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To cite this version:
Louisa Lyon, Philip Burnet, James Kew, Corrado Corti, J.N.P Rawlins, et al.. Fractionation of Spatial Memory in GRM2/3 (mGlu2/mGlu3) Double Knockout Mice Reveals a Role for Group II Metabotropic Glutamate Receptors at the Interface between Arousal and Cognition. Neuropsychopharmacology, Nature Publishing Group, 2011, 10.1038/npp.2011.145. hal-00668575

HAL Id: hal-00668575
https://hal.archives-ouvertes.fr/hal-00668575
Submitted on 10 Feb 2012

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Fractionation of Spatial Memory in GRM2/3 (mGlu2/mGlu3) Double Knockout Mice Reveals a Role for Group II Metabotropic Glutamate Receptors at the Interface between Arousal and Cognition

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Running title – Group II mGluRs, spatial memory and arousal

Key Words: hippocampus, schizophrenia, metabotropic glutamate receptor, arousal

Word count: Abstract 250; Introduction 783; Methods 2100
ABSTRACT

Group II metabotropic glutamate receptors (mGluR2 and mGluR3, encoded by GRM2 and GRM3) are implicated in hippocampal function and cognition, and in the pathophysiology and treatment of schizophrenia and other psychiatric disorders. However, pharmacological and behavioural studies with group II mGluR agonists and antagonists have produced complex results. Here, we studied hippocampus-dependent memory in GRM2/3 double knockout (GRM2/3\(^{-/-}\)) mice in an iterative sequence of experiments. We found that they were impaired on appetitively motivated spatial reference and working memory tasks, and on a spatial novelty preference task that relies on animals’ exploratory drive, but were unimpaired on aversively motivated spatial memory paradigms. GRM2/3\(^{-/-}\) mice also performed normally on an appetitively motivated, non-spatial, visual discrimination task. These results likely reflect an interaction between GRM2/3 genotype and the arousal-inducing properties of the experimental paradigm. The deficit seen on appetitive and exploratory spatial memory tasks may be absent in aversive tasks because the latter induce higher levels of arousal, which rescues spatial learning. Consistent with an altered arousal-cognition relationship in GRM2/3\(^{-/-}\) mice, injection stress worsened appetitively motivated, spatial working memory in wild-types, but enhanced performance in GRM2/3\(^{-/-}\) mice. GRM2/3\(^{-/-}\) mice were also hypoactive in response to amphetamine. This fractionation of hippocampus-dependent memory depending on the appetitive-aversive context is to our knowledge unique, and suggests a role for group II mGluRs at the interface of arousal and cognition. These arousal-dependent effects may explain apparently conflicting data from previous studies, and have translational relevance for the involvement of these receptors in schizophrenia and other disorders.
INTRODUCTION

Group II metabotropic glutamate receptors (mGluRs) comprise mGluR2 (also called mGlu2), encoded by GRM2, and mGluR3 (mGlu3), encoded by GRM3. They are grouped together on the basis of sequence homology and a shared signal transduction mechanism, coupling via G\textsubscript{o} and G\textsubscript{i} proteins to inhibit adenylate cyclase (Niswender and Conn, 2010). mGlu2 receptors are neuronal, particularly in the pre-terminal regions of axons, away from the active zone of neurotransmitter release, whereas mGlu3 receptors are found both pre- and post-synaptically, and in glia. A key function of the pre-synaptic group II mGluRs is to inhibit neurotransmitter release, and mGluR2s primarily act as autoreceptors that negatively modulate glutamate release (Cartmell and Schoepp, 2000; Schoepp, 2001).

Group II mGluRs are being studied for their involvement in, and as potential treatment targets for, a range of psychiatric conditions including schizophrenia, anxiety and stress-related disorders, and substance misuse (Swanson et al, 2005; Witkin and Eiler, 2006; Markou 2007; Krystal et al, 2010). Their putative involvement in cognition and schizophrenia was stimulated by the seminal paper by Moghaddam and Adams (1998). These authors showed that an mGluR2/3 agonist, LY354740, reversed a spatial working memory (SWM) deficit in rats which had been produced by the NMDA receptor antagonist, phencyclidine (PCP). Krystal et al (2005) described comparable findings in human subjects and, notably, Patil et al, (2007) reported that an mGluR2/3 agonist was an effective antipsychotic in a clinical trial. Although unconfirmed, this study has encouraged further investigations of this therapeutic approach and its underlying rationale. In parallel, evidence has emerged that allelic variation in GRM3 affects human hippocampal function and activation (Egan et al, 2004; de Quervain and Papassotiropoulos, 2006), probably via genotype effects on glutamatergic transmission (Egan et al, 2004; Marenco et al, 2006). There is also modest
evidence that GRM3 may be a gene for schizophrenia, in its own right or via epistatic and gene-environment interactions (Egan et al, 2004; Nicodemus et al, 2007; see Harrison et al, 2008). Finally, the expression and function of mGlu2 and mGlu3 may be altered in the disorder (e.g. Gupta et al, 2005; Gonzales-Maeso et al, 2008; Sartorius et al, 2008; Ghose et al, 2009).

Together, these studies suggest that group II mGluRs are involved in cognition and in psychosis (Moghaddam, 2004; Moreno et al, 2009; Krystal et al, 2010). However, many uncertainties remain, not least since pharmacological studies in rodents aimed at elucidating the role of mGluR2/3 in hippocampus-dependent cognition have revealed a varied picture. Thus, in contrast to the improvement in SWM seen with an mGluR2/3 agonist in PCP-treated animals (Moghaddam and Adams 1998), mGluR2/3 agonists impair performance on this task when administered to non-pretreated rodents (Aultman and Moghaddam, 2001; Gregory et al, 2003) or non-human primates (Spinelli et al, 2005). The effects of mGluR2/3 antagonists on hippocampus-dependent memory tasks are also varied: LY341495 produced a small improvement in both T-maze SWM (Gregory et al, 2003) and spatial reference memory (SRM) in the Morris water maze (Higgins et al, 2004), whereas performance on a passive avoidance task was impaired (Sato et al, 2004). Moreover, other studies have found no effect of mGluR2/3 ligands on cognitive tasks (e.g. Schlumberger et al, 2009; Amitai and Markou, 2010). This complex pattern of results suggest that the effects of mGlu2/3 on cognition may be sensitive to various factors (e.g. glutamatergic ‘tone’, prior experience, or the nature of the task chosen), but the matter has not been clearly explored.

