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Involvement of AMPA Receptor GluR2 Subunits in Stimulus–Reward Learning: Evidence from Glutamate Receptor gria2 Knock-Out Mice

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Presence of the glutamate receptor 2 (GluR2) subunit prevents calcium influx through AMPA-receptor complexes; deletion of this subunit results in enhanced hippocampal long-term potentiation. We investigated whether mice lacking the GluR2 subunit [gria2 knock-out (KO) mice] displayed impairments in learning stimulus–reward associations, and the subsequent ability of reward-paired cues to control motivated behavior. Both gria2 KO and wild-type (WT) mice learned to associate a light/tone stimulus with food delivery, as evidenced by approach toward the food magazine after the presentation of the cues (pavlovian conditioning). Subsequently, the cues also served to reinforce an operant response in both KO and WT mice (conditioned reinforcement), although response rates were greater in gria2 KOs. Responding for conditioned reinforcement was enhanced after 0.5 mg/kg amphetamine administration in WT mice, but not in KO mice. The ability of the cues to elicit approach behavior (conditioned approach) and to enhance responding for the reward (pavlovian-to-instrumental transfer; PIT) were also impaired in gria2 KO mice. This pattern of behavior resembles that seen after lesions of the central nucleus of the amygdala (CeA), an area rich in GluR2-containing AMPA receptors. Immunostaining revealed reduced GluR1 expression within both the basolateral amygdala and the CeA, suggesting that the behavioral deficits observed were unlikely to be caused by compensatory changes in GluR1. These results suggest that GluR2-containing AMPA receptors, possibly within the CeA, are critical for the formation of stimulus–reward associations necessary for PIT and conditioned approach, but are not involved in the plastic processes underlying the attribution of motivational value to the conditioned stimulus (CS).

Key words: learning; pavlovian association; conditioned reinforcement; pavlovian to instrumental transfer; pavlovian approach; discriminated approach; AMPA receptor; GluR-A; GluR-B; gria1; amygdala; amphetamine

Introduction
Conditioned associations between environmental stimuli and rewarding events are important in controlling and maintaining appropriate behavioral responses; they may also contribute to aberrant behaviors, including drug addiction. Cues associated with drug taking initiate and control behavior in both abstaining addicts and animal models of drug abuse, increasing drug craving (Childress et al., 1999) and drug seeking (Shaham et al., 2003). Consequently, treatment strategies for relapse prevention include the removal of an addict from situations in which exposure to drug-paired cues is likely (O’Brien et al., 1998). Therefore, understanding the mechanisms that underlie the formation and expression of such associations has both theoretical and practical importance.

Models of associative learning implicate glutamatergic neurotransmission through ionotropic NMDA and AMPA receptors as the molecular basis of learning, long-term potentiation (LTP) (for review, see Nicoll, 2003). In particular, the expression of LTP is associated with enhanced glutamatergic transmission through AMPA receptors, of which GluR1 (encoded by the gria1 gene) and GluR2 (encoded by the gria2 gene) subunits are major components. Because drug addiction has been viewed as a form of aberrant learning (Everitt et al., 2001), the glutamate system is a prime candidate for studies of processes underlying addiction. Previously, we have reported that mice lacking the GluR1 subunit of the AMPA receptor [gria1 knock-out (KO) mice] display specific deficits in stimulus–reward learning (Mead and Stephens, 2003). Although they are capable of forming an association between a discrete cue and the presentation of a food reward, as evidenced by their ability to use the cue as a discriminative stimulus (pavlovian conditioning), show approach to the cue (conditioned approach; Tomie et al., 1999), and are responsive to the ability of the cue to enhance ongoing operant behavior [pavlovian-to-instrumental transfer (PIT); Dickinson, 1994], when presented with the opportunity to obtain the cue through an instrumental response (conditioned reinforcement; Mackintosh, 1974), gria1 KOs failed to respond (Mead and Stephens, 2003). This deficit, an inability to attribute motivational value to the cue, was also reflected in a deficit in responding under a second-order schedule of reinforcement (Mead and Stephens, 2003), a task also dependent on the conditioned reinforcing...
properties of the cue (Whitelaw et al., 1996). This pattern of deficits mirrors that seen after the occurrence of lesions of the basolateral region of the amygdala (BLA) (Everitt et al., 2000), an area also rich in GluR1 expression (McDonald, 1996), leading us to suggest that the deficits observed in gria1 KO mice may be caused by impaired neurotransmission in or via the BLA (Mead and Stephens, 2003).

