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Incentive Learning Underlying Cocaine-Seeking Requires mGluR5 Receptors Located on Dopamine D1 Receptor-Expressing Neurons

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Understanding the psychobiological basis of relapse remains a challenge in developing therapies for drug addiction. Relapse in cocaine addiction often occurs following exposure to environmental stimuli previously associated with drug use (Stewart et al., 1984). The enduring control over relapse by cocaine-paired stimuli reflects the ability of addictive drugs to hijack neural substrates of associative reward-learning and memory that normally enable environmental stimuli paired with natural rewards (e.g., food or water) to guide adaptive behaviors (Robinson and Berriidge, 1993; Berke and Hyman, 2000; Kauer and Malenka, 2007). However, associative reward-learning can be dissociated into a variety of psychologically and neurobiologically distinct processes (Everitt et al., 2001). Consequently, understanding the psychobiological basis of relapse is of considerable importance for developing effective treatments for cocaine addiction.

A common neuronal substrate of associative reward-learning processes involves striatal medium spiny neurons (MSNs), which integrate mesostriatal dopaminergic signals and glutamatergic inputs arising from cortical and limbic regions (Kauer and Malenka, 2007; Goto and Grace, 2008). MSNs provide the sole striatal output to motivational and motor systems and can be divided into two functionally distinct populations, expressing either dopamine D1 (D1-MSNs) or D2 (D2-MSNs) receptors (Gerfen et al., 1990; Heiman et al., 2008; Valjent et al., 2009). However, the relative contributions of D1- and D2-MSNs to motivational output and the molecular events in MSNs underpinning associative reward-learning processes that contribute to relapse-like behaviors remain elusive.

The metabotropic glutamate receptor, mGluR5, is particularly interesting in this context. It is involved in several forms of plasticity in striatal MSNs that are proposed to mediate associative learning and memory processes (Sung et al., 2001; Gubellini et al., 2003; Malenka and Bear, 2004; Hyman et al., 2006; Schottanus and Chergui, 2008), and which are affected by cocaine expe-
rience (Martin et al., 2006; Kauer and Malenka, 2007; Kourrich et al., 2007; Bellone et al., 2008; Anwy, 2009; Moussawi et al., 2009). Although mGluR5 is densely expressed on both D1- and D2-MSN populations (Tallaksen-Greene et al., 1998), converging lines of research would suggest that mGluR5 located specifically on D1-MSNs is ideally positioned to influence associative reward-learning processes that may underpin relapse triggered by drug-paired stimuli. First, there is evidence that striatal dopamine D1 receptors (D1R) play a critical role in both the consolidation of associative reward-learning memories (Dalley et al., 2005) and many of the long-term effects of addictive drugs (Anderson and Pierce, 2005) and second, mGluR5 appears to interact closely with D1Rs to regulate striatal neurotransmission (Paolillo et al., 1998; Voulalas et al., 2005; Schotanus and Cher- gue, 2008).

Here, we determine the role of mGluR5 located on dopamine D1 receptor (D1R)-expressing neurons, in behaviors influenced by drug- or natural reward-paired stimuli, by generation of a novel mouse line in which mGluR5 is selectively knocked-down in neurons expressing the D1R. These mice reveal a necessary role of mGluR5 located on D1R-expressing neurons for highly specific associative reward-learning processes underlying cue-induced reinstatement of cocaine-seeking.

Materials and Methods

Mouse generation

Short hairpin RNAs were designed using the sFold (sTarMir) and BLOCK-I-T Prime Mir Designer (Invitrogen) software packages and tested in cell culture for knock-down (KD) efficiency of mGluR5 mRNA. BLOCK-I-T Pol II miR RNAi Expression vector kit with GW/EmGFP-miR Vector (Invitrogen) was used to insert synthetic oligos to artificial miRNA context (Fig. 1B). The construct was recombined into a bacterial artificial chromosome (BAC; RP24–17E13; Children’s Hospital Oakland Research Institute, Oakland, CA) harboring the mouse D1R gene following a procedure previously described (Parkin et al., 2009) (Fig. 1A). The BAC was purified, the vector sequences were removed, and the transgene was injected into the pronuclei of fertilized oocytes from a C57BL/6J mouse. Experimental animals were generated by backcrossing of mGluR5<sup>−/−</sup> transgenic mice to C57BL/6J inbred. Transgenic animals were genotyped using the following primers: ACCTGAACGAGCCA-CAAGTGC, GAATGCCAACGAGCCAA-CAAGTG, ATGGCCAGAGTCT-ACGTCGCTTACC, and CTCTAGAACTG-GAGGGACCAG. Probe was generated using a DIG RNA Labeling Kit (Roche). Brains were sectioned using 4% paraformaldehyde at 4°C for 24 h and 50 μm free-floating vibratome sections were hybridized with the DIG-labeled probe at 70°C overnight. Signal was developed using alkaline phosphatase-conjugated antigen binding fragments and the DIG-labeled probe at 70°C overnight. The membrane was developed with substrate ECL plus Western Blotting Reagents Mix (GE Healthcare).