Mice lacking GRM2 (GRM2−/− mice) or GRM3 (GRM3−/− mice) provide a complementary means to assess the roles of these receptors, but although they have been studied in various domains, assessment of hippocampus-dependent cognition has been limited to a report that GRM2−/−-mice perform normally in the water maze (Yokoi et al, 1996). In any event, the phenotype of GRM2−/− and GRM3−/− mice may be affected by a degree of redundancy between these two
homologous receptors and the possibility of compensatory changes. Indeed, a small but significant increase in GRM2 expression is observed in the hippocampus of GRM3<sup>−/−</sup> mice and vice versa (Lyon et al., 2008). Here, we have studied double knockout mice in which both GRM2 and GRM3 have been deleted (GRM2/3<sup>−/−</sup> mice), precluding such compensation. Using an iterative series of experiments, we found that GRM2/3<sup>−/−</sup> mice display a striking fractionation of hippocampus-dependent spatial memory, with impaired performance on appetitive and spontaneous/exploratory spatial memory tasks but normal performance on equivalent aversive tasks. We present evidence that the observed pattern of deficits reflects an interaction between GRM2/3 genotype and the arousal-inducing properties of the experimental paradigms. This fractionation is not only unique, to our knowledge, but may be of importance with regard to the interface between cognition and motivation, and to their involvement in schizophrenia and other disorders.

MATERIALS AND METHODS

Subjects

Age-matched, male, wild-type and GRM2/3<sup>−/−</sup> mice were obtained from GlaxoSmithKline, Harlow, UK (see Supplementary Material for details). Behavioral experiments were conducted under the auspices of U.K. Home Office Project and Personal licenses held by the authors, and the study was approved by the local ethics committee. Four cohorts of mice were used. For full details of subjects and testing order see Supplementary Material.

Overview of experimental design for assessment of hippocampus-dependent spatial memory

A series of experiments was conducted, in which results from one set of experiments were used to generate hypotheses, which were then tested in subsequent experiments. For clarity, this iterative,
chronological sequence is summarised here before individual experiments are described in detail. Our initial experiments revealed that GRM2/3⁻/⁻ mice were impaired on appetitively motivated tasks measuring either spatial reference memory (SRM) and/ or spatial working memory (SWM) (Expts. 1-3; Y-maze, T-maze, radial arm maze). GRM2/3⁻/⁻ mice were, however, unimpaired on the aversively motivated Morris watermaze SRM task (Expt. 4). Further experiments were thus conducted to explore this dissociation.

The open field watermaze differs from the “dry land” mazes not only in terms of its aversive character, but also in terms of other procedural aspects. To reconcile these differences, we directly compared experimentally naïve GRM2/3⁻/⁻ and wild-type mice on appetitive and aversive versions of the Y-maze SRM task using a fully counterbalanced design (Expt. 5). These data confirmed that the observed dissociations were not merely consequences of differences in the mazes used, the spatial cues available, or the order of testing. We then performed further experiments to explore further the nature of the deficit in the GRM2/3⁻/⁻ mice. Experiment 6 examined whether the deficit extended to appetitively motivated, hippocampus-independent, behavioural paradigms by assessing performance on a non-spatial, visual discrimination learning task. Experiment 7 examined spatial memory performance in a spontaneous, exploratory task with neither a food reward nor an escape contingency. Finally, amphetamine challenge experiments were used to explore the effects of pharmacologically manipulating arousal levels on locomotor activity and cognition (Expts. 8-9).

Detailed methods, in addition to those presented below, can be found in the Supplementary Material.

Experiment 1: Appetitively motivated SRM on the elevated Y-maze

Each mouse was assigned a goal arm, defined by its position relative to extramaze spatial cues, which was baited with sweetened condensed milk on all trials. On each trial, the mouse was placed
at the end of one of the two non-baited arms (the “start arm”), and allowed to choose one of the remaining arms. If it chose the goal arm, it was allowed to consume the milk reward before being returned to the home cage. Mice that chose incorrectly were returned to the home cage immediately. Previous work in this laboratory has demonstrated that this task is hippocampus-dependent (Deacon et al., 2002). Mice received ten trials per day for nine days, with an inter-trial interval (ITI) of approximately five minutes.

**Experiment 2: Appetitively-motivated SWM on the elevated T-maze**

Each trial consisted of a sample run followed by a choice run. On the sample run, mice were forced either left or right by the presence of a wooden block, closing off one of the goal arms. At the end of the goal arm the mouse collected a reward of sweetened condensed milk. The block was then removed and the mouse placed back in the start arm, facing the experimenter, for the choice run. The mouse could now select either goal arm but was rewarded only for choosing the arm that had not been visited on the sample run, i.e., for alternating (non-matching to place). The interval between the sample and choice runs was approximately 5 seconds. Mice received five trials per day, with an ITI of approximately ten minutes. SWM performance on the T-maze is also dependent on the hippocampus (Deacon et al., 2002).

**Experiment 3: SRM and SWM in the six-arm radial maze**

A two-stage, within-subjects and within-task design was used (Schmitt et al., 2003).

*SRM acquisition:* In the initial acquisition phase, three of the six arms were baited with sweetened condensed milk. At the start of each trial, the mouse was placed on the central platform and given a free choice of arms. After visiting the food well in the chosen arm, the mouse would return to the central platform. The door to the visited arm was then closed and remained so for all
subsequent choices on that trial, preventing the mouse from re-entering that arm. After 10 seconds, all other doors were opened and the mouse allowed a second choice. This was repeated, with five doors open for the second choice, four for the third, and so on, until the mouse had visited all three baited arms. Each entry into an un-baited arm was scored as a SRM error. SRM acquisition on the radial maze is prevented by cytotoxic hippocampal lesions (Schmitt et al., 2003).

Simultaneous assessment of SRM and SWM: When all mice had successfully acquired the SRM task, the SWM component was introduced. Entries into previously visited arms were now no longer prevented but the food rewards were not replaced within a trial. Three types of error were scored (Jarrard, 1993), including SRM errors in which the mouse visited a never-baited arm, and SWM errors in which the mouse visited a baited arm that had already been visited on that trial. If second and subsequent visits to a never-baited arm occurred, these were scored separately as SRM-repeat errors.

Experiment 4: Aversively motivated SRM in the open field Morris watermaze
To escape from the water, mice had to locate a platform, hidden approximately 1 cm below the surface, which for a given mouse remained in the same position on all trials. Each mouse was given four trials per day for nine days. On days 7 and day 11, transfer (probe) tests were conducted. The platform was removed from the pool and the mouse allowed to swim freely for 60 seconds. The percentage of time spent in each quadrant was recorded, together with the number of times the mouse swam across the former location of the platform. We have previously shown in this same laboratory, with the same spatial cues, that acquisition of this fixed location, hidden platform watermaze task is prevented by cytotoxic hippocampal lesions (Deacon et al., 2002).

Experiment 5: Comparison of appetitively and aversively motivated SRM on the Y-maze
The aim of this experiment was to compare performance of wild-type and GRM2/3−/− mice on appetitively and aversively motivated versions of the Y-maze SRM task, with the order of testing and the experimental rooms in which the tests were performed (and therefore the spatial cues available), fully counterbalanced in order to control for practice effects or differences in the salience of the available spatial cues. Thus, 50% of wild-type and 50% of GRM2/3−/− mice first performed the appetitive Y-maze SRM task described in experiment 1 (same apparatus and protocol), in room A. These animals then performed an aversively motivated Y-maze SRM task (see below), in room B. Conversely, the remaining 50% began with the aversively motivated Y-maze task in room A, followed by the appetitively motivated Y-maze task in room B.

