-oxide (CNO) was obtained from the NIDA Drug Supply Program, stored at 4°C in powder aliquots with desiccant, protected from light. For systemic injection, CNO (5 mg/kg) was dissolved daily for IP injection in 5% dimethyl sulfoxide (DMSO; Sigma Aldrich) in 0.9% saline. For microinjections, CNO (1mM; 0.5 μ l/side over

60s) was dissolved in artificial cerebrospinal fluid (Fisher) with 0.5% DMSO, stored in aliquots at -20°C, and thawed/vortexed just before use.

Adolescent THC and Washout

All rats received daily IP injections of THC (5 mg/kg) or vehicle (VEH; 5% Tween-80 in saline) from PD30-43, followed by a washout period of 21+ days, allowing full THC clearance in both sexes (Lee et al. 2022) prior to behavioral testing in adulthood (**Fig. 1a**).

Experimental Design

Following adolescent THC/VEH treatment, 48 adult (PD 70+) rats ($n = 29$ males, 21 females) underwent strategy set-shifting training, followed by training on probabilistic discounting until a group displayed stable levels of choice for 3 consecutive days, determined with previously established criteria (St. Onge and Floresco 2009). Thereafter they underwent 3 counterbalanced amphetamine challenge tests, each (0, 0.25, 0.5 mg/kg IP) delivered 5 min before behavioral testing. Following the challenge rats were retrained for at least 2 days before receiving their next challenge.

For chemogenetic experiments, another adolescent THC/VEH-treated group ($n = 33$ males, 13 females; $n = 28$ TH:Cre+, 18 wildtype (WT)) underwent stereotaxic VTA virus injection of a Cre-dependent hM3Dq vector at ~PD65, and intra-mPFC bilateral cannulae implantation at least 45 days later, at least 8 days prior to the first microinjection. Following recovery from surgery, they were trained on the probabilistic discounting task to stability, then subjected to a series of counterbalanced tests, again with re-stabilization training occurring between them. First, rats received 4 counterbalanced IP injections of the DREADD agonist CNO (5 mg/kg) or VEH, 30 min prior to behavioral testing. Both CNO and VEH were administered twice on separate days, and data was combined between both tests for analysis. Next, they underwent 4 additional counterbalanced tests of probabilistic discounting, each held 5 min after intra-mPFC microinjections of CNO or VEH. Results from intra-mPFC VEH and CNO tests were again averaged to increase reliability of findings. For choice data, the raw number of risky choices in each block made on both VEH or CNO tests were summed and divided by the total number of choices made in those respective blocks on the two VEH/CNO tests, thereby factoring out trial omissions. Win-stay/lose shift values were combined in a similar manner, dividing the sum total of “stays” or “shifts” by the sum total of “wins” and “losses” on the two respective tests. Latency and omission values were averaged across the tests.

Behavioral Methods

Operant Boxes

121 Training and testing took place in Med-Associates rat operant conditioning chambers (30.5 x 24 x 21 cm; St Albans, VT)
122 within sound-attenuating boxes, equipped with two retractable levers with white lights above them, and white house light.

124 *Operant Pretraining*

125 Methods for both strategy set shifting and reward probabilistic discounting tasks closely followed prior reports (Floresco et
126 al. 2008; St. Onge and Floresco 2009). Rats were first homecage-habituated to highly palatable, 45 mg banana-flavored
127 reward pellets (Bio-Serv catalogue #: F0059), then given 2 days of magazine training in the operant boxes, where they
128 received 38 pellets at variable intervals over a 60 min session. They were next sequentially trained to press each of two
129 levers to receive a pellet over 2-5 days. On each lever training day, a lever extended into the chamber (side counterbalanced
130 across rats), and one pellet was delivered on a fixed ratio 1 (FR1) schedule. Once they reliably pressed 50+ times in 30 min
131 on a lever, they were transitioned to learning to press the other lever on the subsequent day, and training continued until
132 meeting this criterion. Thereafter, they entered the next phase of the task, in which each lever was periodically extended
133 into the chamber throughout a 30 min session. Levers extended in a pseudorandom order such that there were 45 left-lever
134 trials and 45 right-lever trials, but no more than two consecutive trials on which the same lever was extended. Each lever
135 extension was accompanied by illumination of a house light signaling the start of a trial (90 trials/session), and trials occurred
136 every 20s. If the rat pressed the lever within 10s of the start of a trial, the lever was retracted and a pellet was delivered,
137 and the house light remained on for 4s. If the rat failed to press in 10s, the lever retracted and the house light extinguished
138 until the next trial, and the trial was considered an omission. Importantly, cue lights present above each lever were never
139 illuminated during this training phase. Rats were trained in this manner for at least 5 days, or until they omitted less than 5
140 trials per session. Since individual rats frequently display idiosyncratic lever position biases that can influence interpretation
141 of subsequent behavior (Brady and Floresco 2015), and to minimize such impacts of stochastic side-preference on
142 subsequent tests, lever side preference was next assessed using a published protocol (Brady and Floresco 2015; Floresco
143 et al. 2008).

145 *Visual Cue Discrimination Training*

146 Next, rats were trained to respond on only one of the two levers extended on each trial—whichever was signaled as the
147 correct response by illumination of a cue light just above it (**Fig. 1b**). Sessions began with both levers retracted and the
148 house light off. Every 20s, one of the two stimulus cue lights was illuminated in a randomized order, 3s later both levers
149 extended, and the house light turned on. If the rat pressed the lever that had a cue light illuminated above it within 10s of
150 lever insertion, it received a pellet, the levers retracted, the stimulus light was extinguished and the house light remained
151 illuminated for 4s. If the rat did not choose a lever in 10s, the trial ended and levers retracted until the next trial, and the

152 trial was scored as an omission. This training proceeded for at least 30 trials, ending when rats either made 10 consecutive
153 correct responses, or after 150 trials had elapsed. If rats did not achieve 10 consecutive correct responses, they received
154 a second identical training session on the following day.

156 *Strategy Set Shifting Test*

157 After learning to follow a cue light to respond for reward in the prior training phase, rats then underwent a single 40 min
158 session on which the response rule was suddenly shifted; a procedure analogous to the Wisconsin Card Sorting Task used
159 in humans to aid diagnosis of PFC dysfunction (Owen et al. 1991; Pantelis et al. 1999). On this day, the non-preferred lever
160 (determined in side-bias training described above) became the correct response, and pressing it during trials yielded a pellet
161 and commencement of the next inter-trial interval (**Fig. 1h**). Light cues above one of the levers were presented just before
162 and during trials as in the prior visual cue discrimination training, but now their location was irrelevant to the receipt of
163 reward. Instead, rats needed to recognize that this old rule (follow the cue light) no longer worked, and that pressing of the
164 previously less-preferred lever, regardless of cue light position, was now the correct strategy for obtaining reward. Trials
165 continued until rats performed 10 consecutive correct responses, or after 150 trials had occurred. Errors during set-shift
166 were categorized into two subtypes as in (Floresco et al. 2008): perseverative errors, where rats responded on the incorrect
167 lever when the previously-relevant visual cue was illuminated above it, and never-reinforced (or non-perseverative) errors,
168 where rats pressed the incorrect lever despite when the visual cue was illuminated above the correct lever.

170 *Probabilistic Discounting*

171 In this task rats chose between a lever that always delivers a small (1 pellet) “certain” reward, or another that delivers a
172 large (4 pellet) “risky” reward, delivered at various probabilities throughout the 52.5 min session (**Fig. 2a**). The probability
173 of receiving 4 pellets upon pressing the risky lever decreases in each of five sequential choice blocks during each session
174 (100%, 50%, 25%, 12.5%, 6.25%). Each block began with 8 forced trials (35s apart), in which the houselight was illuminated
175 and 3s later, one of the two levers extended one at a time for 10s (4 trials for each lever, randomized in pairs). This permitted
176 rats to sample the probability of reward receipt upon pressing of the risky lever in the upcoming certain/risky lever choice
177 phase of the block. During the subsequent choice phase of each block, both levers were extended, and after rats pressed
178 one of them, both levers retracted. Certain lever presses always yielded 1 tasty banana pellet, and risky lever choices
179 delivered 4 or 0 banana pellets, at a probability that varied across blocks. Failure to respond on a lever within 10s on any
180 trial resulted in lever(s) retraction and extinguishing of the house light, and the trial was scored as an omission. Animals
181 were trained on this task for 21 days, at which point choice behavior had stabilized across groups. For rats tested with
182 systemic amphetamine during probabilistic discounting, training on this task commenced following strategy set shifting

183 training and testing described above. Rats in chemogenetic experiments did not undergo visual cue discrimination or set
184 shifting tests. Instead, these rats received pellet habituation, magazine training, initial FR1 training, and retractable lever
185 training prior to training on the probabilistic discounting task.

186
187 For rats that received acute amphetamine challenges (0, 0.25, 0.5 mg/kg IP) on risky decision making, 3 counterbalanced
188 tests were conducted on separate days, with re-training between tests to reestablish stable baseline responding. When rats
189 in chemogenetic experiments achieved stable baseline responding, they similarly received 4 counterbalanced tests, 2 with
190 CNO and 2 with VEH (data from both tests combined for analysis as described above), all injected IP 30 min prior to
191 probabilistic discounting testing, with retraining between tests. After completing these systemic CNO/VEH tests, rats were
192 implanted with mPFC cannulae, allowed to recover, and re-stabilized on discounting performance for 2+ days. They were
193 then tested on the discounting task in 4 additional counterbalanced tests with intra-mPFC microinjections of CNO (1mM/0.5
194 μ l; 2 tests) and VEH (2 tests).

196 **Chemogenetic Methods**

197 *Virus Surgery*

198 Rats were anesthetized with ketamine (56.5 mg/kg) and xylazine (8.7 mg/kg) and given meloxicam (1.0 mg/kg) for pain
199 prophylaxis. An AAV2 vector containing a Cre-dependent, mCherry-tagged hM3Dq excitatory DREADD (hSyn-DIO-hM3Dq-
200 mCherry; titer: 6×10^{12} vg/ml; Addgene catalog #: 44361) was injected bilaterally into VTA (relative to bregma (mm): AP: -
201 5.5, ML: ± 0.8 , DV: -8.1; 0.75 μ L/hemisphere) using a Picospritzer and glass micropipette (Martinez et al. 2023). Injections
202 occurred steadily over 1 min, and the pipette was left in place for 5 min after injection to limit spread. Both TH:Cre and WT
203 rats were injected with the active hM3Dq DREADD virus. Colocalization of mCherry expression to TH+ VTA neurons was
204 verified in each TH:Cre rat, and lack of mCherry expression was confirmed in each WT rat. At least 3 weeks elapsed
205 between virus injections and the first CNO administration. In a second surgery held following behavioral training and
206 systemic CNO tests (to maximize accuracy of placement) guide cannulae (22 ga, 2MM, Plastics One) were implanted
207 bilaterally in the mPFC (relative to bregma (mm): AP: 2.7, ML: ± 1 , DV: -3.1) to allow intra-mPFC CNO injection upon VTA
208 dopamine neuron axon terminals, and occluded with steel stylets between tests.

210 *Histological Validation*

211 Following behavioral testing, chemogenetic rats were perfused with chilled 0.9% saline and 4% paraformaldehyde, brains
212 were cryoprotected in 20% sucrose-azide, and they were sectioned coronally at 40 μ m. VTA virus expression was amplified
213 with mCherry immunohistochemistry, and expression verified to be in dopamine neurons via co-staining of tyrosine

214 hydroxylase. VTA sections were then blocked in 3% normal donkey serum PBST, tissue was incubated overnight at room
215 temperature in rabbit anti-DSred (Clontech; 1:2500) and mouse anti-TH (Immunostar; 1:2000). After washing, sections were
216 incubated in dark at room temperature for 4 hours in AlexaFluor-donkey anti-Rabbit 594 and donkey anti-Mouse 488
217 (Thermo Fisher Scientific; 1:500). Sections were mounted and cover slipped (Fluoromount; Thermo FisherScientific),
218 mCherry/TH expression was imaged at 10X on a Leica DM4000 epifluorescent scope, and expression or lack thereof in
219 VTA was verified in each animal. To verify cannula placement within mPFC, sections were nissl stained with cresyl violet,
220 and cannula tracks were mapped using a rat brain atlas (Paxinos and Watson 2006).