Apparatus for cocaine and associative learning studies

Behavioral training and testing were performed in mouse conditioning chambers (Med Associates), individually housed within sound and light attenuating cubicles. Each chamber was equipped with a pellet dispenser connected to a recessed food magazine. A retractable lever was located on each side of the magazine and a cue light was positioned above each lever. A tone generator was situated between the cue lights and a house light was positioned on the wall opposite to the food magazine. Conditioning chambers were controlled and responses were recorded using a computer running Med-PC IV (Med Associates). The use of laboratory animals; experiments in Germany were approved by the local animal care committee (Karlsruhe, Germany); experiments in the UK were performed in accordance with the United Kingdom 1986 Animals (Scientific Procedures) Act, following institutional ethical review.

Animals for behavioral analysis

Cocaine studies were conducted in Mannheim, Germany while associative learning studies took place in Brighton, UK. In both laboratories, male Wistar and CD-1 mice (minimum 8 weeks old) were maintained on a 12-h light-dark cycle (with lights on at 7:00 AM) under controlled temperature (21 ± 2°C) and humidity (50 ± 5%) conditions. All experiments took place during the light phase. For cocaine studies, mice were single housed and for all other studies, mice were single or group housed. For all studies, body weights were maintained at ~85% of ad libitum feeding weight except for the cocaine self-administration phase during which mice received ad libitum access to food. Experiments were conducted in accordance with European Union guidelines on the care and use of laboratory animals; experiments in Germany were approved by the local animal care committee (Karlsruhe, Germany); experiments in the UK were performed in accordance with the United Kingdom 1986 Animals (Scientific Procedures) Act, following institutional ethical review.

In situ hybridization

An ~900-bp-long digoxigenin (DIG)-labeled riboprobe was used to detect mGluR5 mRNA detection. The DNA template was synthesized using the primers: ACCCTATCTGCTCTTCCTACC and GTCTACTGAATG-T1GCATGAGGGACCAG. Probe was generated using a DIG RNA Labeling Kit (Roche). Brains were sectioned using 4% paraformaldehyde at 4°C for 48 h and 50 μm free-floating vibratome sections were hybridized with the DIG-labeled probe at 70°C overnight. Signal was developed using alkaline phosphatase-conjugated antigen binding fragments and the DIG-labeled probe at 70°C overnight. The membrane was developed with substrate ECL plus Western Blotting Reagents Mix (GE Healthcare).

Removal of ribosomal RNA was verified on RNA LabChips. Small RNAs were detected by quantitative PCR using MicroRNA Reverse Transcription Kit (Applied Biosystems) and microRNA detection assays: mmu-miR-9 (part #4373371), hsa-miR-15a (4373123), hsa-miR-16 (4373121), mmu-miR-124a (4373150), hsa-miR-138 (4373175), snoRNA-234 (4380915) on 10 ng of the small RNA sample. Removal of ribosomal RNA was verified on RNA LabChips. Small RNAs were detected by quantitative PCR using MicroRNA Reverse Transcription Kit (Applied Biosystems) and microRNA detection assays: mmu-miR-9 (part #4373371), hsa-miR-15a (4373123), hsa-miR-16 (4373121), mmu-miR-124a (4373150), hsa-miR-138 (4373175), snoRNA-234 (4380915) on 10 ng of the small RNA sample.

Immunohistochemistry and immunofluorescence

Immunohistochemistry with anti-GFP antibody (1:10 000, Invitrogen, A11122, Lot 50434A) was performed using avidin-biotin-peroxidase complex (ABC) amplification and diaminobenzidine as a substrate. For immunofluorescence we used: rabbit anti-GFP (1:1000, Invitrogen, see above), donkey anti-rabbit Alexa Fluor 488 (1:1000, Invitrogen, A21206), chicken anti-GFP (1:1000, Abcam, ab13970), donkey anti-chicken Alexa Fluor 488, rabbit anti-prepro enkephalin (Neuromics, RA15125), goat anti-rabbit Alexa Fluor 594, mouse anti-DARP-32 (BD Transduction Laboratories, 611520), mouse anti-NeuN (1:400, Millipore Bioscience Research Reagents, MAB377, Lot 0604027006), Cy5-conjugated antimouse (Jackson Immunoresearch), chicken anti-mouse Alexa Fluor 594 (1:100, Invitrogen, A21201). Image analyses were performed with the ImageJ (v1.37, Wayne Rasband, National Institutes of Health, Bethesda, MD) and Creative Suite CS4 (Adobe) software. GFP and NeuN-positive cells were counted on 8 consecutive striatal sections per animal.