In the deep-water escape (aversive) Y-maze, mice could escape from the water by climbing onto a platform hidden approximately 1.5 cm below the water surface. Mice received five trials per day for six days. On each trial the mouse was allowed 90 seconds to find the platform; any that failed to do so were guided there by the experimenter. On day seven, a transfer test was performed, analogous to that used in the watermaze. The platform was removed from the maze and the mouse allowed to swim freely for 30 seconds. Time spent searching in each arm was recorded. Previous work within this laboratory has confirmed that the “swimming Y-maze” task, like the appetitive Y-maze, is hippocampus-dependent.

Experiment 6: Appetitively motivated visual discrimination

One possible explanation for this pattern of results is that GRM2/3−/− mice have reduced appetitive motivation for the milk reward. If this were the case then one would expect the mice to be impaired on other appetitively motivated tasks.

Visual discrimination learning for a food reward was examined using a T-maze with a light gray start arm, plus two removable goal arms. Successful performance involves learning an
association between a visual pattern and reward, but does not require the hippocampus (Morris et al., 1986; Murray and Ridley, 1999). The walls and floor of one of the goal arms were painted with black and white stripes, while those of the other goal arm were plain dark gray. Each mouse was assigned a particular goal arm (“black/white stripes” or “gray”), which was baited with sweetened condensed milk on all trials. On 50% of trials, the rewarded goal arm was positioned to the right of the start arm, and on the remaining 50%, to the left (according to a pseudorandom sequence). On each trial, the mouse was placed at the end of the start arm and allowed to choose one of the goal arms. If the mouse chose the correct arm, it was allowed to consume the milk reward before being returned to the home cage. Mice that chose incorrectly were returned to the home cage immediately. Mice received ten trials per day for five days.

Experiment 7: Spontaneous spatial novelty preference task

GRM2/3/- and wild-type mice were also compared on a spontaneous, spatial novelty preference task in which behavior is driven, not by an overt unconditioned stimulus (e.g. a food reward), but instead relies upon animals’ natural exploratory drive. This task therefore provides a non-aversive experimental context but performance does not rely on the motivating or rewarding effects of food.

The apparatus used was identical to the swimming Y-maze, but without the water. Each mouse was assigned two arms (the “start arm” and the “other arm”) to which they were exposed during the first phase of the task (the “exposure phase”). During this 5-minute “exposure” phase, the entrance to the third, “novel”, arm was closed off by the presence of a large Perspex block. At the end of five minutes, the mouse was removed from the maze and returned to the home cage for one minute. The mouse was then returned to the start arm for the test phase. This consisted of two minutes free exploration during which the mouse could now enter all 3 arms. The amount of time that the mouse spent in each arm, and the number of entries into each arm, were recorded, during
both the exposure and test phases. For the test phase, a discrimination ratio \[\frac{\text{novel arm}}{\text{novel + other arm}}\] was calculated. Previous work has demonstrated that wild-type mice display a marked preference for the novel arm during the test phase, which is abolished by cytotoxic hippocampal lesions (Sanderson et al., 2007).

**Experiment 8: Spontaneous and amphetamine-induced locomotor activity**

Spontaneous locomotor activity was first measured in transparent plastic photocell activity cages over 2 hr. The number of beam breaks made by each mouse was recorded in 24 bins of 5 minutes. A separate cohort of mice were then tested in a threshold activity monitoring system for locomotor activity over a 70 hr period (1 hr bins).

A further separate cohort of drug-naïve mice, with no previous experience of the photocell activity boxes, was used for the amphetamine challenge experiments. Mice were habituated to the activity boxes for 20 minutes prior to receiving an i.p. injection of either saline or 2.5 mg/kg amphetamine (injection volume 10 ml/kg body weight). Activity levels (number of beam breaks) were then measured for 2 hours.

Higher doses of amphetamine are known to induce stereotypy, which reduces locomotor activity. An additional cohort of drug-naïve mice was habituated to the activity boxes for 20 minutes prior to receiving an i.p. injection of either saline or 10 mg/kg amphetamine (10 ml/kg body weight). Mice were returned immediately to the activity boxes and their activity levels measured for 7 hours. Every 10 minutes throughout the first 4 hours, stereotypy was scored for each animal, using an established rating scale (Creese and Iversen, 1974).

**Experiment 9: Effect of injection stress and a low dose of amphetamine on appetitively motivated T-maze SWM**
To assess the effect of a low dose of amphetamine (0.5 mg/kg) on cognitive performance, we trained drug-naïve wild-type (n = 12) and GRM2/3\(^{-/-}\) (n = 11) mice on a T-maze rewarded alternation SWM task (as in Expt. 2). When the performance of both groups was stable, animals received a session of 10 trials with no injections. The following day, they were injected with either 0.5 mg/kg amphetamine or saline (i.p; 10 ml/kg body weight), according to a counterbalanced, within-subjects design. Each animal received a block of 10 trials on the T-maze SWM task 20 minutes after each injection, with drug and vehicle injections separated by 24 hours. Finally, 24 hours later, all animals received a final block of 10 trials with no injection.

**RESULTS**

*Experiment 1: Impaired appetitive SRM on the elevated Y-maze*

All GRM2/3\(^{-/-}\) mice (n = 14) and wild-type (n = 18) mice were able to learn the spatial location of the sweetened condensed milk reward on the Y-maze, a task that has previously been shown to be sensitive to hippocampal lesions (Deacon et al., 2002). However, the rate of acquisition of this appetitive SRM task was slower in GRM2/3\(^{-/-}\) mice (Fig. 1A). Two-way RM-ANOVA revealed an effect of group ($F_{(1,30)} = 8.88; p = 0.006$) and block ($F_{(8,240)} = 82.98; p < 0.001$), as well as a group*block interaction ($F_{(8,240)} = 5.06; p < 0.001$). Duncan’s multiple pairwise comparisons revealed that GRM2/3\(^{-/-}\) mice were impaired in blocks 3-6 inclusive (all $p < 0.05$). Post-choice reinforcement in block 9 confirmed that mice were not locating the milk by virtue of its odour.

*Experiment 2: Impaired SWM on the elevated T-maze*

Wild-type mice (n = 18) successfully performed this task, alternating their spatial responses in order to obtain a condensed milk reward. While GRM2/3\(^{-/-}\) mice (n = 14) did improve marginally over the course of training, they remained significantly impaired relative to their wild-type counterparts.
throughout (Fig. 1B; effect of group: $F_{(1,30)} = 16.43; p < 0.001$; effect of block: $F_{(4,120)} = 5.33; p < 0.001$; no group*block interaction: $F_{(4,120)} = 1.54; p = 0.196$).