222 **Data Analysis**

223 Effects of adolescent treatment (AdoTX: THC or VEH) and Sex (M or F) on learning and cognition employed 2 x 2 ANOVA
224 and Sidak posthoc tests. Effects of amphetamine (0 x 0.25 x 0.5mg/kg) on probabilistic discounting were tested with
225 repeated measures ANOVA, and interactions of these acute treatments with AdoTX and Sex were examined by adding
226 these between subjects variables to multivariate General Linear Model ANOVAs. Simple-main effects analyses conducted
227 after observing statically significant interactions included one-way ANOVA models using the appropriate error term from
228 the overall multifactor analyses. Effects of CNO versus VEH were treated as within-subjects variable, while Genotype
229 (TH:Cre x WT), AdoTX, and sex were treated as between subjects variables in multivariate General Linear Model ANOVAs.
230 Prior to testing, rats were verified to display stable patterns of risky choice by examining discounting performance over at
231 least two consecutive prior training days. Rats of both sexes were tested in chemogenetic experiments, but sample sizes
232 for each sex were insufficient to allow formal analysis of this variable (Fig. S1). Six rats ($n = 2$ males, 4 females; $n = 1$ VEH,
233 5 THC) were excluded from set shifting analyses for failure to meet training criteria for instrumental or visual cue rule
234 performance, and nine rats ($n = 9$ females; $n = 2$ VEH, 7 THC) tested with amphetamine were excluded for failing to achieve
235 stable performance on the probabilistic discounting task. Nine rats ($n = 3$ males, 6 females; $n = 4$ VEH, 5 THC) were
236 excluded from the DREADD experiments for failure to stabilize on the probabilistic discounting, death, or inability to confirm
237 virus expression.

239 **Results**

240 **Effects of Adolescent THC on Strategy Set Shifting in Each Sex**

241 *Initial Instrumental Training:* AdoTHC did not affect acquisition of instrumental food seeking behavior during initial training
242 (no main effect of AdoTX: $F_{s(1, 45)} < 0.40$, $ps > 0.53$), and acquisition was similar in both sexes (No main effect of sex: $F_{s(1,$
243 $45)} < 3.60$, $ps > 0.06$; no Sex x AdoTX interaction: ($F_{s(1, 45)} < 1.12$, $ps > 0.30$).

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Visual Cue Discrimination: Rats were next trained to press whichever lever that had a cue light illuminated above it. AdoTHC exposed rats were more likely to learn the visual cue rule in one session rather than two, relative to AdoVEH rats ($X^2: p = 0.04$). This was driven by AdoTHC females, as evidenced by these animals requiring fewer trials and making fewer errors to reach criterion performance of 10 consecutive correct choices (**Fig. 1c-g**, AdoTX x Sex interaction; trials to criterion: $F_{(1,44)} = 5.31, p = 0.03$; errors to criterion: $F_{(1,44)} = 5.03, p = 0.03$; Šídák's posthoc for trials $p = 0.02$, errors $p = 0.02$). In addition, relative to controls, AdoTHC females were slower to respond ($F_{(1,44)} = 4.55, p = 0.04$) but also made fewer omissions: ($F_{(1,44)} = 8.46, p = 0.01$). In contrast, no differences were observed on these measures between AdoTHC males and controls (trials to criterion: $p = 0.87$; errors: $p = 0.89$; latency: $p = 0.68$, omissions: $p = 0.72$). More generally, we also saw some sex-differences on performance measures irrespective of treatment, as females were slower to respond (Sex: $F_{(1,44)} = 13.02, p < 0.001$) and omitted more trials ($F_{(1,44)} = 5.60, p = 0.02$) compared to males.

Strategy Set Shifting: Following training on the initial visual cue discrimination, rats were then trained to use an egocentric spatial rule (always press the left/right lever regardless of cue location), and tested in a single-session (Fig. 1i-m; Brady and Floresco 2015). AdoTHC had few effects on the ability to learn this new response rule with no change in either sex seen on the number of trials to reach criterion on the new rule (No main effect of AdoTX: $F_{(1,44)} = 0.42, p = 0.52$; or Sex: $F_{(1,44)} = 0.01, p = 0.92$; or interaction: $F_{(1,44)} = 0.80, p = 0.38$). Total errors to criterion were also unaffected (AdoTX: $F_{(1,44)} = 0.97, p = 0.33$; Sex: $F_{(1,44)} = 0.32, p = 0.57$; AdoTX x Sex interaction: $F_{(1,44)} = 0.19, p = 0.66$), as were errors of either a perseverative or never-reinforced subtype (AdoTX: $F_{(1,44)} = 1.56, p = 0.22$; Sex: $F_{(1,44)} = 0.13, p = 0.72$; AdoTX x Sex: $F_{(1,44)} = 0.58 \times 10^3, p = 0.98$). Additionally, we saw no effect of AdoTX on response latency or on omissions (AdoTX: $F_{(1,44)} < 2.62, ps > 0.11$; AdoTX x Sex: $F_{(1,44)} < 0.01, ps > 0.94$). Again, we saw that females took longer to respond (Sex: $F_{(1,44)} = 7.52, p = 0.01$) and omitted more trials ($F_{(1,44)} = 6.20, p = 0.02$) compared to males. Since visual discrimination learning was more efficient in AdoTHC females than in AdoVEH females, but shifting to a spatial rule was equivalent in both groups, we wondered if performance on rule #1 could have led to more robust or persistent learning that could have indirectly impacted performance on the set shift. We therefore performed a secondary analysis where we matched the performance of the two female AdoTX groups on the visual discrimination rule #1. We removed the 4 AdoTHC female rats that learned rule #1 quickest, and the 5 AdoVEH females that learned rule #1 slowest, thus yielding equivalent rule #1 performance in the retained rats from both groups ($n=6$ /group; Errors: AdoVEH: M (SEM) = 17.83 (6.18) AdoTHC: M (SEM) = 19.00 (2.94); $t(10) = 0.17, p = 0.87$; Trials: AdoVEH: M (SEM) = 16.83 (22.83); AdoTHC: M (SEM) = 64.67 (9.58); $t(10) = 0.17, p = 0.870$). Additionally, in this subset, performance on the shift to rule #2 remained equivalent (Errors: AdoVEH: M (SEM) = 79.33 (16.07); AdoTHC: M (SEM) = 91.50 (13.04); $t(10) = 0.22, p = 0.83$; Trials: AdoVEH: M (SEM) = 27.17 (6.58); AdoTHC: M (SEM) = 28.83 (3.95);

275 $t(10) = 0.59, p = 0.57$), rather than showing an AdoTHC-induced deficit as would be expected if rule #1 performance
 276 impacted subsequent shift to rule #2.

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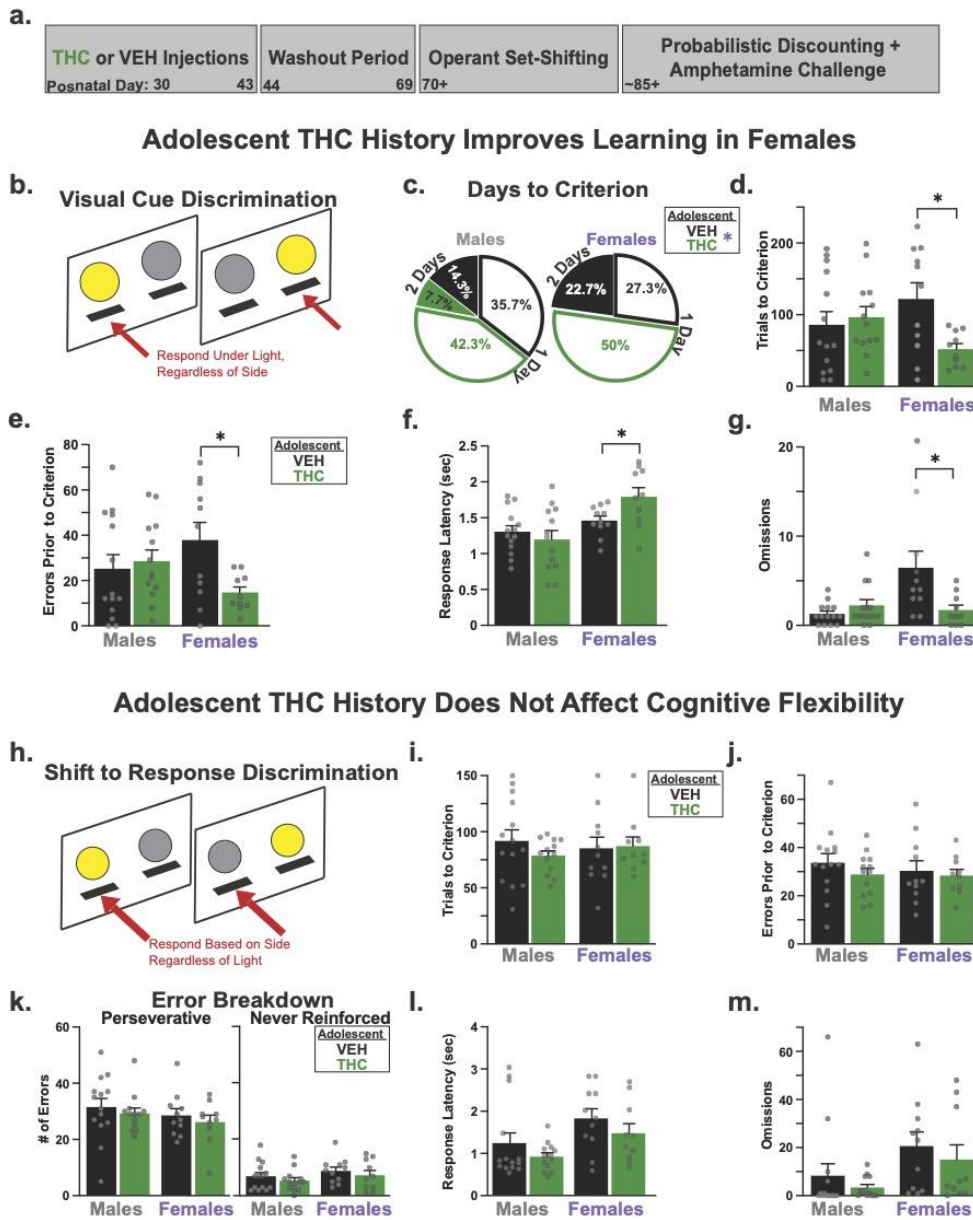


Fig. 1 Adolescent THC history selectively impacts adulthood learning and cognition **a)** Experimental timeline. **b)** Schematic of the visual cue discrimination task. **c)** AdoTHC rats (green wedges) were more likely than AdoVEH rats (black wedges) to acquire the visual cue discrimination task to criterion in only one training session (unfilled wedges), rather than requiring 2 sessions to acquire (filled wedges). Both sexes showed similar patterns. **d, e)** AdoTHC females took fewer trials to meet criterion, and made fewer errors when learning visual cue discrimination than AdoVeh females, no such effects were seen in males. **f, g)** AdoTHC females took longer to respond, and made fewer errors than AdoVEH females during cue discrimination training, without effects in males. **h)** Schematic of the subsequently tested strategy set-shifting task. **i)** AdoTHC did not alter the number of trials to learn the new rule to criterion in either sex. Likewise, AdoTHC did not alter in either sex **j)** the number of errors, **k)** the types of errors, **l)** latency to respond, or **m)** omitted trials during set shifting training. Individual rats shown as grey dots in each graph: AdoVEH ($n = 14$ males, 11 females), AdoTHC ($n = 13$ males, 10 females). $\chi^2 p^* < 0.05$ and repeated measure two-way ANOVA, Sidak post hoc: $p^* < 0.05$. Data presented as mean + SEM.

278 **Effects of Adolescent THC on Probabilistic Discounting in Each Sex:**

279 *Acquisition of Probabilistic Discounting:* The same rats were next trained on a probabilistic discounting task for 21 days. All
 280 rats had acquired stable performance by the last three days of training (no block X day interaction: $F_{(5,498,241.89)} = 1.70$, $p =$
 281 0.13). No effect of AdoTX ($F_{(1,44)} = 0.82$, $p = 0.37$), Sex ($F_{(1,44)} = 0.12$, $p = 0.74$), or AdoTX x Sex interactions ($F_{(1,44)} = 0.59$,
 282 $p = 0.45$) were found, suggesting that rats of both sexes and AdoTX histories acquired the discounting task at similar rates.