Cocaine studies were conducted in Mannheim, Germany while associative learning studies took place in Brighton, UK. In both laboratories, male Wistar and CD-1 mice (minimum 8 weeks old) were maintained on a 12–12 h light-dark cycle (with lights on at 7:00 AM) under controlled temperature (21 ± 2°C) and humidity (50 ± 5%) conditions. All experiments took place during the light phase. For cocaine studies, mice were single housed and for all other studies, mice were single or group housed. For all studies, body weights were maintained at ~85% of ad libitum feeding weight except for the cocaine self-administration phase during which mice received ad libitum access to food. Experiments were conducted in accordance with European Union guidelines on the care and use of laboratory animals; experiments in Germany were approved by the local animal care committee (Karlsruhe, Germany); experiments in the UK were performed in accordance with the United Kingdom 1986 Animals (Scientific Procedures) Act, following institutional ethical review.

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Apparatus for cocaine and associative learning studies

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Cocaine studies

Lever training and surgery. The procedures for lever training, surgery and catheter maintenance were as previously described (Mameli et al., 2009). In brief, to familiarize mice with the action of lever pressing, all mice were trained to lever press for food for a minimum of 14 sessions. The implantation of an indwelling catheter in the right jugular vein occurred 24 h after completion of lever training. Animals were given a minimum of 48 h recovery before cocaine self-administration sessions began.

Figure 1. Knock-down of mGluR5 in striatal dopamine receptor D1-MSNs. A, Design of the transgene expressing GFP as a marker and two interfering RNAs (iRNAs). This construct was inserted after the translational start of the gene encoding the dopamine D1 receptor in a bacterial artificial chromosome. B, Sequences of iRNAs. Interfering sequence is depicted in bold. Red arrows indicate targeted regions of mGluR5 mRNA. C, Expression of the transgene in mGluR5 KD mice (KD) as detected by immunohistochemistry for GFP in a sagittal brain section. Higher magnification showing difference between staining of cell bodies in the caudate–putamen (CPu) and its projections to ventral midbrain nuclei (VMN). D, The transgene (GFP; green) is expressed in ~53% of the striatal neurons (NeuN; red; → indicates examples of GFP-positive neurons and ↑ indicates examples of GFP-negative neurons). E, The expression of the construct is selective for D1-MSNs. Thus, expression is limited to MSNs (DARPP-32; blue) and absent from D2-MSNs (labeled by red immunofluorescent labeling of prepro enkephalin; ppEnk). F, Expression of mGluR5 in the striatum as shown with in situ hybridization. G, Knock-down assessment by quantitative PCR (n = 4–5, p < 0.001) and H, Western-blotting with representative blot example shown (n = 4, p = 0.00112). Data are presented as mean ± SEM, p-value of t test (*p < 0.05, **p < 0.001). Scale bars 20 μm. Cx, Cortex; Acb, nucleus accumbens.

Cocaine self-administration. Once-daily, 90 min, self-administration sessions commenced with the insertion of two levers into the condition-
ing chamber. Responses on one lever (the active lever), under a fixed-ratio 4 schedule (FR4), resulted in a 14–28 μl infusion of cocaine (cocaine hydrochloride; Sigma-Aldrich) delivered by activation of the pump for 1.2–2.4 s. Responses on the alternative lever (the inactive lever) were recorded, but had no scheduled consequence. Each drug infusion was associated with the 20 s presentation of flashing (1 Hz) cue lights (conditioned stimulus (CS)), which also signalled a time-out period during which further lever responses were not reinforced.

For dose–response determination, KD (n = 14) and WT (n = 14) mice were given access to different cocaine doses (0.095–1.5 mg/kg per infusion) in a randomized order during 90 min once daily self-administration sessions. When self-administration behavior was stable for one dose (three consecutive sessions with ±20% variation in the number of infusions earned) mice were given access to a different cocaine dose. Data from the third stable self-administration, from animals with a patent catheter, were used to generate the dose–response curve.