However, deletion of GluR1 subunits in the gria1 KO mouse affects the distribution of other AMPA-receptor subunits in both the hippocampus (Zamanillo et al., 1999) and the BLA (Mead and Stephens, 2003) [although not the neighboring central nucleus of the amygdala (CeA)]. Thus, it was unclear whether the deficits observed in the gria1 KO may be caused by associated changes in GluR2. Therefore, the present set of experiments investigated the effects of GluR2 deletion on stimulus–reward learning, using the gria2 KO mouse (Jia et al., 1996). This mutant shows facilitated AMPA-dependent LTP in hippocampal slices, and enhanced calcium permeability of AMPA receptors, consistent with the known role of GluR2 in preventing calcium influx through AMPA-receptor-gated channels (Jia et al., 1996). Here, we investigate the effects of GluR2 deletion on pavlovian conditioning, conditioned approach, PIT, and conditioned reinforcement, in addition to examining possible changes in GluR1 expression within the amygdala subregions.

Materials and Methods

Animals. Wild-type (WT) and gria2 KO littermates were bred at the University of Sussex from heterozygous parents obtained from The Jackson Laboratory (Bar Harbor, ME; strain name, STOCK Gria2tm1Rodl, stock number, 002913). The genotypes of the offspring were identified using PCR analysis. Mice were housed two or three to a cage and were allowed at least 2 weeks of habituation to the home cages before the beginning of experimental sessions. Holding rooms were maintained at a constant temperature (21 ± 2°C) and humidity (50 ± 10%), and lights were on for 12 hr starting at 7:00 A.M.). Except when specified, mice were allowed access to standard lab chow and water ad libitum. Between five and eight WT and gria2 KO mice were used in each phase of the experiment. All testing took place during the light phase, between 8:00 A.M. and 6:00 P.M. The experiment used a within-subject design, whereby all mice were used for each stage of the study in the order described. All experiments were approved by the institutional ethics committee and were performed under United Kingdom legislation on animal experimental (Animal Scientific Procedures Act, 1986).

Pavlovian conditioning. Mice were food-restricted to maintain their body weight at ~85% of free feeding body weight. During 2 hr sessions, mice were placed into mouse operant chambers (Med Associates, E. Fairfield, VT) with the levers retracted, and 20 mg food pellets (Noyes Precision pellets, Formula P; Research Diets) on a fixed-ratio (FR1) schedule during a 120 min session. The schedule was then progressively increased to an FR10 schedule. After acquisition, the schedule was changed to a VI 30 sec schedule, and increased daily to a VI 120 sec schedule. After 3 d on a VI 120 sec schedule, mice were tested on a series of different VI schedules (VI 30, 60, 120, 240, 360, and 480 sec) in a random order.

PIT. To assess PIT, mice from the instrumental responding stage were retrained on a VI 240 sec schedule for one session of 120 min. Both KO and WT mice responded at a stable rate on this schedule, although the overall rate of lever pressing was higher in KO mice. This VI schedule was chosen for PIT testing because it was at this schedule that WT mice responded at the lowest rates. The following day, mice were again allowed to respond on a VI 240 sec schedule for food pellets for a total of 60 min. In addition, an extended 60 sec CS was presented on a fixed-interval 5 min schedule (cue location and conditions exactly as for pavlovian conditioning stage). The rate of responding was then compared during the presence of the cues (CS period) and the absence of the cues (VI period).

Immunohistochemistry. For immunohistochemical analysis of GluR1, adult mice were anesthetized with tri bromoethanol (Avertin, 20 mg/kg) and transcardially perfused with 4% paraformaldehyde. Brains were removed and stored in 4% paraformaldehyde for 24 hr, before placement in 0.1 M phosphate buffer containing 30% sucrose for 48 hr. Brains were then frozen in isopentane at −45°C, and stored at −80°C until sectioning. Coronal sections (30 μm) were taken using a cryostat, and sections were washed in PBS. Endogenous peroxidase was quenched by immersion in 0.3% hydrogen peroxide, and sections were washed in PBS before blocking in 1.5% normal goat serum (Vector Laboratories, Peterborough, UK). After additional washing in PBS, sections were incubated in 0.25 μg/ml anti-GluR1 (66–306; Upstate UK, Botolph Claydon, UK) overnight. Sections were then washed in PBS, and incubated in a 1:600 dilution of biotinylated secondary antibody (BA–1000; Vector Laboratories) for 60 min, before being washed again. Sections were subsequently incubated in ABC complex (Vectastain ABC elite kit: PK6100; Vector Laboratories), and washed in PBS; staining was visualized using the nickel-DAB glucose (D-5637 and G-2133; Sigma Aldrich, Gillingham, UK).
UK) method. Sections were slide-mounted, dehydrated, and cover-slipped before analysis. For the analysis of sections, images were captured using a Sony (Tokyo, Japan) DSC-S75 digital camera mounted on a Zeiss (Oberkochen, Germany) Axioskop 2 microscope. Negative controls for optical density analysis standardization were run using exactly the same protocol except that the primary antibody was omitted.