283 *Stable Probabilistic Discounting:* Following training, all rats exhibited stable and comparable performance on the task by

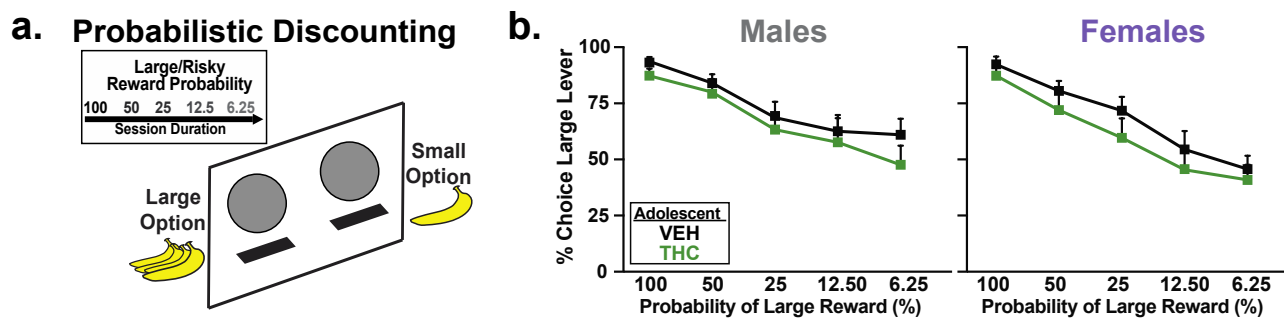


Fig. 2 AdoTHC history does not affect baseline probabilistic discounting **a)** Schematic of the probabilistic discounting task. Sessions consisted of 5 training blocks, in which 2 levers are presented, and rats must choose either a small reward lever that always delivers one pellet, or a large reward-delivering lever which becomes increasingly unlikely to deliver any reward as the session progresses. **b)** Data is shown for males and females, indicating that AdoTHC rats did not differ from their AdoVEH counterparts in probabilistic discounting, with both shifting from nearly exclusively preferring the large reward lever, but appropriately shifting away from it as it became less likely to deliver reward. AdoVEH ($n = 14$ males, 11 females), AdoTHC ($n = 15$ males, 8 females). Data represented as average within each probability block from three consecutive days of stable performance. Mean \pm SEM.

284 the last three days, with decreasing choice of the high-reward lever as delivery of this reward became increasingly unlikely
 285 or “risky” (main effect of Block: $F_{(4,176)} = 75.01$, $p < 0.001$). No effect of AdoTX ($F_{(1,44)} = 0.85$, $p = 0.36$), Sex ($F_{(1,44)} = 0.10$,
 286 $p = 0.75$), or AdoTX x Sex interactions ($F_{(1,44)} = 0.62$, $p = 0.44$) were found, suggesting that rats of both sexes and AdoTX
 287 histories displayed comparable levels of risky choice across blocks (**Fig. 2b**). Likewise, neither AdoTX nor Sex affected win
 288 stay or lose shift choice strategies (no main effect of AdoTX: $F_{(1,44)} < 1.64$, $ps > 0.21$; no main effect of Sex $F_{(1,44)} < 0.55$,
 289 $ps > 0.46$; no AdoTX x Sex $F_{(1,44)} < 1.51$, $ps > 0.23$; data not shown).

290
 291 With respect to other performance measures, AdoTX did not affect choice latencies (no AdoTx: $F_{(1,44)} = 0.16$, $p = 0.69$; no
 292 AdoTX X Sex interaction: $F_{(1,44)} = 0.129$, $p = 0.26$). Additionally, we found that AdoVEH rats omitted more on the “riskier”
 293 probability blocks compared to AdoTHC animals (AdoTX x Sex x Block: $F_{(4,176)} = 3.95$, $p = 0.004$, AdoTX x Block: $F_{(4,176)} =$
 294 4.39 , $p = 0.002$; no AdoTx: $F_{(1,44)} = 0.94$, $p = 0.34$), an effect that was particularly apparent in females (AdoTX x Block: $F_{(4,$
 295 $68)} = 3.19$, $p = 0.02$; data not shown). Overall, we saw that females took longer to respond than males (Sex: $F_{(1,44)} = 40.03$,

$p < 0.0001$) and omitted more trials ($F_{(1,44)} = 63.83, p < 0.0001$), especially in low-probability training blocks at the end of the session (Sex x Block: Latency: $F_{(4,176)} = 4.57, p = 0.002$; Omissions: $F_{(4,176)} = 26.57, p < 0.0001$).

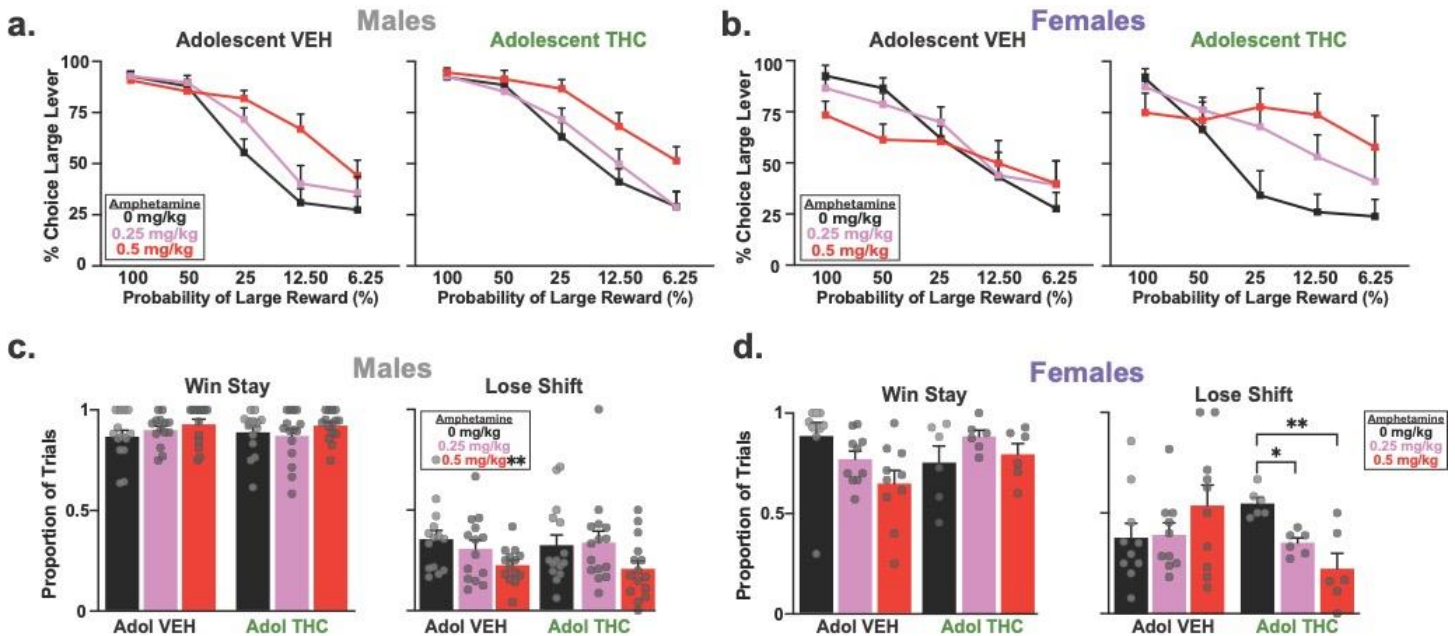


Fig. 3 Amphetamine-induced 'risky' responding is potentiated after adolescent THC. **a)** Data from saline (black line/bar), low dose (0.25mg/kg; pink line/bar) and high dose (0.5mg/kg; red line/bar) amphetamine tests in each sex are shown. When performance patterns were further interrogated, we found that high dose of AMPH in Males increased inflexibility across both AdoTX groups, however in **b)** females, the high dose of AMPH in AdoVEH rats reduced risky responding at high probability blocks not seen in AdoTHC females. **c)** AMPH did not alter win-stay, but reduced lose-shift in AdoTX males. **d)** In AdoVEH females, AMPH decreased win-stay and increased lose-shift, while in AdoTHC AMPH had no effect on win-stay but decreased lose-shift. AMPH = amphetamine. AdoVEH ($n = 14$ males, 10 females), AdoTHC ($n = 15$ males, 6 females). Repeated measure three-way ANOVA; Sidak post hoc: $p^* < 0.05, p^{**} < 0.01$. Data represented as mean + SEM, individual animals shown as grey dots.

Effects of Acute Amphetamine on Probabilistic Discounting: As previously reported (St. Onge et al. 2010), amphetamine increased choice of the large/risky option in a dose dependent manner when analyzed across both groups and sexes (**Fig. 3a**, amphetamine x Block interaction: $F_{(8,328)} = 10.02, p < 0.0001$; main effect of amphetamine: $F_{(2,82)} = 9.88, p < 0.0001$). However, the effects of different doses of amphetamine varied as a function of AdoTX treatment and sex. Analysis of the choice data also revealed significant amphetamine x AdoTX ($F_{(2,82)} = 3.33, p = 0.04$) and amphetamine x AdoTX x Sex interactions ($F_{(2,82)} = 3.20, p = 0.05$). Partitioning this latter interaction by AdoTX group revealed that for control rats, amphetamine exerted different effects on choice in males vs females (amphetamine x Sex: $F_{(2,44)} = 3.73, p = 0.03$). Post-hoc comparisons showed that, in males, amphetamine increased risky choice following treatment with the 0.5 mg/kg dose ($p < 0.02$) but not the lower, 0.25 mg/kg dose ($p = 0.19$). In contrast, the 0.5 mg/kg dose had more deleterious effects on choice in females, reducing risky choice in the high probability blocks and increasing it in the lower ones, while the 0.25 mg/kg dose induced a modest increase in risky choice in the latter block. This yielded an overall lack of effect of amphetamine in control females (**Fig. 3b**, amphetamine: $F_{(2,18)} = 0.57, p = 0.57$). On the other hand, amphetamine induced a more reliable and pronounced increases in risky choice in both males and females AdoTHC animals, with analysis of

311 these data producing significant main effects of amphetamine treatment (**Fig. 3a,b**, $F_{(2,38)} = 12.07$, $p < 0.0001$) in the absence
312 of an interaction with the sex factor ($F_{(2,38)} = 1.56$, $p = 0.22$). When collapsed across sex, post hoc comparisons showed that
313 both the 0.25 and 0.5 mg/kg dose increase risky choice (both $ps < 0.05$), although inspection of Fig. 3b indicates that the
314 effect of the 0.25 mg/kg dose was driven primarily by females. From these data, we conclude that AdoTHC treatment makes
315 rats more sensitive to the ability of amphetamine to increase risky choice, and this effect appears to be more prominent in
316 females.

317
318 Subsequent analyses examined how amphetamine alters sensitivity to recent rewarded or non-rewarded choices by
319 comparing win-stay and lose shift ratios. Analysis of the win-stay data yielded a significant amphetamine x AdoTX x Sex
320 interaction (**Fig. 3c,d**, $F_{(2,82)} = 4.34$, $p = 0.02$). This was driven by the fact that in control rats, the 0.5 mg/kg dose resulted in
321 lower win-stay values in females vs males ($p < 0.01$), although neither group showed significant changes in these values
322 relative to saline (both $F_s < 2.5$, both $ps > 0.10$). Win-stay behavior was unaltered in AdoTHC rats (all $F_s < 2.4$, all $p > 0.10$).
323 In contrast, amphetamine had more pronounced effects on sensitivity to reward omissions, as indexed by changes in lose
324 shift behavior. The analyses here revealed a significant amphetamine x AdoTX interaction ($F_{(2,38)} = 4.84$, $p = 0.01$) and a
325 three-way interaction with the sex factor ($F_{(2,38)} = 5.15$, $p = 0.01$). In controls, amphetamine reduced lose shift behavior in
326 males, but actually increased it in females (**Fig. 3c,d**, amphetamine x Sex: $F_{(2,44)} = 3.86$, $p = 0.03$), whereas in AdoTHC rats,
327 these treatments uniformly reduced lose-shift behavior across sexes (amphetamine: $F_{(2,38)} = 8.87$, $p < 0.001$; amphetamine
328 x Sex: $F_{(2,38)} = 2.61$, $p = 0.08$). From these data, we conclude that AdoTHC treatment makes rats more sensitive to the
329 ability of amphetamine to increase risky choice and reduce sensitivity to losses, and this effect appears to be more prominent
330 in females.