Cue-induced reinstatement. KD (n = 7) and WT (n = 6) mice were trained to self-administer cocaine (0.75 mg/kg per infusion) during 10 consecutive sessions under identical conditions to those described above. In addition, 7 animals (n = 4/3; KD/WT) which received cocaine 0.75 mg/kg per infusion as the final dose of the dose–response study were added to this experimental cohort. After the final cocaine self-administration session, mice received 14, once daily, 90 min extinction sessions in which responses on both levers were recorded but had no scheduled consequence. Prior studies from our laboratory (unpublished) revealed that 14 extinction sessions was sufficient to produce stable lever responding with active lever responses reduced to 50% or less of responses maintained by cocaine, as well as complete loss of discrimination between the active and inactive levers. Reinstatement tests took place 24 h after the last extinction session under conditions identical to the final session of cocaine self-administration, except that cocaine was not available. Thus, responses on the previously active lever triggered the noise of the infusion pump and a brief CS presentation. Responses on the inactive lever were without consequence.

### Procedure

**Magazine training.** To familiarize mice with the food used for conditioning studies (5TUL, catalog #1811142; Test Diet), a small amount of the food was given to all mice in their home cage. Mice also received a single, 30 min, magazine training session in which food pellets were delivered once every 60 s, on average (range of 25 to 95 s).

**Goal-tracking and conditioned reinforcement.** The procedures for Pavlovian conditioning, goal-tracking and CRf tests were as previously described (O’Connor et al., 2010). In brief, KD (n = 12) and WT (n = 9) mice received 11, once daily, 60 min Pavlovian conditioning sessions in which 16 presentations of a 10 s stimulus paired with food delivery (CS+; flashing cue lights) and 16 presentations of a 10 s stimulus paired with no outcome (CS−; the alternative stimulus) occurred. Each stimulus presentation was separated by a variable, no-stimulus ITI (range of 80–120 s; M = 100 s). A single food pellet was delivered 5 s after CS+ onset. For the 45 min sign-tracking test, conducted 24 h after the final conditioning session, two nose-poke holes were inserted into the conditioning chamber. In one hole, 15 × 1 min presentations of a flashing cue light (that is, the CS+) occurred. Each CS+ presentation was separated by a 0.25 min, no-stimulus ITI. No food presentation occurred in the second (control) nose-poke hole and no food was delivered during the test. Entries into each hole were recorded during CS+ presentations, thus providing a measure of sign-tracking responses (that is, approaches) toward the CS+.

**Statistical analysis.** For the assessment of the knock-down efficiency by quantitative PCR and Western blotting, statistical analyses were performed using t test. For cocaine (self-administration and cue-induced reinstatement) and associative learning (goal-tracking, CRf and PIT) studies, data were initially analyzed by mixed-factor ANOVA, where genotype comparisons were represented by the between-subjects factor of genotype (WT, KD). When a significant (≤0.05) main effect or interaction term was found, further analysis was performed using ANOVA and post hoc comparisons by Newman–Keuls or t test. For the sign-tracking test, approaches toward the CS+ or a control nose-poke hole were initially compared for each genotype by Mann–Whitney U test, with comparisons of responding in each nose-poke between genotypes made with Wilcoxon matched pairs test.

### Results

**Generation and validation of mice with knock-down of mGlur5 selectively in D1-expressing neurons.** To test the role of mGlur5 on D1-expressing neurons we generated mice with a selective knock-down of mGlur5 in these cells (mGlur5 K/D mice). We used a construct that expresses two artificial microRNAs targeting mGlur5 mRNA under the control of the D1R promoter (Fig. 1A, B). The coding sequence for green fluorescent protein (GFP) was introduced in tandem with two artificial microRNAs targeting mGlur5 mRNA under the control of the D1R promoter (Fig. 1A, B). The coding sequence for green fluorescent protein (GFP) was introduced in tandem.
with the microRNAs (Fig. 1A), enabling us to easily track expression of the construct. Immunostaining of GFP in brains from mGluR5KD-D1 mice showed that the expression pattern fits with that described for D1Rs, including strong expression in the dorsal striatum and nucleus accumbens (Fig. 1C). A more detailed examination of the striatum confirmed that the transgene (GFP) was expressed in ~53% of the striatal neurons (Fig. 1D, NeuN). Furthermore, expression of the transgene was confined to MSNs (identified by immunostaining against DARPP-32) (Fig. 1E) but the transgene was not expressed in D2-MSNs (identified by immunostaining against preproenkephalin; ppEnk) (Fig. 1E), showing that expression is restricted to D1-MSNs. Next, we analyzed whether expression of the transgene reduces the abundance of the mGluR5 transcript. In situ hybridization revealed reduced numbers of mGluR5-positive cells in the striatum, while the staining-intensity in the cells still expressing mGluR5 was not reduced (Fig. 1F), indicating strong mGluR5 knock-down selectively in the targeted cells. The abundance of mGluR5 transcript was reduced to ~40% in the homogenized striatum (Fig. 1G) with the corresponding protein reduced to ~50% compared with levels in WT mice (Fig. 1H). Since the expression of the construct is restricted to D1-MSNs (Fig. 1E), we estimate that the knock-down efficiency is ~90% in the targeted cells. There was no significant reduction of mGluR5 mRNA in the cerebral cortex or in the hippocampus of mGluR5KD-D1 mice (Fig. 2B) confirming normal function of the endogenous microRNA processing machinery. Together, our data indicate a highly specific and efficient knock-down of mGluR5 mRNA without off-target effects or disruption of endogenous microRNA function.