Statistical analysis

Pavlovian conditioning. Two measures were analyzed to assess pavlovian conditioning behavior. First, the latency between cue onset and reward retrieval (latency) was compared between genotypes; second, the percentage of total food-magazine entries occurring during the CS presentation (CS%) was compared. Two-way ANOVA was performed, with training sessions (within subjects) and genotype (between subjects) as factors. Post hoc analysis was performed using independent-sample *t* tests.

Conditioned approach. For analysis of conditioned approach, nose-poke rates into the CS aperture were assessed during the CS (CS+) and compared with rates when the CS was not presented (CS−). Similarly, nose-poke rates into the CTRL aperture were compared. Data for this measure were square root transformed to gain homogeneity of variance and to allow parametric analysis. Three-way ANOVA was performed, with CS state (either CS+ or CS−), aperture (within subjects) and genotype (between subjects) as factors.

CS components. For the analysis of CS component data, latency and CS% were measured. Two-way ANOVA was performed, with CS condition (within subjects) and genotype (between subjects) as factors. When appropriate, post hoc analysis was performed using the Student–Newman–Keuls test.

Conditioned reinforcement. For the analysis of CR, the total number oflever presses was used as the dependent variable. Three-way ANOVA was performed, with lever and drug treatment (within subjects) and genotype (between subjects) as factors. Post hoc analysis was performed using repeated measures or independent-sample *t* tests when appropriate. In addition, ANOVA was also performed with testing order as a factor. This analysis was performed to rule out the possibility that the order of drug treatment influenced responding for conditioned reinforcement, because previous work has shown that even a single amphetamine administration can enhance responding on subsequent tests for CR (Mead et al., 2003).

Instrumental responding. For the analysis of instrumental responding, the number of food pellets received and total lever presses during the session were taken as the dependent variables. Two-way ANOVA was performed, with VI schedule (within subjects) and genotype (between subjects) as factors. Post hoc analysis was performed using independent-sample *t* tests.

PIT. For the analysis of PIT, response rates during the CS were divided by response rates during the VI period to produce a ratio (for which a ratio of 1 indicates identical response rates during the CS and in the absence of the CS). Two-way ANOVA was performed, with lever (within subjects) and genotype (between subjects) as factors. Post hoc comparisons were made using repeated-measures *t* tests. Magazine approach during the PIT session was also analyzed by comparing nose-poke rates into the magazine during the CS with rates during the intervening VI period. Two-way ANOVA was performed, with CS state (within subjects) and genotype (between subjects) as factors. Post hoc comparisons were made using repeated-measures *t* tests.

Immunohistochemistry. The analysis of GluR1 immunoreactivity was performed by counting the number of GluR1-positive soma and analyzing optical density within a 130 x 170 mm region of the BLA and CeA. The regions selected are indicated in Figure 5G, and represent regions from within the basolateral amygdaloid nucleus and the central amygdaloid nucleus (Franklin and Paxinos, 1997). For the quantitative analysis of GluR1-positive soma, two independent observers scored each section, and were unaware of the condition. For the analysis of optical density of GluR1 staining, mean optical density values were obtained using Scion-Image (Scion Corp., Frederick, MD), rating each pixel with a value of 0 to 255. Mean optical densities for each section were then corrected for nonspecific binding by subtracting optical density values from negative controls. Two-way ANOVA was then performed with region (within subjects) and genotype (between subjects) as factors. Post hoc comparisons were performed using independent-sample *t* tests when appropriate.