In a separate cohort of wild-type ($n=17$) and GRM2/3$^{-/-}$ mice ($n=22$) we replicated this experiment, again showing that GRM2/3$^{-/-}$ mice ($71.4 \pm 1.4 \%$ correct) were impaired relative to wild-type controls ($77.6 \pm 2.0\%$ correct; $F(1,37) = 6.63; p < 0.05$). In addition, we recorded the latency of the mice to run (i) from the beginning of the start arm to the food well on the sample trial, and (ii) from the beginning of the start arm until making a choice into one of the goal arms on the choice trial. Importantly, the deficit in the GRM2/3$^{-/-}$ mice was not due to the fact that these animals simply ran more slowly and thus experienced an increase in the delay between the sample trial and choosing on the choice trial, increasing the mnemonic demands of the task. In fact, the GRM2/3$^{-/-}$ mice displayed shorter latencies than their wild-type controls on both the sample (median and inter-quartile (s); wild-type = 7.1 (4.3 - 64.0), GRM2/3$^{-/-} = 3.4$ (2.8 – 7.2; Mann-Whitney U-test, $p < 0.05$) and choice runs (median and inter-quartile (s); wild-type = 8.5 (3.3 – 40.6), GRM2/3$^{-/-} = 2.5$ (2.0 – 7.2; Mann-Whitney U-test, $p < 0.005$) during performance on this task.

**Experiment 3: Impaired SRM and SWM on the six-arm radial maze**

To confirm the results of experiments 1 and 2, GRM2/3$^{-/-}$ and wild-type mice were compared on a six-arm radial maze task, which permits simultaneous assessment of SRM and SWM. In the first stage, animals had to learn to discriminate between three baited and three non-baited arms (SRM). GRM2/3$^{-/-}$ mice displayed slower acquisition of this SRM component than wild-type animals (Fig. 1C). Two-way RM-ANOVA revealed an effect of block ($F_{(25,600)} = 25.91; p < 0.001$), as well as a group*block interaction reflecting the differing rates of acquisition ($F_{(25,600)} = 4.25; p < 0.001$). The main effect of group failed to reach significance ($F_{(1,24)} = 3.08; p = 0.092$), but Duncan’s multiple
pairwise comparisons confirmed that the GRM2/3\(^{-}\) mice were significantly impaired in blocks 6 – 13 inclusive (all \( p < 0.05 \)). The performance of the two groups was well-matched from block 14 onwards (all \( p > 0.05 \), with the exception of block 20 in which GRM2/3\(^{-}\) mice actually made fewer errors than wild-type animals, \( p < 0.05 \)). These additional blocks of trials were conducted to ensure that both groups of mice were well-matched for SRM performance, and thoroughly familiar with the location of the three rewarded arms, prior to the introduction of the SWM component.

In the second stage of this task, entries into already visited arms were now no longer prevented. Comparison of SWM error rates revealed that, as with T-maze rewarded alternation (Expt. 2), GRM2/3\(^{-}\) mice were again less able to remember which arms they had already visited on a given trial (Fig. 1D; main effect of group: \( F_{(1,24)} = 38.91; \ p < 0.001 \); no effect of block: \( F_{(3,72)} = 0.74; \ p = 0.531 \); no group*block interaction: \( F_{(3,72)} = 1.34; \ p = 0.268 \)). By this stage of training, both groups of mice were maintaining similarly low levels of SRM errors (no effect of group: \( F_{(1,24)} = 2.55; \ p = 0.124 \); no effect of block: \( F_{(3,72)} = 1.02; \ p = 0.387 \); no group*block interaction: \( F_{(3,72)} = 1.53; \ p = 0.214 \)). SRM-repeat errors were therefore very infrequent in both groups of mice (average \(< 0.2 \) per trial) (no effect of group: \( F < 1; \ p > 0.20 \); or block: \( F_{(3,72)} = 1.80; \ p = 0.155 \); and no interaction: \( F < 1; \ p > 0.20 \)) (data not shown).

Overall therefore, GRM2/3\(^{-}\) mice displayed slower but ultimately successful acquisition of the initial SRM phase, consistent with their slower acquisition but eventual success on the SRM Y-maze task (Expt. 1). GRM2/3\(^{-}\) mice also exhibited a persistent SWM impairment, consistent with their impaired performance on T-maze rewarded alternation (Expt 2).

Experiment 4: Unimpaired aversive SRM in the open field Morris watermaze
The performance of GRM2/3\(^{-}\) mice was examined in the standard, fixed location, hidden platform Morris watermaze task. In contrast to the appetitive spatial memory tasks, GRM2/3\(^{-}\) mice were
completely unimpaired (n = 18 wt; n = 13 GRM2/3\(^{-/-}\); Fig. 2). Since GRM2/3\(^{-/-}\) mice swam more slowly than wild-types (0.19 ± 0.004 cm/s vs. 0.21 ± 0.003 cm/s; t\(_{(29)}\) = 4.62; p < 0.001), distance traveled to reach the platform (path length) was taken as the performance measure during training. Two-way RM-ANOVA revealed an effect of day (F\(_{(8,232)}\) = 59.22; p < 0.001), but no effect of group, nor group*day interaction (both F < 1; p > 0.20).

Probe tests were conducted on days 7 and 11 (24 hours after training trials 24 and 36 respectively), in which the platform was removed from the pool and the animal allowed to swim freely for 60 s. In both probe tests, wild-type and GRM2/3\(^{-/-}\) mice concentrated their search in the quadrant that had previously held the platform (Fig. 2B,C). In the analysis of the distribution of time spent searching the four quadrants, the numerator term in the degrees of freedom was reduced by one to control for the fact that the quadrant dwell times were not independent. This analysis confirmed an effect of quadrant in both probe tests (day 7, F\(_{(2,87)}\) = 31.16; p < 0.05; day 11, F\(_{(2,87)}\) = 49.81; p < 0.05), but no group*quadrant interaction on either (day 7, F < 1; p > 0.20; day 11, F\(_{(2,87)}\) = 1.91; p = 0.13). A between-subjects t-test comparing the proportion of time spent searching in the training quadrant confirmed that both groups had learned the location of the platform to the same extent (day 7, t\(_{(29)}\) = 1.01; p = 0.32; day 11, t\(_{(29)}\) = 1.39; p = 0.18).

The number of “annulus crossings” made by the two groups (i.e., the number of times the mouse swam across the former location of the platform, along with the number of crossings of the equivalent positions in the other quadrants), were also compared. The results revealed equivalent performance in wildtype and GRM2/3\(^{-/-}\) mice in both probe tests (day 7, effect of quadrant: F\(_{(3,87)}\) = 21.34; p < 0.001, but no effect of group: F\(_{(1,29)}\) = 1.73; p = 0.198, and no group*quadrant interaction: F < 1; p > 0.20; day 11, effect of quadrant: F\(_{(3,87)}\) = 28.06; p < 0.001, but no effect of group: F < 1; p > 0.20, and no interaction: F\(_{(3,87)}\) = 1.89; p = 0.137). This was confirmed by
comparison of the number of annulus crossings in the training quadrant only (day 7, U = 210.5; p = 0.602; day 11, t(29) = 1.14; p = 0.263; see Supplementary Material, Figure S1).