331
332 With respect to other performance measures, amphetamine increased choice latency and number of omissions across all
333 probability blocks (amphetamine: $F_{s(2,82)} < 15.19$, $ps < 0.001$, no amphetamine x Block interaction: $F_{s(8,328)} < 0.98$, $ps >$
334 0.45). Analysis of the latency data revealed a significant amphetamine x AdoTX x Block ($F_{(8,328)} = 2.27$, $p = 0.02$) and
335 amphetamine x AdoTX x Sex x Block interaction ($F_{(2,82)} = 2.67$, $p = 0.01$). Partitioning this latter interaction by AdoTX group
336 revealed that in AdoVEH rats, females took longer to respond than males across the session (Sex x Block: $F_{(4,88)} = 3.02$, p
337 $= 0.02$, Sex: $F_{(1,22)} = 44.13$, $p < 0.001$). In AdoTHC rat, females took longer to respond compared to males (Sex: $F_{(1,19)} =$
338 21.44 , $p < 0.001$; no Sex X Block: $F_{(4,76)} = 2.41$, $p = 0.06$). Further analysis of the omission data revealed that overall females
339 omitted more on trials compared to males (Sex x Block: $F_{(4,164)} = 12.93$, $p < 0.001$; Sex: $F_{s(1,41)} < 71.39$, $p < 0.001$).

341 Chemogenetic Dopamine Neuron Stimulation During Probabilistic Discounting

342 The above experiments showed that AdoTHC enhances visual cue learning in females but had few other effects on cognitive
343 flexibility or probabilistic discounting under basal conditions. However, when treated with the monoamine-enhancing drug
344 amphetamine, we found evidence for stronger enhancement of “risky” responding in AdoTHC rats, relative to AdoVEH
345 controls. We therefore next asked whether this effect relates to changes in the functions of VTA dopamine neurons in
346 particular, by using Gq-coupled DREADDs to acutely stimulate VTA dopamine neurons or VTA dopamine neuron projections
347 to mPFC in rats with both AdoTX histories.

348
349 *Initial Training:* Rats in this experiment did not undergo strategy set shifting training prior to probabilistic discounting training,
350 so we confirmed that AdoTX again did not affect initial acquisition of instrumental food seeking behavior during initial training
351 (no main effect of AdoTX: $F_{S(1,46)} < 0.78$, $ps > 0.38$). We also confirmed that genotype (Geno: TH:Cre+ or WT littermate)
352 did not alter instrumental training ($F_{S(1,46)} < 2.55$, $ps > 0.12$; No AdoTX x Geno interaction: $F_{S(1,46)} < 0.63$, $ps > 0.43$).

353
354 *Probabilistic Discounting Training:* Likewise, all included rats (TH:Cre and WT) successfully learned the discounting task
355 (Main effect of Block: $F_{(4,184)} = 129.26$, $p < 0.001$, no main effects of, or interactions involving AdoTX or Geno: $F_{S(1,46)} < 0.88$,
356 $ps > 0.35$). Additionally, AdoTX nor genotype affected latency, omissions, or win-stay/lose-shift choice strategies (no main
357 effects of, or interactions involving AdoTX or Geno: $F_{S(1,46)} < 3.34$, $ps > 0.07$).

359 **Impact of Acutely Activating VTA Dopamine Neurons on Probabilistic Discounting**

360 Despite the robust changes in locomotion and reward seeking that is seen following chemogenetic VTA dopamine neuron
361 manipulations in TH:Cre rats (Boekhoudt et al. 2016; Boekhoudt et al. 2018; Halbout et al. 2019; Lawson et al. 2023; Mahler
362 et al. 2019; Runegaard et al. 2018), we found few effects of VTA dopamine neuron stimulation on the probabilistic
363 discounting task. First, we looked at VTA dopamine neuron stimulation in only TH:Cre rats. All rats displayed normal
364 discounting profiles (TH:Cre+: main effect of Block: $F_{(4,84)} = 105.93$, $p < 0.001$), and as in the prior experiment, no effect of
365 AdoTX alone was seen on discounting (AdoTX x Block: $F_{(4,84)} = 0.91$, $p = 0.46$). Moreover, when we tested the effects of
366 systemic CNO to activate DA neurons, we did not observe any effects of this treatment in either control or AdoTX groups
367 (**Fig. 4e**, no main effect of, or interactions involving AdoTX or CNO: $F_s < 2.27$, $ps > 0.07$). Additionally, we did not see any
368 effects of CNO, AdoTX, nor interactions therein on choice latency, omissions, or win-stay/lose-shift choice strategy ($F_{S(1,21)}$
369 < 3.00 , $ps > 0.10$; data not shown). These findings are inconsistent with our hypothesis that increased excitability of
370 mesolimbic dopamine neurons would be sufficient to recapitulate the ability of amphetamine to increase risky/perseverative
371 responding in either control or in AdoTHC-experienced rats.

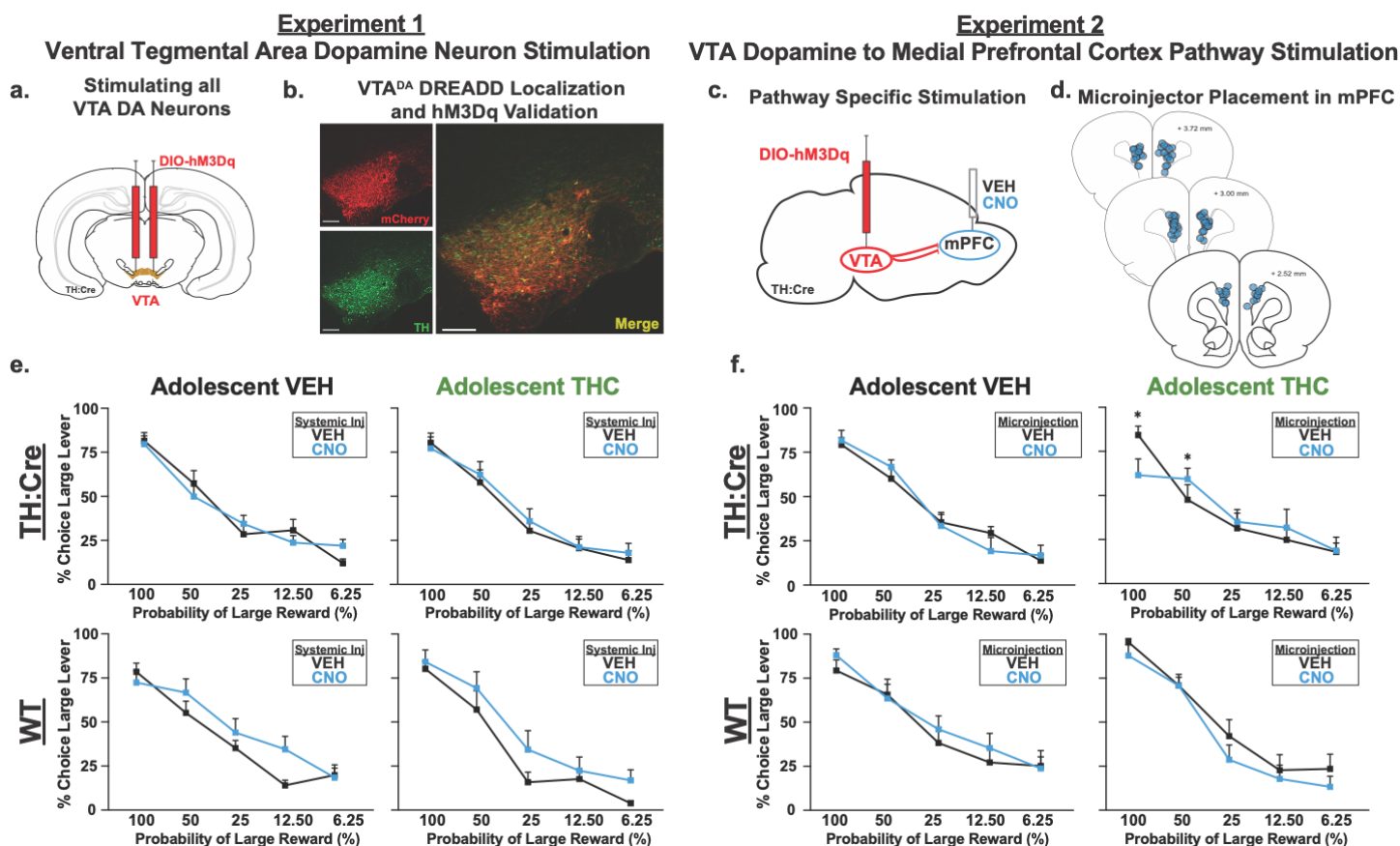


Fig. 4 Stimulation of VTA dopamine neurons, or VTA dopamine projections to mPFC does not affect probabilistic discounting a) Bilateral Cre-dependent hM3Dq DREADD AAV injections were made in VTA of TH:Cre rats, and of wildtype (WT) littermates. b) Example hM3Dq DREADD expression (red) is localized to tyrosine hydroxylase+ (TH; green) neurons within VTA (yellow=merge). Scale bar, 300 μ m. c) For pathway-specific stimulation of VTA dopamine projections to mPFC, Cre-dependent hM3Dq DREADDs were injected into VTA as in Experiment 1, and cannulae targeting mPFC allowed CNO microinjection (1mM, 0.5 μ l) upon DREADD-expressing dopamine neuron axons in this pathway. d) Cannula placements of each rat in pathway stimulation Experiment 2 is shown. e) Neither VTA dopamine neuron stimulation induced by systemic CNO in TH:Cre rats, nor f) stimulation of the VTA dopamine projection to mPFC induced by mPFC CNO microinjections in TH:Cre rats robustly altered probabilistic discounting in AdoTHC or AdoVEH rats. Likewise, lower panels of e,f indicate that CNO did not have robust effects on WT rats without DREADDs. Data is represented as average % choice of the large reward lever in each probability block across the two CNO, and two VEH tests conducted in each rat, mean \pm SEM, $p^* < 0.05$. TH:Cre+: VTA dopamine stimulation; AdoVEH ($n = 8M, 3F$) AdoTHC ($n = 8M, 4F$); VTA dopamine to mPFC: AdoVEH ($n = 10M, 4F$) AdoTHC ($n = 9M, 5F$). Wildtype: VTA dopamine stimulation; AdoVEH ($n = 5M, 3F$) AdoTHC ($n = 5M$); VTA dopamine to mPFC: AdoVEH ($n = 9M, 4F$) AdoTHC ($n = 8M$).

372

373

Impact of Selectively Activating VTA Dopamine Projections to mPFC

374

We next asked whether selectively stimulating VTA dopamine neuron projections to mPFC would recapitulate the potentiation of amphetamine effects in AdoTHC rats. We did so by locally applying CNO upon DREADD-expressing axons of VTA dopamine neurons in mPFC. We have shown that this pathway-specific stimulation approach is capable of potentiating both axonal dopamine release and motivated reward seeking (Halbout et al. 2019; Mahler and Aston-Jones 2012; Mahler et al. 2019). We found few effects of this manipulation on probabilistic discounting. On the probabilistic discounting task, "risky" responding across the session varied across CNO and AdoTX (Fig. 4; AdoTX x CNO x Block interaction: $F_{(4,104)} = 2.99$, $p = 0.02$; no main effect of AdoTX: $F_{(1,26)} = 0.15$, $p = 0.70$; no main effect of CNO: $F_{(1,26)} = 0.000$

380

381 $p = 0.10$), in AdoVEH animals there were no measurable effects of or interactions with CNO (all $F_s < 1.80$, $p_s > 0.14$).
382 However, in AdoTHC rats CNO modestly reduced risky responding at the 100% probability block followed by an increase
383 at the 50% probability block (CNO x Block: $F_{(4,52)} = 2.99$, $p = 0.03$). Rats performed similarly on choice latency, omissions,
384 and win-stay/lose-shift regardless of AdoTX, CNO, or interactions therein with block (All $F_s < 2.00$, $p_s > 0.10$).