Results

Pavlovian conditioning

Figure 1A indicates that when trained to associate a tone/light cue with the delivery of a food reward, by presenting the cues imme-
mediately before the randomly timed delivery of a reward, both WT and KO mice learned the association. This is indicated by an increase in the percentage of total nose-pokes into the food magazine during the CS presentation (CS%) (main effect of session, \(F_{(13,182)} = 25.14, p < 0.01\)). Differences were observed between genotypes during the later sessions only (session by genotype interaction, \(F_{(13,182)} = 3.12, p < 0.01\)), and post hoc analysis revealed that the CS% was significantly lower in KO mice for sessions 11–14 (\(p < 0.05\)). However, the analysis of total nose-pokes into the food magazine during the CS and intervening random interval (RI) periods revealed that the differences observed in sessions 11–14 were caused by a small difference in the number of irrelevant nose-pokes during the RI period, rather than a reduction in nose-pokes during the CS period (Fig. 1C). Analysis of reward retrieval latency after the CS onset (Fig. 1B) also indicated acquisition of the CS–unconditioned stimulus (US) relationship across sessions (main effect of session, \(F_{(13,182)} = 7.37, p < 0.01\)), although no between-genotype differences were observed for this measure.

**Conditioned approach**

Once the cue light had been relocated in the wall opposite to the food magazine, WT mice displayed conditioned approach toward the CS, specifically during the CS presentation (Fig. 2A). This effect was also specific to the CS location, because approaches into a control aperture were unaffected by the CS state (CS state by aperture interaction; \(F_{(1,7)} = 9.40, p < 0.05\)). In contrast, KO mice displayed no selective approach toward the CS aperture, and any approaches made were not related to the CS state (main effect of aperture, \(F_{(1,7)} = 0.39\), main effect of CS state, \(F_{(1,7)} = 0.51\); CS state by aperture interaction, \(F_{(1,7)} = 1.13\; \text{all not significant}\).

**CS components**

To determine whether the failure of KO mice to approach the CS was caused by a sensory impairment, we examined the ability of the individual cue components (light alone or tone alone) to elicit discriminated approach. The analysis of retrieval latency after the CS onset (Fig. 2B) revealed that the individual CS elements were equally effective as discriminative stimuli as the compound CS used previously (no significant effect of CS type, \(F_{(2,28)} = 1.86\), not significant; or CS by genotype interaction, \(F_{(2,28)} = 1.24\), not significant). The analysis of CS% revealed differences between genotypes and CS types, but no interaction (main effect of CS, \(F_{(2,28)} = 5.67, p < 0.01\); main effect of genotype, \(F_{(1,14)} = 4.95, p < 0.05\)). The reduced CS% observed in the KO mice for all CS types was consistent with that seen in the final four sessions of the discriminated approach stage. Post hoc analysis also revealed that both WT and KO mice displayed a reduced CS% when CS elements were presented alone (lights alone or tone alone) than when the compound CS was presented (tone plus lights). Although these results suggest that the compound cue is more effective as a discriminative stimulus than the individual cue components alone, they indicate that both WT and KO mice are capable of using either the visual or auditory elements of the CS as discriminative stimuli, and that the deficit observed in the KOs during the conditioned approach task was unlikely to be caused by general sensory deficits in these mice.

**Conditioned reinforcement**

The analysis of lever pressing for the presentation of the CS indicated that the CS attained conditioned reinforcing properties in both WT and KO mice (main effect of lever, \(F_{(1,12)} = 43.19, p < 0.01\) (Fig. 3A)). ANOVA also indicated a three-way interaction between genotype, drug treatment, and lever (\(F_{(1,12)} = 4.86, p < 0.05\)). Additional investigation of this interaction revealed that WT mice responded at higher levels on the CR lever after amphetamine administration than after saline administration, without concurrent changes in responding on the NCR lever (drug treatment by lever interaction; \(F_{(1,7)} = 9.98, p < 0.05\)). However, in KO mice, there was no effect of drug treatment on response.
levels on either lever (main effect of drug treatment, \( F_{(1,8)} = 0.22 \); drug treatment by lever interaction, \( F_{(1,8)} = 0.69 \), both not significant). Independent-sample \( t \) tests revealed that after saline administration, the response rate on the CR lever was significantly higher in KO mice than in WT mice (\( t_{(12)} = 2.80, p < 0.05 \)) whereas response rates on the NCR lever did not differ (\( t_{(12)} = 1.90, p < 0.05 \)). However, after amphetamine administration, response rates on either the CR or NCR lever did not differ between genotypes (\( t_{(12)} = 1.09, p < 0.05 \), respectively, not significant). Importantly, KO mice displayed equal rates of magazine approaches during the CS as a percentage of total approaches (CS%) although this may reflect a performance deficit (Fig. 4B, CR lever in the WT sal condition). There were no differences between genotypes (main effect of genotype, \( F_{(1,8)} = 0.19, p < 0.05 \); genotype-by-schedule interaction, \( F_{(1,8)} = 1.51 \); order by genotype interaction, \( F_{(1,8)} = 0.67, p < 0.05 \); both not significant).