Finally, the data were used to calculate accuracy scores, based on a comparison of the number of platform crossings in each of the four quadrants (accuracy score = TRA – [(Opp + AdjL + AdjR)/3], where TRA, Opp, AdjL, and AdjR are the number of platform crossings in the training, opposite, adjacent left and adjacent right quadrants respectively. Again, there were no group differences in either of the probe tests (day 7, F < 1; p > 0.20; day 11, F(1,29) = 2.40; p = 0.132).

**Experiment 5: Impaired appetitively motivated SRM but spared averishly motivated SRM in the Y-maze**

Experimentally naïve GRM2/3−/− (n = 25) and wild-type (n = 25) mice were compared on two different versions of the Y-maze SRM task: one appetitively motivated (as in Expt. 1) and the other, an averishly motivated “swimming Y-maze (water escape)” task. All mice were motivated to escape from the water and readily learned the aversive task. GRM2/3−/− mice were unimpaired in the aversive version of the Y-maze SRM task (no effect of group: F < 1; p > 0.20; effect of block: F(5,240) = 25.61; p < 0.001; no group*block interaction: F < 1; p > 0.20) (Fig. 3A). In a probe test conducted on day 7, in which the platform was removed from the maze and the animals allowed to swim freely for 30 s, both groups of mice concentrated their search in the arm that had previously held the platform: wild-types spent 55.2 ± 2.0 % of total search time in the correct arm, t(24) = 2.53; p = 0.018 compared to chance; GRM2/3−/− mice spent 61.8 ± 2.6 % of total search time in the correct arm, t(24) = 4.49; p < 0.001 compared to chance). In fact, there was a trend for GRM2/3−/− mice to spend a greater proportion of time in the correct arm than wild-type animals (t(48) = 2.00; p = 0.051) (Fig. 3A; inset).
In contrast, GRM2/3−/− mice once again displayed slower acquisition of the appetitive Y-maze SRM task (effect of group: $F_{(1,48)} = 19.01; p < 0.001$; effect of block: $F_{(8,384)} = 173.15; p < 0.001$; group*block interaction: $F_{(8,384)} = 7.91; p < 0.001$; n = 25 in each group; Fig. 3B). These results are consistent with a dissociation between appetitive and aversive spatial memory task performance in GRM2/3−/− mice.

**Experiment 6: Unimpaired appetitive visual discrimination learning**

To examine whether the GRM2/3−/− deficit extended to an appetitively motivated, non-spatial task, GRM2/3−/− (n = 14) and wild-type (n = 18) mice were compared on a simple “gray” versus “black/white stripes” visual discrimination task in the T-maze. A two-way RM-ANOVA revealed an effect of block ($F_{(4,120)} = 42.22; p < 0.001$), but no effect of group and no group*block interaction (both $F < 1; p > 0.20$) (Fig. 4A). Post-choice reinforcement in block 5 confirmed that mice were not locating the milk by virtue of its odor. This demonstrates therefore that GRM2/3−/− mice do not display a general deficit on all appetitively motivated maze tasks.

**Experiment 7: Impaired spontaneous spatial novelty preference**

This test of spatial memory is neither appetitively nor aversively motivated, but relies upon animals’ natural exploratory drive. If the deficit in GRM2/3−/− mice simply reflects a reduced appetitive motivation for food rewards, then one would expect normal memory performance on this spatial novelty preference task. During the initial exposure phase, GRM2/3−/− (n = 13) and wild-type (n = 18) mice spent a similar amount of time exploring the two available arms (the “start” and “other” arms) ($F_{(1,29)} = 1.632; p = 0.212$; no group*arm interaction: $F < 1; p > 0.20$). There was a tendency for GRM2/3−/− mice to make fewer arm entries in total, however ($t_{(29)} = 2.05; p = 0.05$), suggesting that they were less active. GRM2/3−/− mice made fewer entries into the “start” arm than
wild-type animals \( t_{(29)} = 2.43; \ p = 0.022 \), but importantly the number of entries into the “other” arm did not differ between the groups \( t_{(29)} = 1.57; \ p = 0.128 \).

In the subsequent test phase, both groups displayed a preference for the previously unvisited, “novel” arm over the now familiar, “other” arm (Fig. 4B). This preference was stronger in the wild-type mice. Statistical analysis of the discrimination ratio \([(\text{novel arm} / (\text{novel} + \text{other arm})] confirmed that wild-type mice showed a preference for the “novel” arm that was above chance \( (\text{chance} = \text{discrimination ratio of 0.50}) \), both in terms of arm entries \( t_{(17)} = 9.54; \ p < 0.001 \) and time spent in arms \( t_{(17)} = 9.63; \ p < 0.001 \). The discrimination ratio for the GRM2/3\(^{-/-}\) mice was also above chance, again in terms both of arm entries \( t_{(12)} = 3.59; \ p = 0.004 \) and time spent in arms \( t_{(12)} = 2.94; \ p = 0.012 \). Importantly, the “novel” arm preference was stronger in the wild-type mice than in GRM2/3\(^{-/-}\) mice: wild-type mice made a higher proportion of “novel” arm entries \( t_{(29)} = 2.64; \ p = 0.013 \) and spent a significantly greater proportion of time in the “novel” arm \( t_{(29)} = 3.05; \ p = 0.005 \) than GRM2/3\(^{-/-}\) mice. Taken in conjunction with the result of experiment 6, this shows that the GRM2/3\(^{-/-}\) cognitive deficit cannot be attributed to a reduction in appetitive motivation.

**Experiment 8: GRM2/3 knockout mice display reduced spontaneous and amphetamine-induced locomotor activity**

GRM2/3\(^{-/-}\) mice were hypoactive relative to wild-type controls. A photocell beam breaks test, conducted in a novel cage environment and lasting for 2 hr (24 x 5 min blocks), revealed less spontaneous locomotor activity in GRM2/3\(^{-/-}\) mice \( (\text{effect of group: } F_{(1,30)} = 8.20; \ p = 0.008; \ \text{effect of block: } F_{(23,690)} = 32.66; \ p < 0.001; \ \text{no group*time interaction: } F_{(23,690)} = 1.35; \ p = 0.127; \text{18 wild-types vs. 14 GRM2/3\(^{-/-}\) mice; Fig. 5A}).