**Cocaine self-administration and cocaine-seeking in mGluR5KD-D1 mice**

To explore the consequence of the specific knock-down of mGluR5 for behaviors related to cocaine addiction, we first examined the propensity of mGluR5KD-D1 mice to self-administer cocaine. When given access, in a randomized order, to five different doses of cocaine under a fixed-ratio (FR4) schedule of reinforcement, WT and mGluR5KD-D1 mice displayed comparable self-administration behavior (Fig. 3A). Responses on the ‘active’ lever, which resulted in cocaine infusions and the concomitant presentation of a simple light stimulus, exhibited comparable inverted U-shape curves between genotypes, demonstrating that mGluR5KD-D1 mice were able to adapt their responding to the dose of cocaine available. Moreover, when trained to self-administer cocaine (0.75 mg/kg per infusion) for 10 consecutive sessions, both WT and mGluR5KD-D1 mice rapidly acquired and maintained stable responding on the active lever (Fig. 3B). Collectively, these results indicate that the primary reinforcing effects of cocaine are unaffected by knock-down of mGluR5 on D1R-expressing cells.

The ability of the stimulus associated with cocaine infusions to reinstate extinguished cocaine-seeking was then assessed. Following stable responding on the active lever during cocaine self-administration sessions, cocaine-seeking responses were extinguished by withholding further drug infusions and stimulus presentations. During extinction sessions, both genotypes significantly reduced responding on the active lever (Fig. 3C). During the test of cue-induced reinstatement of cocaine-seeking, mGluR5KD-D1 mice made significantly fewer responses than WT mice on the active lever that now resulted in presentation of the previously cocaine-paired stimulus, but not cocaine itself (Fig. 3D). These findings indicate that mGluR5 located on D1R-expressing cells is intimately involved in the reinstatement of cocaine-seeking maintained by a cocaine-paired stimulus.

**Associative learning in mGluR5KD-D1 mice**

Through associative learning, a stimulus paired with reward (CS) can acquire informative or predictive properties that serve to signal the availability and/or location of the reward (goal-tracking) and can also acquire incentive motivational properties enabling CSs to attract attention (sign-tracking), energize ongoing reward-seeking (Pavlovian-instrumental transfer), and/or directly reinforce instrumental behaviors (conditioned reinforcement) (Rescorla, 1988; Robinson and Flagel, 2009). In principle, any of these neurobiologically distinct learned properties could contribute to the effects of drug-paired stimuli on drug-seeking and relapse (Everitt and Robbins, 2005). The next series of experiments examined the consequence of mGluR5 knock-down on D1R-expressing cells for these different aspects of associative reward-learning processes.

Using Pavlovian conditioning procedures, cohorts of hungry mice were presented with a stimulus associated with food delivery (CS+) and a second stimulus associated with no outcome (CS−) (conditioning data from conditioned reinforce-
ment/goal-tracking cohort shown in Fig. 4A). There was no genotype difference in the learning of predictive properties of the CS+ that enable it to signal the availability and location of reward, as indicated by an increase across conditioning sessions in the number of head-entries into the food-delivery magazine that occurred following onset of the CS+, but before food delivery (goal-tracking responses; Fig. 4B). mGluR5 KD-D1 mice were also able to attribute incentive properties to the CS+ necessary for energizing ongoing reward-seeking, as demonstrated by the ability of noncontingent CS+ presentations to enhance responding on a lever previously associated with food delivery (Pavlovian-instrumental transfer test; Fig. 4D).