386 Minimal DREADD-independent Effects of CNO

387 Neither systemic nor intra-mPFC CNO had major behavioral effects in WT rats lacking DREADD expression (**Fig. 4e,f**).
388 Systemic CNO in WT rats seemed to promote a more risk-prone phenotype, based on increased preference for the risky
389 lever across all blocks (Main effect of CNO in WT rats: $F_{(1,11)} = 9.42$, $p = 0.01$; no interactions of CNO, AdoTX, or Block:
390 $F_{s(4,44)} < 2.04$, $p_s > 0.11$), but did not affect response latency, omissions, or win-stay/lose-shift choice strategies ($F_{s(1,11)} <$
391 2.96 , $p_s > 0.11$). Intra-mPFC CNO in WT rats altered risk responding such that “risky” choice was decreased in AdoTHC
392 rats, whereas it was increased in AdoVEH animals (AdoTX x CNO: $F_{(1,16)} = 5.40$, $p = 0.03$). These effects were driven by
393 changes in reward sensitivity, as CNO differentially affected win-stay behavior in the two groups ($F_{(1,160)} = 5.78$, $p = 0.03$; no
394 main effect of CNO $F_{(1,16)} = 0.73$, $p = 0.41$). Thus, CNO in AdoTHC WT rats were less likely to follow a risky win with another
395 risky choice, while CNO had the opposite effects in AdoVEH WT (data not shown). Additionally, we saw that AdoTHC
396 animals had lower risky responding in the higher probability blocks, that then increased in the latter probability blocks
397 (AdoTX x Block: $F_{(4,64)} = 2.52$, $p = 0.05$). We did not see any effect of AdoTX or CNO on lose-shift strategies (CNO x AdoTX:
398 $F_{(1,16)} = 0.67$, $p = 0.43$; no main effect of CNO $F_{(1,16)} = 1.71$, $p = 0.21$) or response latency and omissions ($F_{s(1,16)} < 3.74$, p_s
399 > 0.07 ; data not shown).

401 Discussion

402 Here we show that administration of a well-characterized, human-relevant dose of THC (Ruiz et al. 2021a; Torrens et al.
403 2022; Torrens et al. 2020) during adolescence has subtle effects on behavioral tests of instrumental learning, without
404 measurably altering performance on mPFC dopamine-dependent strategy set shifting or risk/reward decision making
405 assessed with a probabilistic discounting task. However, when monoamine signaling was acutely enhanced with systemic
406 amphetamine, an underlying effect of AdoTHC history was revealed. Relative to AdoVEH controls, amphetamine in AdoTHC
407 rats were more sensitive to the ability of lower doses of amphetamine to increase perseverative responding for a large
408 reward option when this choice was unlikely to result in reward. However, this potentiation of amphetamine effects in
409 AdoTHC rats was not recapitulated by more specific chemogenetic stimulation of VTA dopamine neurons, or of their
410 projections to mPFC in particular. This could suggest non-VTA dopaminergic mechanisms underlying potentiation of this

411 cognitive effect of amphetamine in AdoTHC rats, suggesting potential impacts of THC on adolescent development of other
412 systems upon which amphetamine acts. Alternatively, the enhanced effect of amphetamine reported here may be driven by
413 alterations at the level of the dopamine terminal rather than changes in dopamine cell excitability, as dopamine receptor
414 antagonism is able to block this amphetamine-induced risky responding (St. Onge and Floresco 2009). Further, we
415 thoroughly characterize sex differences in decision-making across these behavioral tasks, some of which mediate the
416 persistent impacts of AdoTHC on behavior. Results open new directions for investigating long-term impacts of AdoTHC on
417 non-VTA dopaminergic modulation of cognition, and may inform associational studies of the long-term impacts of adolescent
418 cannabis use in humans.

419 420 **Adolescent THC Exposure Enhances Initial Discrimination Learning**

421 AdoTHC rats learned a visually cued instrumental response rule quicker than AdoVEH-treated controls, and this effect was
422 most robust in females. This finding adds to the literature on potentially pro-learning/cognitive effects of AdoTHC (Hernandez
423 et al. 2021; Stringfield and Torregrossa 2021b). This said, previous studies using other adolescent cannabinoid exposure
424 models have not found analogous increases in initial rule discrimination learning (Freels et al. 2024; Gomes et al. 2015;
425 Hernandez et al. 2021), though this might be due to the fact that few studies included female subjects, and the THC
426 administration protocols employed were quite different.

427
428 We found no clear effects of AdoTHC across our set shifting performance metrics, including trials to criterion, number of
429 errors made, or error types on the set shift day. These findings are consistent with others that found no major AdoTHC-
430 induced changes in cognitive flexibility, as quantified in with different set shifting tasks, including the intra/extra dimensional
431 “digging” task and an operant-based task similar to the one used here (Gomes et al. 2015; Hernandez et al. 2021; Poulia
432 et al. 2021). However, in a study utilizing the same strategy set-shifting task as used here, females exposed to self-
433 administered cannabis extract vapor during adolescence took longer to learn the new rule, and made more errors on the
434 set-shift day, though this deficit was not seen after experimenter-administered cannabis vapor (Freels et al. 2024). It is
435 presently unclear whether differences in patterns of results reflect differences in route of THC administration or dose, the
436 specific timing of THC exposure during adolescence, or other experimental details. As such, this remains an important topic
437 for future research intended to probe how adolescent cannabinoid exposure may alter executive functioning.

438
439 Though untested here, it is possible that AdoTHC altered other related cognitive processes such as transitioning between
440 tasks, mental sets, or rule structures, which depend upon more lateral PFC subregions such as orbitofrontal cortex (OFC;
441 Birrell and Brown 2000; Floresco et al. 2008; McAlonan and Brown 2003). For example, adolescent pubertal administration

of the potent CBR agonist WIN 55,212 alters OFC-dependent reversal learning in and attentional set-shifting task (Gomes et al. 2015), though adolescent vaporized cannabis or cannabis smoke did not impact this same type of reversal learning (Freels et al. 2024; Hernandez et al. 2021). Again, discrepancies between studies may reflect differences in effects of cannabinoid drugs, doses, exposure timing, and washout period; further underscoring the need for a consistent, rationally-designed AdoTHC exposure model in the field—we argue that the model used here is the best-characterized to date in the field (Halbout et al. 2023; Lee et al. 2022; Lee et al. 2024; Lin et al. 2023; Ruiz et al. 2021a; Torrens et al. 2022; Torrens et al. 2020). This said, the possibility that OFC is even more sensitive to disruption by AdoTHC than mPFC should be directly tested in future studies.

Adolescent THC Does Not Alter Basal Probabilistic Discounting

In the mPFC-dependent probabilistic discounting task (St. Onge and Floresco 2010), rats choose between a small reward that is always delivered when chosen (1 palatable banana pellet), and a larger reward that becomes increasingly unlikely to be delivered over the course of the ~1h session (providing either 4 or 0 pellets). Efficient performance on this task demands evaluation of both risk and opportunity, and is dependent on intact functioning of both the mPFC and mesocortical and mesoaccumbens dopamine transmission (Jenni et al. 2017; St. Onge et al. 2012; St. Onge et al. 2010; Stopper et al. 2013). Impaired PFC activity results in deficits in adjusting choice in response to changes in reward probabilities, loss assessment, and a diminished ability to appropriately compare and favor larger rewards even when the probability of receiving them is higher (Bercovici et al. 2023; St. Onge and Floresco 2010). We found no major impacts of AdoTHC history on acquisition of, or stable performance on this task, as measured by choices of the ‘risky’ lever, choice strategies following rewarded vs unrewarded trials, latency to decide, and decisions to omit trials. One prior study (Jacobs-Brichford et al. 2019) found that administration of WIN 55,212 in both sexes during adolescence elevated preference for the ‘risky’ lever at lower reward probabilities, which we did not see in either sex as a result of our AdoTHC exposure model. Though several other differences exist between this study and the present one, we note that several prior reports have also shown more severe lasting effects of synthetic CBR agonists versus THC when administered in adolescents (Higuera-Matas et al. 2015; Renard et al. 2014; Stringfield and Torregrossa 2021a).

Adolescent THC History Potentiates Amphetamine Effects on Probabilistic Discounting

Dopamine markedly influences mPFC-dependent cognition (Floresco and Magyar 2006; Goldman-Rakic 1995; Goto et al. 2007; Seamans and Yang 2004), including in the probabilistic discounting task employed here (Floresco and Whelan 2009; Islas-Preciado et al. 2020; Jenni et al. 2017; St. Onge et al. 2012; St. Onge et al. 2010; St. Onge and Floresco 2009). In the present study we replicated prior findings that pharmacologically challenging monoamine systems with amphetamine

473 increased perseveration of responding for a large reward in both males and females (Islas-Preciado et al. 2020; St. Onge
474 and Floresco 2009). This change was accompanied by a reduction in lose-shift behaviors following 'risky' losses, indicating
475 that enhancing DA levels attenuates the impact that non-rewarded actions exert over subsequent choice. Interestingly, we
476 saw that amphetamine led to an overall increase in choice latency, and omitted trials. This may have been driven in part by
477 increased psychomotor activity induced by amphetamine that may have displaced animals from the levers, thus delaying
478 choice.

479
480 Moreover, we found that this effect of amphetamine was more potent in AdoTHC, relative to AdoVEH rats, especially in
481 females. At the highest dose of amphetamine, AdoTHC rats chose the 'risky' lever more when probabilities of that reward
482 were low. These effects were also markedly sex-dependent. In males, amphetamine reduced overall lose-shift behavior. In
483 females, the highest dose of amphetamine induced a disruptive reduction in win-stay behavior only in AdoVEH females,
484 whereas it reduced lose-shift behaviors only in AdoTHC females, contributing to the heightened increased 'risky' responding
485 observed in the AdoTHC groups. While we refer to this behavior as 'risky' responding (St. Onge et al. 2010), this behavior
486 may also be interpreted as amphetamine causing AdoTHC animals to be less adaptive to changing contingencies, or more
487 liable to perseverate on the initially prepotent large reward lever. Regardless, these results suggest that although
488 probabilistic discounting under basal conditions is not altered by AdoTHC, underlying differences were nonetheless revealed
489 upon acute activation of monoaminergic signaling. Amphetamine blocks and reverses the transporter for dopamine,
490 norepinephrine, serotonin, and changes in one or more of these systems could be responsible for the potentiated response
491 to amphetamine we see in this task in AdoTHC rats (especially females).