In a separate cohort of wild-type \( (n=17) \) and GRM2/3\(^{-/-}\) mice \( (n=22) \) locomotor activity was tested over a 70 hr period \( (70 x 1 \text{ hr blocks}) \) in a threshold activity monitoring system (see
ANOVA revealed an overall main effect of group \((F(1,37) = 5.94; \ p < 0.05)\) which reflected the fact that wild-type mice were more active overall than the knockouts. There was also a main effect of block \((F(69, 2553) = 46.1; \ p < 0.0001)\) which reflected the changes in diurnal activity over the course of the experiment, and a significant group by block interaction \((F(69, 2553) = 2.09; \ p < 0.0001)\). Analysis of simple main effects revealed that the activity levels of the groups differed significantly between 1-4 hr after first being placed in the novel activity cages, and then again consistently during the middle of the dark phase on each of the three days of continual testing in these cages (see Supplementary Material and Supplementary Figure S2 for full details). On 13 out of 14 time blocks in which the two groups differed, the wild-type mice were more active than the GRM2/3\(^{-/-}\) mice.

A moderate dose of amphetamine (2.5 mg/kg) increased activity levels in both GRM2/3\(^{-/-}\) \((n = 10)\) and wild-type \((n = 12)\) mice by approximately 100%. GRM2/3\(^{-/-}\) mice therefore remained hypoactive relative to wild-type controls after this amphetamine treatment \((\text{effect of group: } F_{(1,20)} = 4.85; \ p = 0.039; \text{effect of drug: } F_{(1,20)} = 15.94; \ p < 0.001; \text{no group*drug interaction: } F < 1; \ p > 0.20)\) (Fig. 5B(i) and (ii)).

Following injection of 10 mg/kg amphetamine, wild-type mice \((n = 5)\) displayed an initial burst of activity (as did wild-type saline-injected controls; \(n=6\); Fig. 5C). Activity levels then fell sharply, well below the activity levels in the wild-type saline controls, concomitant with an increase in observed stereotypical behaviors (Supplementary Material, Supplementary Fig. S3). Approximately 2.5 hours post-injection, activity levels in the wild-type amphetamine group began to rise again as stereotypy decreased. For the next 3-4 hrs activity levels in the wild-type amphetamine mice were considerably greater than in the wild-type saline animals, before returning back to wild-type saline levels between 6 and 7 hr post-injection. The initial burst of activity was
notably reduced in the GRM2/3−/− mice (amphetamine: n = 6; saline: n = 6) compared to wild-types. Similar to wild-type mice, however, activity levels in amphetamine treated GRM2/3−/− mice declined sharply as stereotypical behaviors increased, remaining low until approximately 2.5 hours post-injection, when they then began to increase again as stereotypies reduced, paralleling the change in the wild-type mice. The increase in locomotor activity observed between 2.5 and 6.5 hr post-injection of amphetamine was substantially less in the GRM2/3−/− mice compared to wild-types and, importantly, at no point did locomotor activity in GRM2/3−/− mice exceed that of wild-type mice. Statistically, these observations reflected an overall effects of drug (F(1,19) = 14.07; p = 0.0014), genotype (F(1,19) = 8.23; p = 0.0098), and time bin (F(27, 513) = 5.81; p < 0.001). There was a drug*time bin interaction (F(27,513) = 8.27; p < 0.001), but neither drug*genotype nor genotype*time bin interactions (both F < 1; p > 0.20). There was, however, a drug*genotype*time bin interaction (F(27,513) = 2.00; p = 0.002), indicating that wild-type and GRM2/3−/− mice responded differentially to 10mg/kg amphetamine at different time points post-injection.

Experiment 9: Injection stress has differential effects on T-maze SWM performance in GRM2/3−/− mice and wild-type mice

Finally, we examined whether an amphetamine-induced increase in arousal would affect spatial memory performance in the T-maze rewarded alternation task. A low dose (0.5 mg/kg) of amphetamine was used to avoid any risk of aphagia. A within-subjects analysis comparing performance after either amphetamine or saline injection revealed a main effect of genotype (F(1,21) = 6.24; p = 0.02), as GRM2/3−/− mice performed less well than wild-type controls in both injection conditions. There was, however, no effect of drug condition (i.e., amphetamine versus saline - F < 1; p > 0.2), and no genotype*drug condition interaction (F < 1; p > 0.2).
While the effects of 0.5 mg/kg amphetamine on SWM performance did not differ from those of saline, we observed that the injection stress itself differentially affected the performance of wild-type and GRM2/3⁻/⁻ mice, producing impairment in the former but facilitating performance in the latter. We therefore performed a further comparison of “injected” (including both saline and amphetamine data) versus “no-injection” performance (including (i) the last day of training on the task prior to any injection and (ii) the final day of testing performed 24 hr after the last injection). Thus, injected and no-injection performance were compared using a counterbalanced “A-B-B-A” design (where “A” = no injection and “B” = injected). This comparison revealed a main effect of genotype (F(1,21) = 18.95; p = 0.0003), as GRM2/3⁻/⁻ mice remained impaired overall relative to wild-types, but no main effect of condition (i.e., injected versus no-injection) (F < 1; p > 0.20). Importantly, however, there was a significant genotype*condition interaction (F(1,21) = 5.64; p = 0.027), indicating that the two groups were responding differentially to the injection stress. Simple main effects analysis showed that the opposite directions of changes in performance in wild-type versus GRM2/3⁻/⁻ mice post-injection were responsible for the significant interaction, as there was no significant effect of condition for either group individually (wild-types: F(1,21) = 1.96; p = 0.18; GRM2/3⁻/⁻: F(1,21) = 3.80; p = 0.065; Fig. 6).

**DISCUSSION**

GRM2/3⁻/⁻ mice demonstrated a novel fractionation of hippocampus-dependent spatial memory. They were impaired in appetitively motivated spatial tasks (Expts. 1-3) but not in the aversively motivated open field water maze (Expt. 4). This dissociation was confirmed using experimentally naïve mice in fully counterbalanced appetitive and aversive versions of the Y-maze SRM task (Expt. 5). GRM2/3⁻/⁻ mice were not impaired on all appetitively motivated tasks, however, displaying normal acquisition of a non-spatial, visual discrimination task (Expt. 6). Furthermore,
GRM2/3⁻/⁻ mice showed reduced spatial memory for recently visited arms in a Y-maze test of spontaneous, spatial novelty preference that does not involve a food reward, but instead relies on animals’ exploratory drive (Expt. 7). Therefore, the deficits do not simply reflect reduced appetitive motivation for the milk reward. Instead, the pattern of results suggests a change in the arousal-cognition function in GRM2/3⁻/⁻ mice. This fractionation of spatial memory in GRM2/3⁻/⁻ mice is, to our knowledge unique. It is, for example, very different from the phenotypes seen in ionotropic glutamate receptor subunit knockout mice which exhibit fractionated spatial memory on the basis of different aspects of information processing, rather than on the arousing properties of the experimental paradigms. For example, deletion of GluA1 (Reisel et al, 2002; Schmitt et al, 2003) or GluN2A (Bannerman et al, 2008) dissociate SWM from SRM.