However, when a CS+ was presented contingent upon a novel instrumental response, mGluR5 KD-D1 mice made significantly fewer responses on the lever that resulted in CS+ presentations than WT mice (conditioned reinforcement test; Fig. 4E). In this test, there were no genotype differences in responses on the lever that resulted in CS− presentations, or the latency to explore either lever (lever, genotype, and lever × genotype interaction, F < 1). The specific impairment in CS+ reinforced lever responding could not be attributed to a general inability of mGluR5 KD-D1 mice to acquire an instrumental response, because they readily acquired instrumental responding when it was reinforced by the primary food reward (see food self-administration training data from Pavlovian-instrumental transfer cohort, Fig. 4C). Together, these data indicate a necessary role of mGluR5 on D1-expressing neurons for incentive learning that enables a CS+ to serve as a conditioned reinforcer.

Finally, the ability of the CS+ to attract behavior was assessed by relocating a discrete light CS+ behind a nose-poke hole and measuring approach responses toward it. mGluR5 KD-D1 mice made significantly fewer approaches toward the light CS+ than WT mice and there were no significant genotype differences in responses into the control nose-poke hole (sign-tracking test; Fig. 4F). Thus, in addition to the aforementioned deficit in conditioned reinforcement, mGluR5 knockdown on D1-expressing neurons resulted in a deficit in the attribution of incentive properties to the CS+ necessary for the CS to become highly salient and attractive (Robinson and Berridge, 1993; Tomie et al., 2008).

Discussion

Using cell type-specific RNA interference, we have generated a novel mouse line in which the metabotropic glutamate receptor, mGluR5, is selectively knocked-down on cells that express dopamine D1 receptors. We identify this mGluR5 population as playing a dissociable role in the primary versus secondary (that is, conditioned) reinforcing effects of cocaine, as revealed by normal cocaine self-administration but impaired cue-induced reinstatement of cocaine-seeking in mGluR5 KD-D1 mice. A detailed assessment of reward-learning in these mice reveals specific deficits in learning processes necessary for the attribution of incentive motivational properties to reward-paired stimuli that enable them to directly reinforce behaviors (conditioned reinforcement) and to become highly salient and attractive (sign-tracking). However, other aspects of reward learning were normal in mutant mice, including learning about the predictive properties of reward-paired stimuli which serve to signal the availability and location of reward (goal-tracking) and incentive learning that enables the reward-paired stimulus to energize responding directed toward obtaining a reward (Pavlovian-instrumental transfer). Collectively, our data indicate that mGluR5 located on D1R-expressing neurons play a central role in specific associative reward-learning processes, which are engaged following cocaine experience and thereby enable environmental stimuli associated with cocaine to exert a prolonged and pervasive influence over relapse susceptibility.

To interfere with the expression of mGluR5 selectively in D1R-expressing neurons we used a BAC-based construct in

Figure 3. Cocaine self-administration and cue-induced reinstatement in mGluR5 KD-D1 (KD) and control (WT) mice. Lever-press responses during cocaine self-administration (A, B), extinction (C), and cue-induced reinstatement test phases (D). A, Self-administration: cocaine-reinforced (●, active) and nonreinforced (▲, inactive) responses across five different doses of cocaine did not differ between genotypes (lever × genotype × dose interaction, \( F_{(4,102)} = 0.125; p > 0.05 \)). B, Similarly, lever responses across 10 consecutive sessions with a 0.75 mg/kg per infusion training dose did not significantly differ between WT and KD mice (session × lever × genotype, \( F_{(5,198)} = 1.56; p > 0.05 \)). During the training phase, the presentation of a CS was associated with each cocaine infusion. C, Extinction: responses on the active lever during the last 3 sessions of cocaine self-administration (C1-3) and 14 subsequent extinction sessions did not differ between the two genotypes (session × genotype, \( F_{(16,288)} = 1.27; p > 0.05 \)). D, Reinstatement: lever responses during the last extinction session (Ext) and the cue-induced reinstatement test (Reinst). Contingent presentation of the CS increased the number of responses on the active lever over extinction performance, in mice from both genotypes. However, reinstatement of the cocaine-seeking response was significantly lower in KD mice (genotype × lever × condition, \( F_{(1,36)} = 5.12; p < 0.05 \)). *Post hoc significant difference (\( p < 0.05 \)) from WT. $Significant difference (\( p < 0.01 \)) from the active lever responses during extinction. #Significant difference (\( p < 0.01 \)) from active lever responses during reinstatement. Responses are plotted as mean (±SEM).
Conditioned reinforcement: both genotypes preferentially responded on a lever that led to CS together with a very recent report (Garbett et al., 2010), our findings reflect off-target (Olive, 2009), anhedonic (Ba¨ckstro¨m and Hyytia¨, 2006) anisms in which targeted mutagenesis is not feasible. Previous use of RNAi-based approaches have raised our awareness that excessive levels of short RNAs may oversaturate exportin 5 and thus block the processing of endogenous short RNAs leading to perturbed cellular homeostasis (Grimm et al., 2006). This is not the case for the mGluR5KD-D1 mice, where matura-