493 **Effects of Chemogenetic VTA Dopamine Neuron, and mPFC Dopamine Projection Stimulation on Probabilistic** 494 **Discounting**

495 In our next experiments, we sought to determine whether, as predicted, changes in VTA dopamine neuron signaling are
496 responsible for the potentiated effect of amphetamine in AdoTHC rats. We therefore prepared transgenic TH:Cre rats with
497 excitatory DREADDs, allowing stimulation of either all VTA dopamine neurons, or of dopamine neuron projections to mPFC
498 in particular. Surprisingly, no major effects of chemogenetically stimulating all VTA dopamine neurons were found on
499 probabilistic discounting. In this regard, it bears mentioning that pharmacological stimulation of dopamine D2 or D1 receptors
500 in the nucleus accumbens (a main target of the VTA dopamine projection) either had no effect on risky choice or led to more
501 optimal decision making, respectively- an effect distinct from that induced by amphetamine (Stopper et al. 2013).
502 Furthermore, when we stimulated VTA dopamine neuron projections to mPFC, this did not markedly affect probabilistic
503 discounting in AdoVEH rats, which is similar to a prior finding examining manipulations of VTA dopamine projections to PFC

(Verharen et al. 2018). On the other hand, chemogenetic stimulation of mPFC dopamine terminals in AdoTHC rats attenuated risky responding during the 100% probability block, but then increased it in the subsequent 50% block. This effect loosely resembles that induced by intra-PFC D₂ receptor stimulation on probabilistic discounting, which flattened the discounting curve (St. Onge et al. 2011). This is consistent with the possibility that AdoTHC exposure causes a slight enhancement in mesocortical dopamine D₂ receptor signaling—a possibility that should be directly investigated. Moreover, no other differential effects of DREADD stimulation were seen in AdoTHC versus AdoVEH groups. The absence of significant effects in this experiment may also relate to the relatively small number of TH:Cre females, especially since amphetamine effects on the same behaviors seem to be driven by females. Additionally, the lack of behavioral effects of chemogenetic stimulation is unlikely to have resulted from inefficient DREADD stimulation or infection. Extent of viral expression, and selectivity of expression in TH+ VTA neurons was equivalent to our prior reports using these transgenic rats and viral vectors. In these reports we have shown this DIO-hM3Dq vector has >97% selectivity to TH+ neurons, and it is capable of driving dopamine neuron firing rates in vitro, increasing the number of them active in vivo, and increasing cFos expression in them (Mahler et al. 2019). Accordingly, we and others have found very robust behavioral effects of VTA dopamine neuron stimulation in a range of tasks, including cocaine intake and seeking, food reward intake, locomotion, and motivation (Boekhoudt et al. 2016; Boekhoudt et al. 2018; Halbout et al. 2019; Lawson et al. 2023; Mahler et al. 2019; Runegaard et al. 2018). We also previously confirmed that pathway-specific stimulation of VTA dopamine projections enhances dopamine release, and stimulation of mPFC projections is behaviorally efficacious for stimulating cue-induced cocaine seeking as well (Mahler et al. 2019). Therefore, it was particularly striking that we did not see any notable behavioral effects of dopamine neuron or dopamine projection stimulations on probability discounting, despite the fact that this task is sensitive to mPFC dopamine receptor manipulations. This may imply that alterations of VTA and non-VTA dopamine caused by amphetamine might simply be qualitatively distinct from manipulations of VTA dopamine neuron activity and dopamine release, as were conducted here chemogenetically (Mahler et al. 2019; Mahler et al. 2014). In this regard, performance of this probabilistic discounting task is associated with fluctuations in mesocortical and mesoaccumbens dopamine that relate to changes in the amount of rewards received and adjustments in risky choice biases (St. Onge et al, 2012). Amphetamine would cause robust increases in tonic dopamine levels, leading to a more static dopamine signal that impedes flexible adjustments in choice biases, whereas chemogenetic increases in dopamine neuron excitability would be a comparatively more subtle manipulation that could still permit variations in dopamine signaling over a session. Alternatively, amphetamine-induced changes in risk taking seen in AdoTHC rats may not solely depend upon VTA dopamine neurons themselves, but may instead depend upon other neural systems targeted by amphetamine, such as nigrostriatal dopamine projections to the dorsomedial striatum (Schumacher et al. 2021), other non-mPFC targets of VTA dopamine neurons, or other monoamine systems affected by amphetamine. Regardless, this intriguing finding refines our understanding of the neural

535 substrates of probability-based decision making and underscores the need for further exploration to identify the specific
536 mechanisms involved.

538 **Sex Differences**

539 We conducted these studies in female as well as male rats, and found several consistent sex differences, both overall and
540 in interaction with other experimental variables. Consistent with existing literature (Gargiulo et al. 2022; Islas-Preciado et al.
541 2020), we saw sex differences in response characteristics during operant set-shifting as well as probabilistic discounting,
542 with females showing longer deliberation times (response latency), and more omitted trials than males. Strikingly, these
543 specific sex differences consistently surfaced in the probabilistic discounting task across various experiments and cohorts,
544 underscoring the robustness of the response latency and trial omission effects in females. Some of these sex differences
545 may derive from females attaining satiety on the palatable reward quicker than males, but since both omissions and longer
546 latencies at least partially emerged in females early in sessions during otherwise apparently vigorous reward seeking, this
547 finding may also reflect different strategies taken by females relative to males (Chen et al. 2021; Orsini and Setlow 2017).
548 For example, female rats have previously been shown to be more risk-averse than males on average (Orsini et al. 2016),
549 and may exhibit prolonged response times and higher omission rates due to aversion to the risk of losing a reward, as
550 demonstrated by a heightened sensitivity to loss (van den Bos et al. 2012). Alternatively, extended choice latencies and
551 omissions may signify females' tendency to take more time in learning about the probability distribution of the outcomes.
552 This aligns with both rodent and human findings, indicating that females take longer to develop a preference for the more
553 advantageous option when learning about probability distributions of reward versus punishment (van den Bos et al. 2013;
554 van den Bos et al. 2012). Both possibilities warrant further exploration in future.

556 **Limitations**

557 The present report has a number of limitations that should be considered. A single dose of THC (5mg/kg IP) was
558 administered to adolescents daily from postnatal days 30 to 43, and persistent effects of THC are known to be dose-
559 dependent (Amal et al. 2010; Freeman-Striegel et al. 2023; Irimia et al. 2015), and dependent upon the specific
560 developmental stage at which it is experienced (Cha et al. 2006; Gorey et al. 2019; Mokrysz et al. 2016; Murray et al. 2022;
561 Schramm-Sapyta et al. 2007; Torrens et al. 2022; Torrens et al. 2020). The low-dose THC regimen used here may capture
562 the effects of frequent, low-potency cannabis use (Cooper and Haney 2009; Huestis et al. 1992) rather than high-intensity
563 or chronic cannabis exposure, use of high-potency cannabis products, or escalating use over time. Further work should
564 examine how THC dose, dosing pattern, and means of administration (i.e. oral vs intraperitoneal vs inhalation) impact long-

565 term function of dopamine-dependent cognition. Also relevant to dose, we and others often observe more potent behavioral
566 effects of THC in female, relative to male adolescent rats, which is likely attributable in part to differences in THC metabolism,
567 in particular with regard to a higher blood and brain levels in female adolescent and adult rats and mice of the THC
568 metabolite 11-OH-THC, a CB1 agonist (Baglot et al. 2021; Craft et al. 2019; Ruiz et al. 2021a; Ruiz et al. 2021b; Torrens
569 et al. 2022; Torrens et al. 2020; Tseng et al. 2004; Wiley and Burston 2014), and this sex difference is also seen in humans
570 after cannabis use (Arkell et al. 2022; Matheson et al. 2020; Sholler et al. 2021). This metabolic sex difference may at least
571 in part account for the more robust persistent effects of adolescent THC in females than males seen here, and in numerous
572 prior reports (Cha et al. 2007; Le et al. 2022; Rubino and Parolaro 2011; Rubino et al. 2009; Ruiz et al. 2021a; Ruiz et al.
573 2021b; Tseng et al. 2004). We found that in females, AdoTHC had more pronounced effects than in males, including
574 enhancement of visual cue learning, and stronger potentiation of amphetamine's effects on probability discounting
575 behavior—but no effect on other behaviors like set shifting and baseline probability discounting. These results support the
576 increasingly apparent possibility that AdoTHC's effects are both highly sex- and task-dependent. Further studies
577 investigating the reasons why AdoTHC seems to influence some behaviors and not others are needed, and may provide
578 important new evidence relevant to cognitive and anatomical development. Another limitation of this report is the lack of
579 data on complementary cognitive tasks, which are dependent upon other cortical regions like OFC that are notably sensitive
580 to adolescent cannabinoid exposure (Egerton et al. 2005; Gomes et al. 2015; Klugmann et al. 2011), and which are mediated
581 by ECB-dependent processes that may be particularly sensitive to AdoTHC's long-lasting impacts.

582 583 **Summary**

584 In sum, this paper provides novel information about the impact of AdoTHC on the developing brain of males, and especially
585 of females. Results point to long-lasting changes that influence cue learning, and the ability of acute amphetamine to alter
586 risky decision making. A key finding is the pervasive nature of sex-differences—both in behavioral strategies employed by
587 rats in these tasks, and in the severity of long-lasting consequences of AdoTHC.

588

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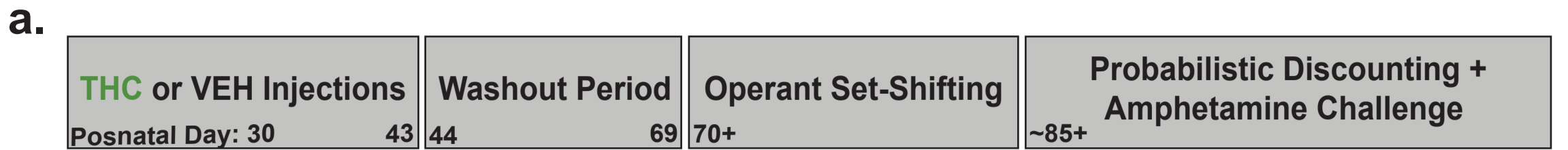
864 Figure Legends

865 **Fig. 1 Adolescent THC history selectively impacts adulthood learning and cognition** **a)** Experimental timeline. **b)**
866 Schematic of the visual cue discrimination task. **c)** AdoTHC rats (green wedges) were more likely than AdoVEH rats (black
867 wedges) to acquire the visual cue discrimination task to criterion in only one training session (unfilled wedges), rather than
868 requiring 2 sessions to acquire (filled wedges). Both sexes showed similar patterns. **d, e)** AdoTHC females took fewer trials
869 to meet criterion, and made fewer errors when learning visual cue discrimination than AdoVeh females, no such effects
870 were seen in males. **f, g)** AdoTHC females took longer to respond, and made fewer errors than AdoVEH females during
871 cue discrimination training, without effects in males. **h)** Schematic of the subsequently tested strategy set-shifting task. **i)**
872 AdoTHC did not alter the number of trials to learn the new rule to criterion in either sex. Likewise, AdoTHC did not alter in
873 either sex **j)** the number of errors, **k)** the types of errors, **l)** latency to respond, or **m)** omitted trials during set shifting training.
874 Individual rats shown as grey dots in each graph: AdoVEH ($n = 14$ males, 11 females), AdoTHC ($n = 13$ males, 10 females).
875 $X^2 p^* < 0.05$ and repeated measure two-way ANOVA, Sidak post hoc: $p^* < 0.05$. Data presented as mean + SEM.
876

877 **Fig. 2 AdoTHC history does not affect baseline probabilistic discounting** **a)** Schematic of the probabilistic discounting
878 task. Sessions consisted of 5 training blocks, in which 2 levers are presented, and rats must choose either a small reward
879 lever that always delivers one pellet, or a large reward-delivering lever which becomes increasingly unlikely to deliver any
880 reward as the session progresses. **b)** Data is shown for males and females, indicating that AdoTHC rats did not differ from
881 their AdoVEH counterparts in probabilistic discounting, with both shifting from nearly exclusively preferring the large reward
882 lever, but appropriately shifting away from it as it became less likely to deliver reward. AdoVEH ($n = 14$ males, 11 females),
883 AdoTHC ($n = 15$ males, 8 females). Data represented as average within each probability block from three consecutive days
884 of stable performance. Mean \pm SEM.
885

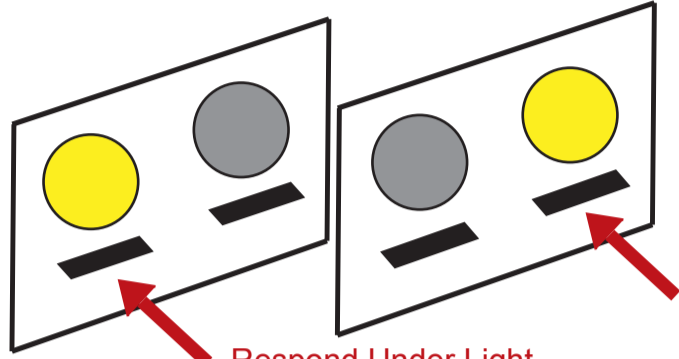
886 **Fig. 3 Amphetamine-induced 'risky' responding is potentiated after adolescent THC.** **a)** Data from saline (black
887 line/bar), low dose (0.25mg/kg; pink line/bar) and high dose (0.5mg/kg; red line/bar) amphetamine tests in each sex are
888 shown. When performance patterns were further interrogated, we found that high dose of AMPH in Males increased
889 inflexibility across both AdoTX groups, however in **b)** females, the high dose of AMPH in AdoVEH rats reduced risky
890 responding at high probability blocks not seen in AdoTHC females. **c)** AMPH did not alter win-stay, but reduced lose-shift in
891 AdoTX males. **d)** In AdoVEH females, AMPH decreased win-stay and increased lose-shift, while in AdoTHC AMPH had no
892 effect on win-stay but decreased lose-shift. AMPH = amphetamine. AdoVEH ($n = 14$ males, 10 females), AdoTHC ($n = 15$
893 males, 6 females). Repeated measure three-way ANOVA; Sidak post hoc: $p^* < 0.05$, $p^{**} < 0.01$. Data represented as mean
894 + SEM, individual animals shown as grey dots.
895