A well-established inverted U-shaped function relates levels of acute stress and arousal with performance on cognitive tasks; that is, low levels of stress increase levels of attention and arousal, and enhance cognitive performance relative to the “non-stressed” state. Above a certain threshold, however, higher levels of stress become deleterious (Yerkes and Dodson, 1908). By this account, on low stress, appetitive maze tasks, both groups likely sit on the rising phase of the inverted U-shaped function, with arousal levels in GRM2/3⁻/⁻ mice hypothesized to be below those of wild-type animals, resulting in spatial memory deficits. During aversive tasks like the water maze, GRM2/3⁻/⁻ mice move closer to the peak of the inverted U function, improving their performance and catching up with that of the controls. Importantly, injection stress also facilitated performance in GRM2/3⁻/⁻ mice but actually impaired spatial memory performance in wild-type mice (Expt. 9). Thus, injection stress also moved GRM2/3⁻/⁻ mice to the right on the arousal-cognition function, but, at the same time, moved wild-type animals beyond the peak of the inverted U, thereby impairing their performance.
In keeping with the possibility of altered arousal levels in the GRM2/3−/− mice, they displayed locomotor hypoactivity relative to wild-type controls, in both the absence and presence of amphetamine (Expt. 8). Furthermore, in an assessment of diurnal activity levels across a 70 hr period, the GRM2/3−/− mice were consistently less active during the middle of the dark phase. However, the relationship between activity levels and arousal is likely to be complex and it is notable that in a different situation, namely spatial working memory testing on the T-maze, it might be argued that the GRM2/3−/− mice were more active than the wild-types, in demonstrating shorter running latencies on both sample and choice runs. Again, this may be consistent with a biphasic relationship between mGluR activity and behaviour.

A role for mGluR2/3 activation in the inverted U-shaped relationship between arousal and cognition could, in addition to accounting for the results of the present study, also accommodate previous, seemingly paradoxical, pharmacological data. While the mGluR2/3 agonist LY354740 impaired T-maze rewarded alternation in normal animals (Aultman and Moghaddam, 2001), the same drug actually reversed the SWM performance deficit produced by PCP when co-administered on the same task (Moghaddam and Adams, 1998). At first glance, these opposing effects seem counter-intuitive. However, if under normal conditions for appetitive maze tasks, baseline glutamate and arousal levels lie on the ascending limb of the inverted-U function, then an agonist-induced reduction in glutamate release would be expected to impair cognition (Aultmann and Moghaddam, 2001; Gregory et al, 2003; Higgins et al, 2004; Sato et al, 2004; Lin et al, 2005; Daumas et al, 2009). Conversely, under circumstances of enhanced glutamate levels, as produced by PCP, (i.e. to the right of the peak), then the mGluR2/3 agonist-induced reduction of glutamate (Moghaddam and Adams, 1998) would be predicted to improve cognitive performance (Aultmann and Moghaddam, 2001). Of note, Marek et al, (2010) also recently postulated a biphasic dose-effect
function between group II mGluRs and cognition. Our findings provide empirical support for the notion, and extend it by suggesting that arousal may be one of the mediating variables.

Although an ‘inverted U’ is an attractive model, it remains speculative. It will be important to identify the pathways and substrates by which the proposed interaction between arousal and cognition occurs, since the deficits in the GRM2/3−/− mice are unlikely to arise entirely from the hippocampus, nor solely from an effect on glutamate signaling. Thus, although the affected memory tasks are hippocampus-dependent, and it has been suggested that the hippocampus may be critical for the interplay between arousal and cognition (Gray and McNaughton, 2000), interconnected regions may also be involved. Group II mGluRs are expressed widely in limbic and forebrain regions (Gu et al., 2008). Notably, they may also play an important role in modulating arousal through effects on the ascending reticular activating system, and the associated midline and intralaminar thalamic nuclei. Within the thalamus, mGluR2 is preferentially expressed in these nuclei, and mGluR3 is relatively selectively expressed in GABAergic neurons of the reticular nucleus (Ohishi et al, 1993, 1998). Furthermore, a significant portion of neocortical group II mGluRs are on thalamocortical afferents (Marek et al., 2001). Thus, modulation of the reticular formation and associated thalamocortical pathways may be a prime target for mGluR modulation of arousal states.

Similarly, there may be downstream effects on other neurotransmitter systems beyond glutamate. Dopamine is one candidate, given its roles in hippocampal cognition (Lisman and Grace, 2005; Shohamy and Adcock, 2010) and in stress and arousal (Pani et al, 2000; Ungless et al, 2010). More specifically, group II mGluRs modulate dopamine release (Hu et al, 1999; Cartmell et al, 2000; Chaki et al, 2006; Pehrson and Moghaddam 2010), possibly with a biphasic dose relationship (Hu et al, 1999). In addition, there are interactions between GRM3 and the dopamine metabolizing enzyme catechol-O-methyltransferase (COMT) which affect working memory and cortical function.
(Tan et al., 2007). Indeed, dopamine has its own inverted U relationship with cognition, with dopaminergic manipulations, including response to amphetamine, having differential effects depending on the starting position on the curve (Arnsten, 1998; Mattay et al., 2003; Williams and Castner, 2006). It will be of interest to determine whether, and in what respect, the present findings reflect an overlap between the ‘arousal’ and ‘dopamine’ inverted U functions; preliminary data indicate changes in dopamine metabolism in the GRM2/3−/− mice (unpublished observations).

Equally, other potential mediators of the putative altered arousal-cognition relationship in GRM2/3−/− mice should not be overlooked, notably norepinephrine (Swanson et al., 2004; Lorrain et al., 2005; Nikiforuk et al., 2010), serotonin (Cartmell et al., 2000; Nikiforuk et al., 2010), and the HPA axis (Scaccianoce et al., 2003).

The main limitation of this study is that the generation of a sufficiently large number of double knockout animals necessitated, for ethical and financial reasons, the use of separate breeding lines and thus non-littermate wild-type controls. This is a common practice in double knockout mouse studies, but means that we cannot rule out the possibility that subtle differences in genetic background or epigenetic factors between GRM2/3−/− and wildtype mice could have influenced the results. However, several considerations provide some reassurance in this regard: the findings were seen in several cohorts, including the replication of key results in ania arising from two different backcrosses, and the replication of key results across these; the relatively large sample sizes and statistical significance of the key results, and the replication of key results together with and, increases our confidence. A second limitation is that we do not know the relative contributions of GRM2 and GRM3 to the findings. This information would help address some of the mechanistic questions raised above, given the differences in properties and functions of the two receptors (Harrison et al., 2008); however, repeating the present experiments in GRM2−/− and GRM3−/− mice might not prove conclusive, in light of the partial compensation that occurs in
expression of the remaining receptor (Lyon et al., 2008). **Further studies are also needed to determine whether the observed phenotype arises from the developmental consequences of the absence of group II mGluRs, and to clarify the downstream effects on other aspects of glutamate transmission.**

Finally, our results have translational relevance for the range of disorders in which group II mGluRs have been implicated, and for which stress responsivity and motivation are important, such as anxiety, mood, and substance use disorders (Swanson et al., 2005; Witkin and Eiler, 2006; Markou, 2007). The greatest relevance, however, may be for schizophrenia (Moghaddam, 2004; Krystal et al., 2010), a disorder characterized not only by psychotic symptoms, but by disturbances in cognition, attention and motivation. These various domains are sometimes simplistically considered in isolation, but are increasingly appreciated and conceptualized as being intertwined, and likely sharing overlapping substrates (e.g. Kapur, 2003; Barch, 2005; Javitt, 2009). Our results suggest that group II mGluRs may be involved at the interface between these processes.