tions show that this technique can be used successfully in the brain. Compared with conditional gene deletion this approach has the advantage that it involves only one mouse line and offers the perspective to be used, in modified forms, in other organisms in which targeted mutagenesis is not feasible. Previous use of RNAi-based approaches have raised our awareness that excessive levels of short RNAs may oversaturate exportin 5 and thus block the processing of endogenous short RNAs leading to perturbed cellular homeostasis (Grimm et al., 2006). This is not the case for the mGluR5KD-D1 mice, where matura-

which a conventional RNA-polymerase II promoter (the D1R-promoter) drives the expression of artificial microRNAs and a reporter. A similar approach has been reported previously for interference with other genes in nurse cells (Rao et al., 2006) and, together with a very recent report (Garbett et al., 2010), our findings reflect off-target (Olive, 2009), anhedonic (Bäckström and Hyytia¨, 2006) or reinforcing (van der Kam et al., 2009) effects of the pharmacological tools used.

Figure 4. Associative learning in mGluR5KD-D1 (KD) and control (WT) mice. A, Pavlovian conditioning: entries into a food magazine increased during presentations of a stimulus associated with food delivery (CS+), but decreased during presentations of a stimulus associated with no outcome (CS−). B, Goal-tracking: magazine entries that occurred following CS+ onset, but before food delivery (that is, goal-tracking responses), significantly increased across conditioning sessions (main effect of session, F(10,190) = 7.6, p < 0.01), but did not differ between genotypes (session × genotype interaction, $F_{(10,190)} = 0.171, p > 0.05$). C, Food self-administration for PIT cohort: both genotypes responded more on a lever that resulted in food delivery (Act), than an alternate lever on which responding had no consequence (Ina), when food delivery occurred under an FR1 (main effect of lever, $F_{(1,14)} = 54.84, p < 0.001$; lever × genotype interaction $F_{(1,14)} = 1.18, p > 0.05$) or a variable-interval 60 s schedule (VI60) (main effect of lever $F_{(1,14)} = 37.61, p < 0.001$; lever × genotype $F_{(1,14)} = 0.10, p > 0.05$). D, PIT test: responses on a lever that previously led to the delivery of food significantly increased during CS+ presentations, compared with a decrease in responding during CS− presentations (main effect of stimulus, $F_{(1,14)} = 20.93, p < 0.001$). There was no difference in PIT between genotypes (stimulus × genotype, $F_{(1,14)} = 0.125, p > 0.05$). E, Elevation score = lever responses during CS minus responses pre CS. F, Conditioned reinforcement: both genotypes preferentially responded on a lever that led to CS+ presentations, compared with a CS− paired lever (main effect of lever, $F_{(1,19)} = 24.38, p < 0.001$). However, KD mice made significantly fewer CS+ paired lever responses than WT mice (genotype × lever, $F_{(1,19)} = 5.57, p < 0.05$). *p < 0.05, post hoc comparison between genotypes by t test. F, Sign-tracking: both genotypes preferentially approached the location of the CS+ during its presentations. However, KD mice made significantly fewer CS+ approaches than WT mice. *p < 0.05, comparison between genotypes by Wilcoxon matched pairs test.

Using the cue-induced reinstatement model, considered an animal model of relapse vulnerability (Shaham et al., 2003; Sanches-Segura and Spanagel, 2006; Stephens et al., 2010), our current findings add to previous reports indicating a role of mGluR5 in regulating behavioral responses to cocaine (Chiamulera et al., 2001) and cocaine-paired cues (Bäckström and Hyytia¨, 2006) by suggesting a location of mGluR5 necessary for the cue-induced reinstatement of cocaine-seeking, while the primary reinforcing effects of cocaine are unaffected following specific knock-down of mGluR5 on D1R-expressing neurons. Our study also lends mechanistic confidence to previous reports that have used pharmacological tools to identify a role of mGluR5 in behaviors maintained by reward-paired stimuli (Tessari et al., 2004; Bespalov et al., 2005; Bäckström and Hyytia¨, 2006; Schroeder et al., 2008; Gass et al., 2009; Kumaesan et al., 2009; Martin-Fardon et al., 2009; O’Connor et al., 2010), since these reports could have re-