896 **Fig. 4 Stimulation of VTA dopamine neurons, or VTA dopamine projections to mPFC does not affect probabilistic**
897 **discounting** **a)** Bilateral Cre-dependent hM3Dq DREADD AAV injections were made in VTA of TH:Cre rats, and of wildtype
898 (WT) littermates. **b)** Example hM3Dq DREADD expression (red) is localized to tyrosine hydroxylase+ (TH; green) neurons
899 within VTA (yellow=merge). Scale bar, 300 μ m. **c)** For pathway-specific stimulation of VTA dopamine projections to mPFC,
900 Cre-dependent hM3Dq DREADDs were injected into VTA as in Experiment 1, and cannulae targeting mPFC allowed CNO
901 microinjection (1mM, 0.5 μ l) upon DREADD-expressing dopamine neuron axons in this pathway. **d)** Cannula placements of
902 each rat in pathway stimulation Experiment 2 is shown. **e)** Neither VTA dopamine neuron stimulation induced by systemic
903 CNO in TH:Cre rats, nor **f)** stimulation of the VTA dopamine projection to mPFC induced by mPFC CNO microinjections in
904 TH:Cre rats robustly altered probabilistic discounting in AdoTHC or AdoVEH rats. Likewise, lower panels of **e,f)** indicate
905 that CNO did not have robust effects on WT rats without DREADDs. Data is represented as average % choice of the large
906 reward lever in each probability block across the two CNO, and two VEH tests conducted in each rat, mean \pm SEM, $p^* <$
907 0.05. TH:Cre+: VTA dopamine stimulation; AdoVEH ($n = 8$ M, 3F) AdoTHC ($n = 8$ M, 4F); VTA dopamine to mPFC: AdoVEH
908 ($n = 10$ M, 4F) AdoTHC ($n = 9$ M, 5F). Wildtype: VTA dopamine stimulation; AdoVEH ($n = 5$ M, 3F) AdoTHC ($n = 5$ M); VTA
909 dopamine to mPFC: AdoVEH ($n = 9$ M, 4F) AdoTHC ($n = 8$ M).
910



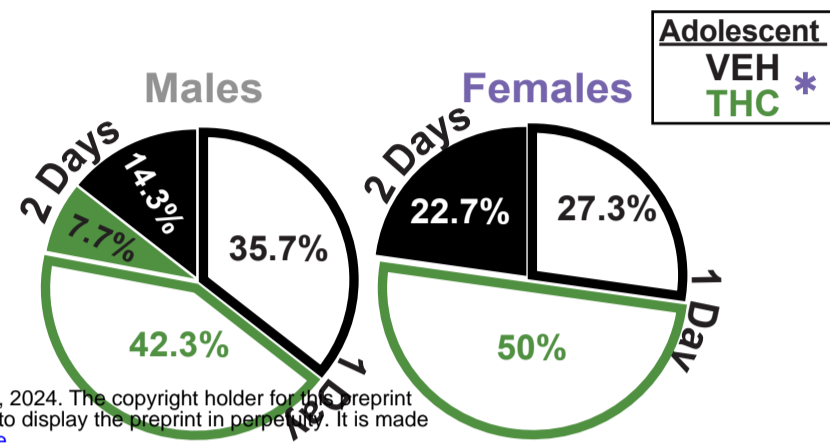
Adolescent THC History Improves Learning in Females

b. Visual Cue Discrimination

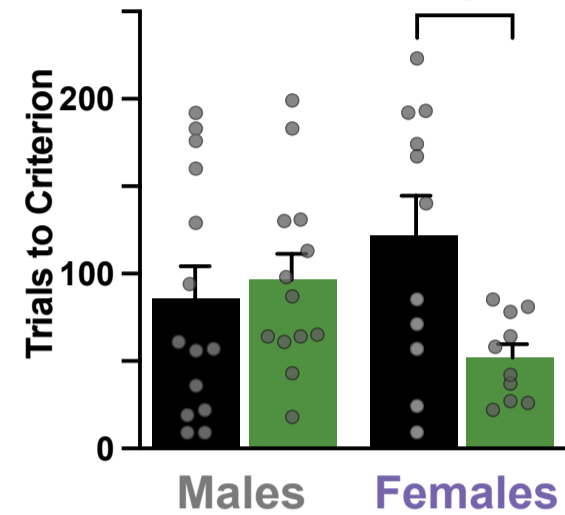


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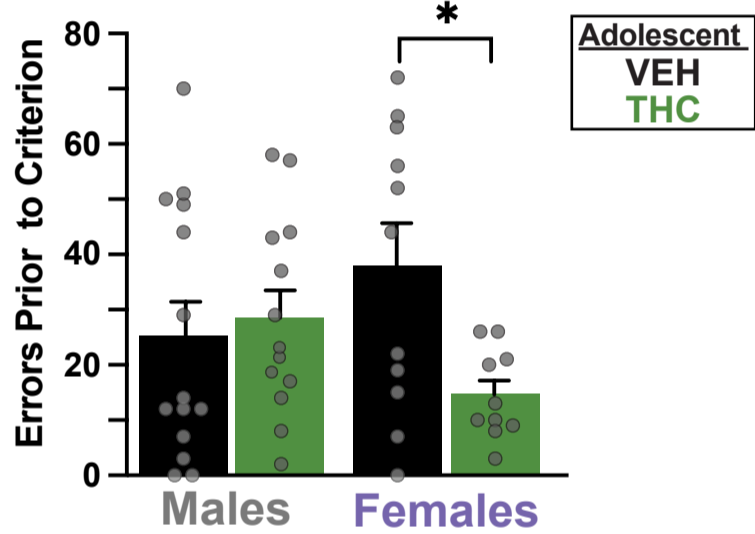
c. Days to Criterion



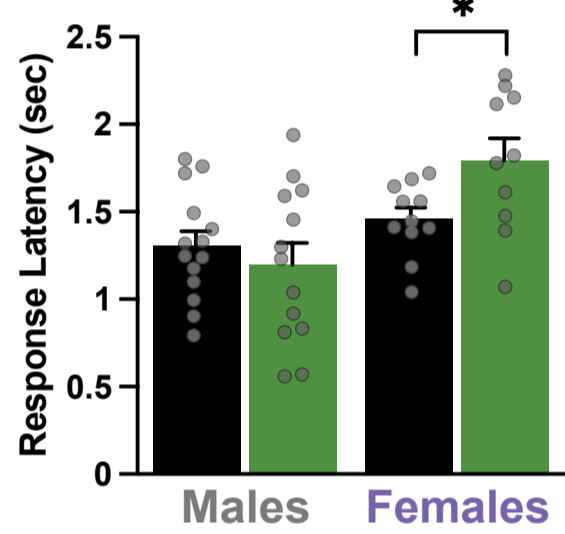
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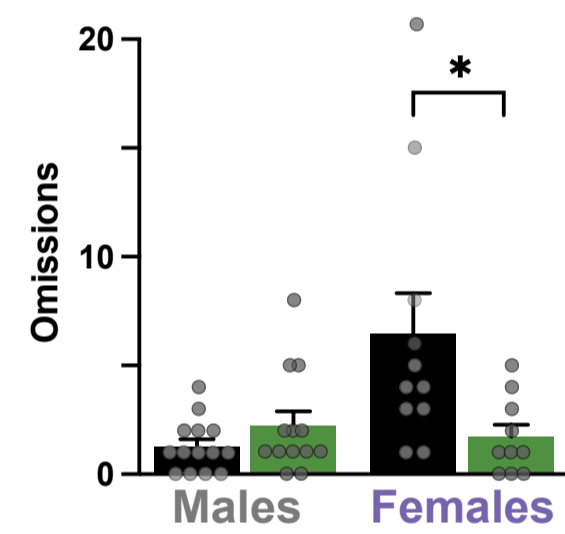
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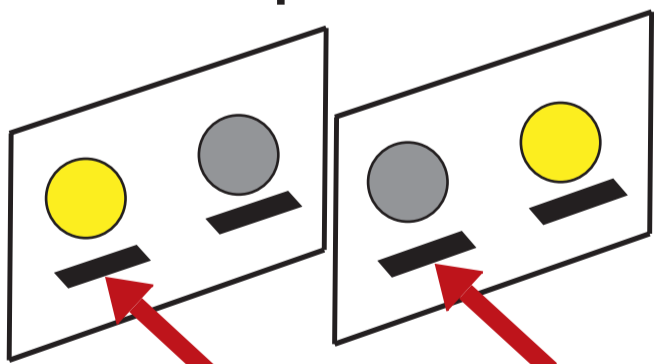


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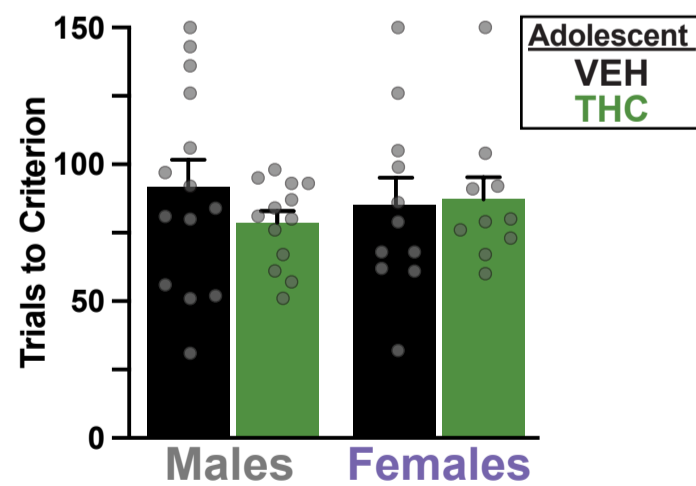