In summary, mice lacking group II mGluRs have impairments in appetitive but not aversive hippocampal memory tasks, and show a differential response to amphetamine and injection stress compared to wildtype mice. These findings suggest a change in the relationship between arousal and cognition, possibly explained in terms of an altered ‘inverted U’. This attractive but speculative proposal, which was recently proposed in a more general form for the impact which stress has upon glutamate pathways and thence memory function (Sandi, 2011), awaits refinement and critical testing. Such a model may help reconcile the seemingly conflicting results of studies using group II mGluR ligands, since their effects will likely be state-dependent. Our findings also bear upon the translational interest in group II mGluRs across a broad range of psychiatric disorders, all of which involve arousal, cognition, and their interplay.
DISCLOSURE/ CONFLICTS OF INTEREST

Mice were bred, genotyped, and supplied by GlaxoSmithKline under an unrestricted agreement with the University of Oxford, where all experiments were conducted. Dr Bannerman is a consultant to Eli Lilly and a member of their Centre for Cognitive Neuroscience. Dr Harrison in the last three years has received honoraria for lectures, chairing meetings, or advisory boards, from AstraZeneca, Bristol Myers Squibb, Janssen, Merck, Otsuka, and Wyeth. Drs Kew and Corti are employees of GlaxoSmithKline.

ACKNOWLEDGEMENTS

Work supported by a Wellcome Trust Ph.D. studentship awarded to Louisa Lyon, and grants to David Bannerman from the Wellcome Trust (074385 and 087736). Additional support from the UK Medical Research Council. The authors thank Prof. Shigemoto Nakanishi for generous provision of GRM2⁻/⁻ mice, used to produce the GRM2/3⁻/⁻ mice.
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Fig. 1. GRM2/3<sup>−/−</sup> mice were impaired in appetitive SRM (left panels, A and C) and SWM tasks (right panels, B and D). A: GRM2/3<sup>−/−</sup> mice (n=14) displayed slower acquisition of a Y-maze SRM task than wildtype (wt; n=18) mice and B: GRM2/3<sup>−/−</sup> mice (n=14) were persistently impaired in an appetitive T-maze rewarded alternation SWM task compared to wild-types (wt; n=18). Data shown are percent correct responses (mean±sem) for each block of 10 trials. Chance performance is indicated by the dashed line. C: GRM2/3<sup>−/−</sup> mice (n=13) displayed slower acquisition of the SRM component of an appetitive six-arm radial-arm maze task than wt (n=13) and D: GRM2/3<sup>−/−</sup> mice subsequently made significantly more SWM errors than wt. Data shown are mean errors (± sem) per trial in each block of 4 trials. Asterisks indicate statistical significance at p < 0.05.

Fig. 2. GRM2/3<sup>−/−</sup> mice (n = 13) did not differ from wild-type mice (wt; n=18) in the water maze SRM task. A: path length for each day of testing during acquisition. B: percentage time spent in the adjacent left (AdjL), training (TRA), adjacent right (AdjR), and opposite (Opp) quadrants in probe test 1 (day 7). C: percentage time spent in each quadrant in probe test 2 (day 11). Chance performance is indicated by the dashed line. Values are mean ± sem.

Fig. 3. GRM2/3<sup>−/−</sup> mice show a dissociation between performance on appetitive and aversive spatial memory tasks. A: GRM2/3<sup>−/−</sup> mice (n=25) learned the swimming SRM Y-maze task at the same rate as wild-type mice (wt; n=25). Data shown are mean percent correct responses (± sem) for each block of 5 trials. Inset: In a probe test on day 7, GRM2/3<sup>−/−</sup> mice spent an equal or greater (p=0.051) proportion of their time searching in the arm that had previously held the platform, relative to wild-type mice. B: GRM2/3<sup>−/−</sup> mice (n=25) displayed slower acquisition of the appetitive SRM Y-maze task than wild-type controls (n=25). Data shown are mean percent correct responses (± sem) for
each block of 10 trials. Order of testing and the rooms in which the tests were performed were fully counterbalanced. Chance performance is indicated by the dashed line. *p < 0.05.

**Fig. 4.** Cognitive deficits in GRM2/3−/− mice cannot be explained simply by reduced appetitive motivation for the milk reward. A: GRM2/3−/− mice (n=14) did not differ from wild-type mice (wt; n=18) in acquisition of an appetitively motivated, non-spatial gray versus black-white stripes visual discrimination task (Expt. 6). Data shown are mean percent correct responses (± sem) for each block of 10 trials. Chance performance is indicated by the dashed line. B: GRM2/3−/− mice (n=13) show a reduced spatial novelty preference compared to wild-type animals (n=18; Expt 7). Data shown are mean discrimination ratios [(novel arm/ (novel + other arm)] (± sem) calculated for the number of arm entries (left) and the time spent in the arms (right) during the test phase. The discrimination ratio was significantly lower for GRM2/3−/− mice on both measures. A discrimination ratio of 0.5 reflects chance performance. (* p < 0.05 and ** p = 0.005).

**Fig. 5.** GRM2/3−/− mice are hypoactive. A: GRM2/3−/− mice (n=14) displayed less spontaneous locomotor activity (LMA) than wild-type mice (wt; n=18). B: 2.5 mg/kg amphetamine increased LMA in both wild-type (n=10) and GRM2/3−/− mice (n=10) mice, such that GRM2/3−/− mice remained hypoactive relative to wild-types. (i) summarises the total number of beam breaks over a 2 hr period while (ii) shows the time course of changes in LMA. C: 10 mg/kg amphetamine induced stereotypy in both wt (n=6) and GRM2/3−/− (n=6) mice, resulting in a decrease in LMA followed later by an increase in activity levels relative to saline-injected controls (n=5 wt and n=6 GRM2/3−/− mice). At all time points GRM2/3−/− mice were hypoactive relative to wild-types.
Fig. 6. Effect of injection stress on the appetitive T-maze SWM task. Injection stress impaired performance in the wild-types (wt; n=12) but facilitated it in GRM2/3^{−/−} mice (n=11).