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is not a unitary process but can be dissociated psychologically, neurobiologically (Everitt et al., 2001) and genetically (Mead and Stephens, 2003a,b). Thus, to determine precisely which reward-learning processes were disrupted in mutant mice, we used Pavlovian conditioning procedures in which a stimulus was paired with the delivery of food [that is, the unconditioned stimulus (US)]. A potential limitation of this approach is that the extent to which neural circuitries that mediate associative learning for natural reinforcers (such as food) overlap with those engaged by drug reinforcers is not fully understood. However, attempts to employ purely Pavlovian conditioning procedures using a “drug US” have been hampered by the negative behavioral effects associated with nonresponse contingent drug delivery (Dowkin et al., 1995; Mitchell et al., 1996; Arroyo et al., 1998). Nevertheless, our findings that cocaine-seeking and specific incentive learning processes were both impaired in mutant mice provide empirical support for multiple contemporary theories of drug addiction, which propose that the ability of drug-paired stimuli to influence drug-seeking and relapse reflect the interactions of addictive drugs with neural systems that normally subserve associative reward-learning processes for natural reinforcers (Stewart et al., 1984; Tiffany, 1990; Robinson and Berridge, 1993; Everitt et al., 2001; Stephens and Duka, 2008; Thomas et al., 2008).

An advantage of the behavioral models used in the present study is that the underlying neural circuitry is relatively well characterized. The nucleus accumbens is crucial for learning necessary for conditioned reinforcement (Parkinson et al., 1999; Ito et al., 2004), the development of sign-tracking responses (Parkinson et al., 2000; Di Ciano et al., 2001) and reinstatement of cocaine-seeking (Fuchs et al., 2004). This strongly suggests that the deficits in associative learning and reinstatement of cocaine-seeking observed in the mGluR5KD-D1 mice are due to the lack of mGluR5 in D1-MSNs in the nucleus accumbens. Moreover, the continued expression of mGluR5 on non-D1 MSNs [that is, D2-MSNs, except the minority expressing both D2R and D1R (Valjent et al., 2009)] was insufficient to support specific incentive learning processes and relapse-like behaviors in mutant mice. Although we cannot formally rule out the contribution of mGluR5 on MSNs in the dorsal striatum or in other D1R-expressing cells, such as those in the hippocampus or cortex, a major contribution from mGluR5 in the latter structures seems unlikely since we saw no significant reduction of mGluR5 in the cortex or hippocampus of mGluR5KD-D1 mice. These observations may suggest that the D1R-promoter is less strong in these regions or that D1 and mGluR5 are not expressed in the same neuronal populations.

Recent reports have highlighted that stimulation of striatal D1R and NMDA receptors, and the resultant activation of extracellular signal–related kinase (ERK) specifically in D1 MSNs, represent critical mechanisms through which the long-term effects of addictive drugs are mediated (Heusner and Palmiter, 2005; Valjent et al., 2005; Bertran-Gonzalez et al., 2008). Moreover, both D1R and NMDA receptors in the accumbens appear critical for the early consolidation of appetitive Pavlovian memories (Dalley et al., 2005). These reports are particularly relevant in the context of our current findings, given the close interactions between mGluR5 and D1R (Paolillo et al., 1998; Voulalets et al., 2005; Schotanus and Chergui, 2008) and NMDA receptors (Pisani et al., 2001; Mao and Wang, 2002; Choe et al., 2006) in the striatum. Thus, it is possible that impaired incentive learning and relapse-like behaviors in mGluR5KD-D1 mice were due, in part, to changes in striatal D1R and NMDA receptor function as a consequence of mGluR5 loss. A future challenge will be to further understand the complex interplay of glutamate and dopamine signaling within striatal circuits, and determine precisely which cellular mechanisms encode appetitive memories and mediate subsequent behavioral responses to environmental stimuli associated with natural and drug reinforcers.

In summary, our present findings, together with a recent report from our laboratory (O’Connor et al., 2010), suggest that mGluR5-mediated neuroplastic events on D1-MSNs are crucial for the formation of psychologically distinct associations between environmental stimuli and rewards that endow reward-paired stimuli with the subsequent ability to both reinforce and attract motivated behaviors. Furthermore, recent reports have revealed that mGluR5-mediated striatal plasticity is involved in, or affected by, cocaine experience (Fourgeaud et al., 2004; Moussawi et al., 2009). Our report provides a psychobiological context for these findings by pointing to glutamate signaling at mGluR5 on striatal D1-MSNs as a key mediator through which repeated cocaine experience (and presumably exposure to other drugs of abuse) produces a persistent increase in the susceptibility to relapse triggered by environmental stimuli associated with drug use.

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