Adolescent THC History Does Not Affect Cognitive Flexibility

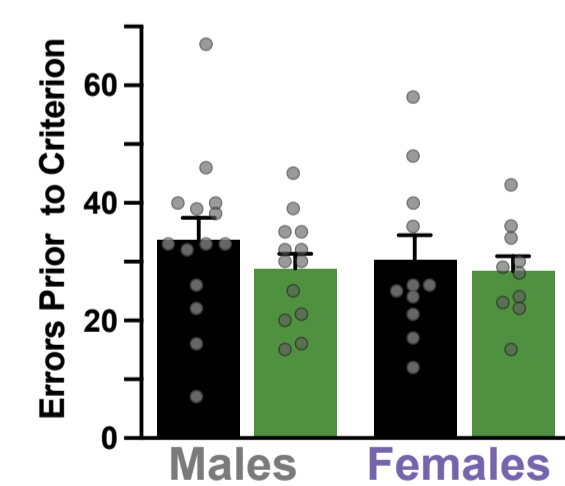
h. Shift to Response Discrimination



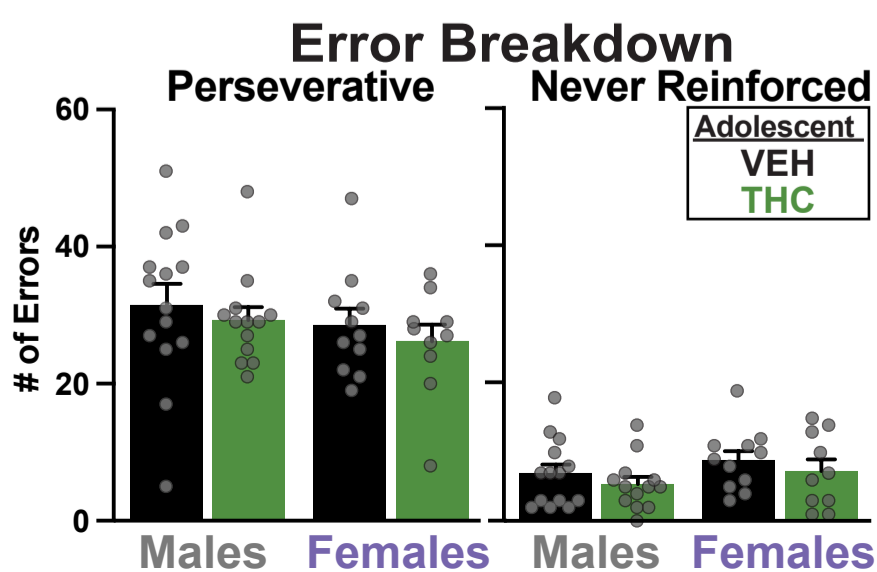
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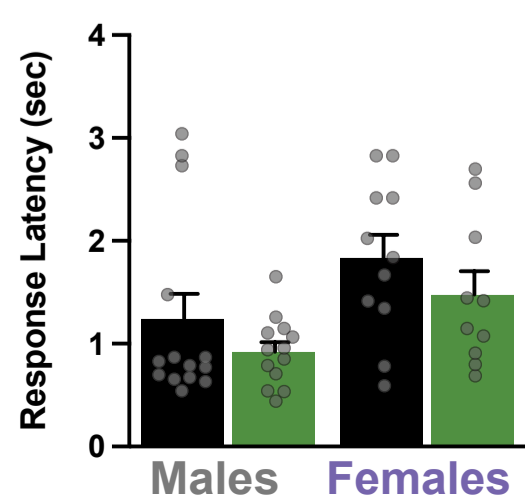
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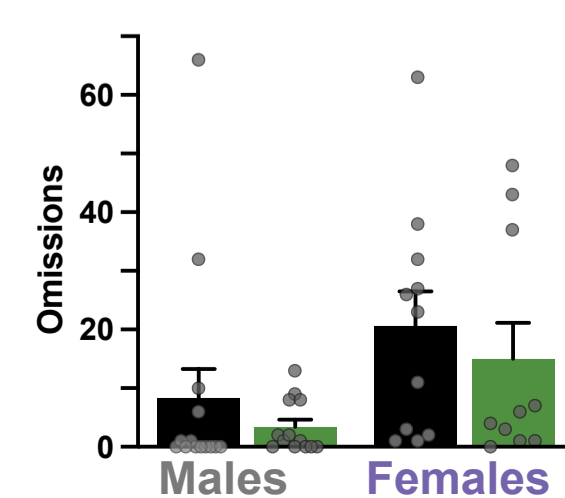
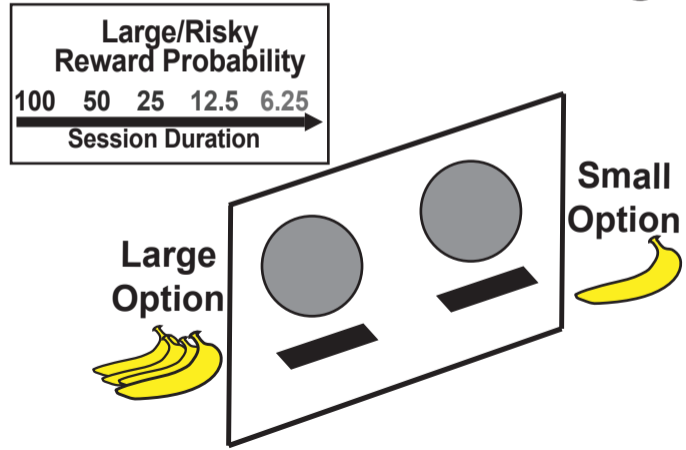


Fig. 2

a. Probabilistic Discounting



b.

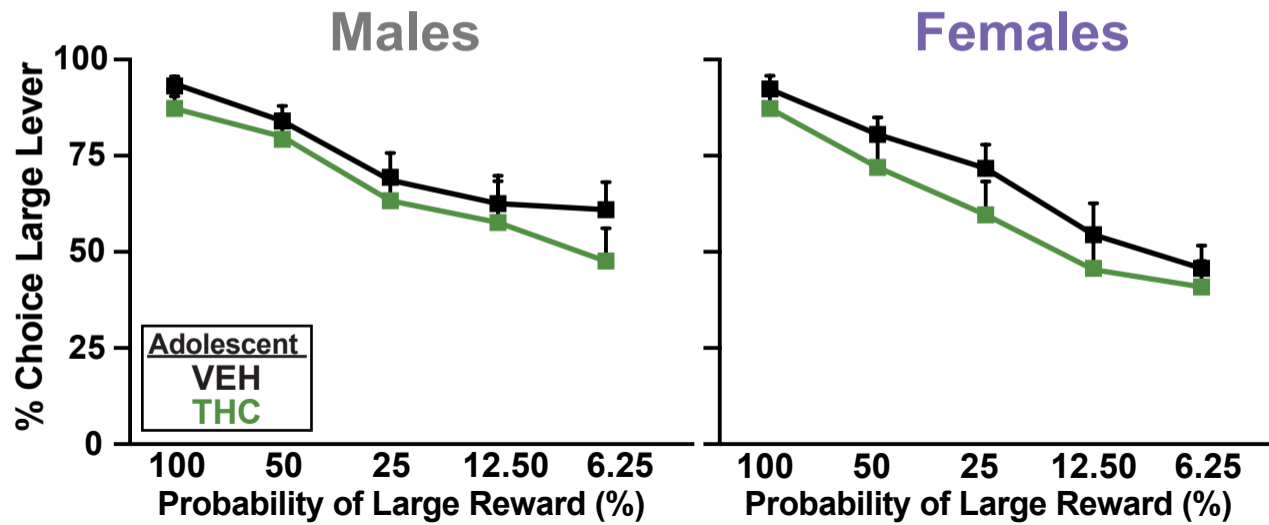


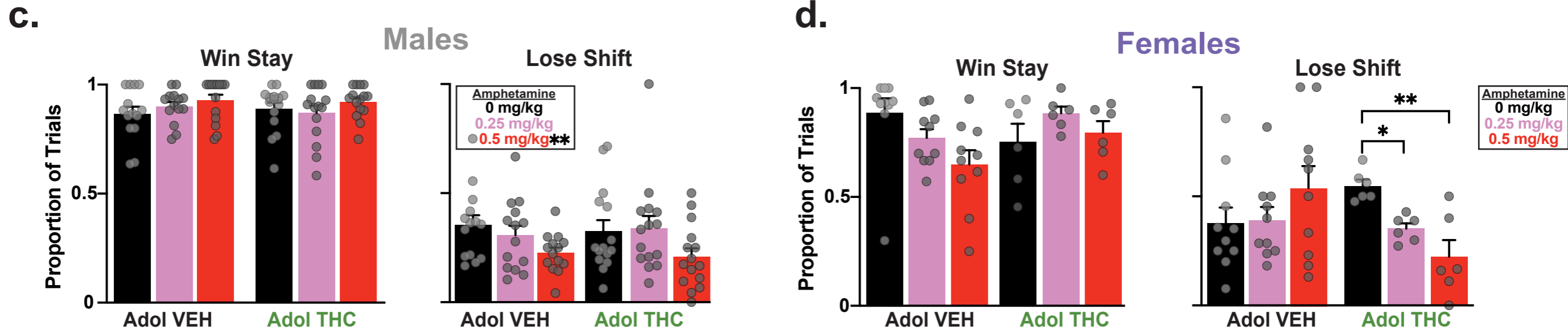
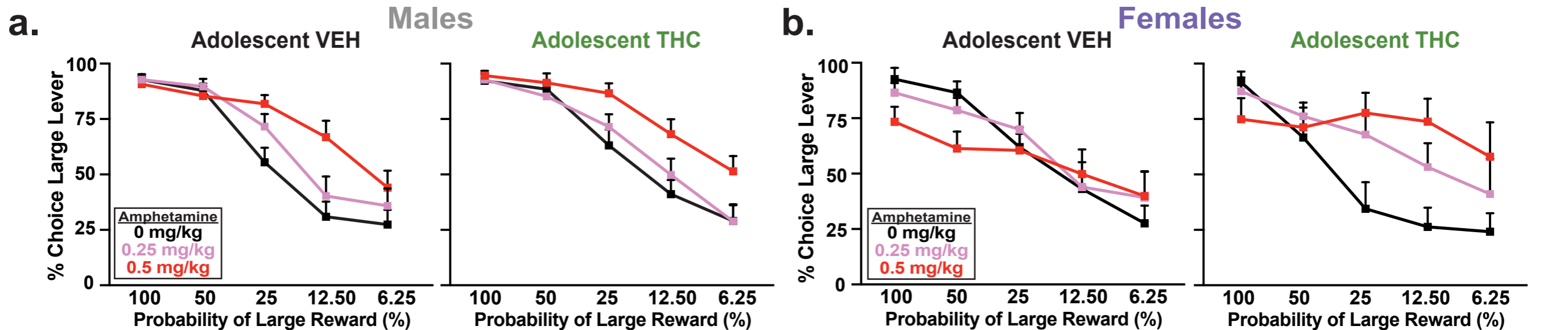
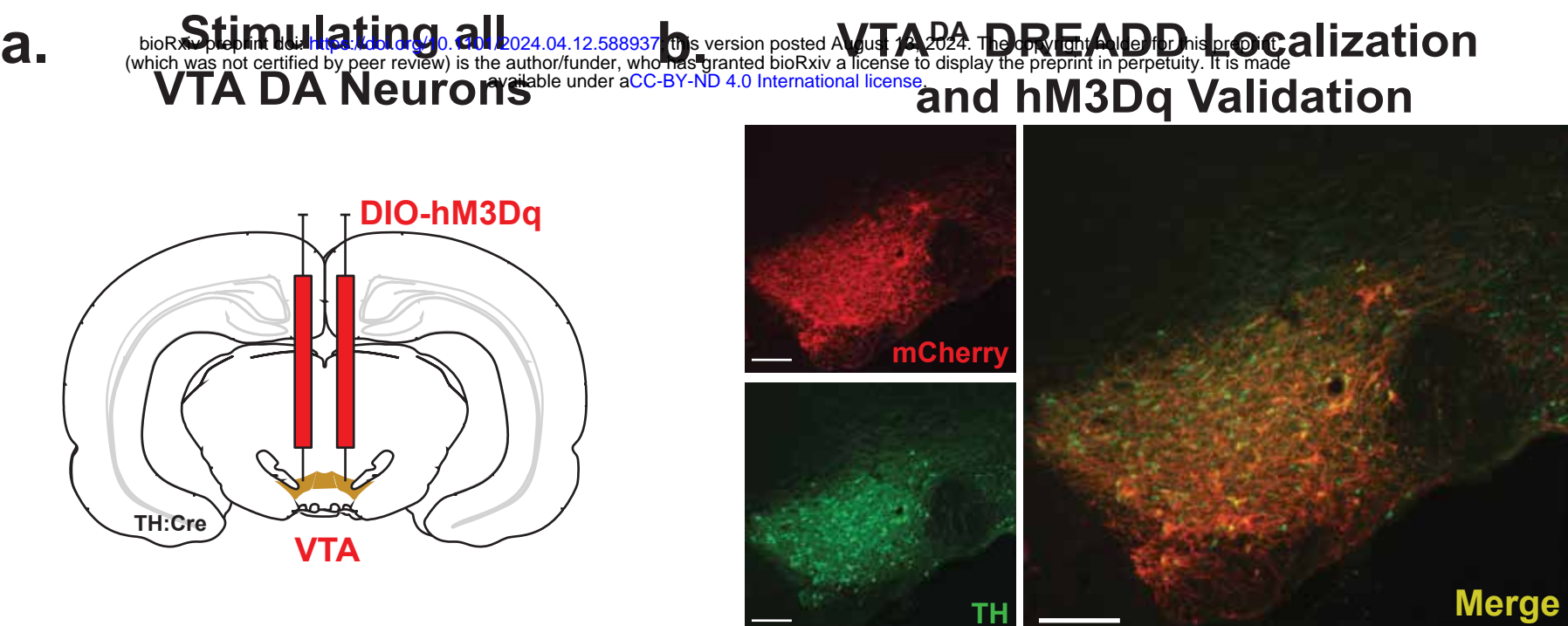
Fig. 3

Fig. 4**Experiment 1****Ventral Tegmental Area Dopamine Neuron Stimulation****Experiment 2****VTA Dopamine to Medial Prefrontal Cortex Pathway Stimulation**