**Instrumental responding**

Response rates for primary reinforcement are shown in Figure 3B. Analysis of the number of reinforcers obtained (top) revealed that as the VI schedule increased, the number of reinforcers earned decreased (main effect of schedule, \( F_{(5,70)} = 219.08, p < 0.01 \)). However, there were no differences between genotypes on this measure. In contrast, analysis of active lever responses (bottom) indicated not only an effect of increasing the VI schedule, but also effects of genotype and a genotype-by-schedule interaction (main effect of genotype, \( F_{(1,14)} = 8.67, p < 0.05 \); genotype-by-schedule interaction, \( F_{(5,70)} = 2.79, p < 0.05 \)). Post hoc analysis revealed that KO mice responded at significantly higher rates on the active lever than WT mice at VI schedules at or above 120 sec. There were no differences in inactive lever responses. Therefore, although KO mice obtained the same number of reinforcers as WT mice, they were emitting a greater number of responses to obtain these reinforcers.

**PIT**

Analysis of PIT was performed by comparing rates of responding during the CS with rates of responding during the inter-CS intervals (VI periods). These data are expressed as a ratio for clarity. Analysis of PIT was performed by comparing rates of responding on a lever resulting in the cues presentation (CR) and on a control lever with no consequences (NCR) during a 3 hr session. Mice received either saline (s) or 0.5 mg/kg amphetamine (amph) before the test session. \( p < 0.05 \) compared with responding on the CR lever in the WT sal condition. Figure 3B. Responding for primary reinforcement in WT and gria2 KO mice. A. Mean square-root responses on a lever resulting in the cues presentation (CR) and on a control lever with no consequences (NCR) during a 3 hr session. Mice received either saline (s) or 0.5 mg/kg amphetamine (amph) before the test session. *p < 0.05 compared with responding on the CR lever in the WT sal condition. B. Responding for primary reinforcement (20 mg food pellet) on VI schedules. Data show mean responses and total reinforcers earned during a 2 hr session. “p < 0.05; **p < 0.01, compared with WT active responses on the indicated schedule.

**Discussion**

In the present experiments, we demonstrate that targeted deletion of the gria2 gene encoding the GluR2 subunit of the AMPA receptor leads to specific deficits in stimulus–reward learning. Namely, deletion of gria2 results in impairments in conditioned approach and PIT, without affecting discriminated approach performance during pavlovian conditioning. In addition, gria2 KO mice display enhanced responding for conditioned reinforcement, but insensitivity to the rate-enhancing effects of amphetamine in this task. Finally, GluR1 expression in amygdala subregions is disrupted after the deletion of gria2.

Gria2 KO mice display normal acquisition of a cue–reward association, as shown by their ability to use the cues as a signal for reward availability (pavlovian conditioning). However, it is not clear which aspects of the cue–reward association are necessary for appropriate responding of this type. After extensive training, magazine approach behavior in gria2 KO mice became less accurate than that seen in WT mice, with regard to the number of magazine approaches during the CS as a percentage of total approaches (CS%). Although this may reflect a performance deficit in the KO mice, it appears more likely that it is a reflection of the motor impairments seen in these mice. First, no deficits were observed on retrieval latency after CS onset, indicating that KO mice were equally capable of using the CS as a signal to retrieve the reward. Second, gria2 KO mice display marked motor coordination deficits, resulting in increased passivity and failure to perform the rotarod task (Jia et al., 1996; Gerlai et al., 1998). Analysis of total magazine entries indicated that the difference in CS% was attributable to a small increase in the number of magazine approaches during the inter-CS periods, rather than decreased approaches during the CS. Although WT mice typically move around the operant chambers during sessions, KO mice were observed to spend most of their time close to the food mag-
in the attribution of motivational value to appetitive cues have been facilitated after the deletion of \textit{gria2}. This is consistent with data showing that AMPA receptors lacking the GluR2 subunit display enhanced calcium permeability (Hollmann et al., 1991; Mishina et al., 1991), and removal of this subunit results in facilitated LTP (Gerlai et al., 1998), perhaps implying enhanced learning ability. If this were the case, it would also be expected that \textit{gria2} KO mice would display enhanced rates of learning on similar tasks, because deletion of GluR2 occurs throughout all brain regions. However, rates of acquisition on pavlovian conditioning did not differ after the deletion of \textit{gria2}, suggesting that these mice do not simply display enhanced learning of all stimulus-reward associations. It is also possible that the increased responding for conditioned reinforcement may be a result of increased motivational value being attributed to the US, because \textit{gria2} KO mice also responded at higher rates for the primary reinforcer (food) when responding was maintained on a VI schedule. Response rates during VI schedules of reinforcement are suggested to be indicative of, and influenced by, the frequency and magnitude of reinforcement, whereby increased rates of responding are associated with increased frequency or magnitude of reinforcement (Herrnstein, 1970; Heyman et al., 1987). However, this explanation for the enhanced responding for conditioned reinforcement relies on the assumption that the motivational value of the US is quantitatively related to the motivational value of an associated CS. Although we are not aware of any direct tests of this assumption, indirect support is provided by the observation that response rates during the first predrug period of a second-order schedule for cocaine are directly related to the cocaine dose associated with the CS (Arroyo et al., 1998).

Despite displaying normal pavlovian conditioning and responding for CR, \textit{gria2} KO mice were clearly impaired on the other aspects of cue-maintained behaviors studied here. Whereas WT mice displayed selective approach toward the stimulus light when it was presented (conditioned approach), \textit{gria2} KOs showed no such approach. This effect could not be attributed to a visual deficit in the \textit{gria2} KOs because these mice were capable of using the light CS as a discriminative stimulus for predicting food delivery as effectively as WT mice. In addition, the ability of the cues to enhance responding on an ongoing operant task (PIT) was abolished after the deletion of GluR2. Although the magnitude of the PIT effect was not as large as we have reported previously in the WT mice in our \textit{gria1} KO experiment (possibly because the background strain of the \textit{gria2} KO mouse was CD1 but whereas that for the \textit{gria1} KO was C57Bl), there was still a clear increase in response rates after the CS onset (Fig. 4 B). One likely explanation for the weaker effect is that WT mice also displayed enhanced magazine approach during the CS, the appropriate response from earlier pavlovian conditioning sessions. This magazine approach resulted in an initial decrease in responding (response competition), followed by an increase (PIT). However, no increase in response rates during the CS was observed in \textit{gria2} KO mice, and because no increase in magazine approach was observed, this lack of PIT cannot be attributed to response competition.

Deficits observed in the \textit{gria1} KO mouse were attributed to impaired BLA function (Mead and Stephens, 2003) because of the similarities in behavioral impairments seen after \textit{gria1} deletion and BLA lesions, a striking similarity is seen between deletion of \textit{gria2} and lesions of the CeA (Table 1). CeA lesions impair conditioned approach and PIT, while leaving intact pavlovian conditioning and responding for conditioned reinforcement (Everitt et al., 2000). Furthermore, CeA lesions abolish the
amphetamine-induced potentiation of responding for conditioned reinforcement (Robledo et al., 1996), as does the deletion of gria2. Therefore, our data indicate a double dissociation in the roles of GluR1 and GluR2 in the behavioral responses to appetitive cues, similar to that reported previously with lesions of the BLA and CeA.

In the case of the gria1 KOs, the deficits could not be conclusively attributed to GluR1 deletion, because we also observed increased GluR2 expression in the BLA of these mice. Previous reports noted no overall changes in levels of GluR1, GluR3, or GluR4 in the brains of gria2 KO mice (Jia et al., 1996), but no studies on possible compensation within specific brain regions have been performed previously. To examine whether compensatory changes in AMPA-receptor subunits occurred after gria2 deletion, we examined GluR1 expression within the amygdala of gria2 KO mice. Although changes in the overall levels of GluR1 were observed in the CeA of gria2 KO mice, no alterations in the number of neurons expressing GluR1 were observed. However, in the BLA, both a reduction in the number of GluR1-positive neurons, and the overall density of GluR1 was found. These observations raise the possibility that the “CeA like” behavioral deficits seen in the gria2 KO mice may have been caused by alterations in GluR1 as well as the deletion of GluR2. However, the fact that a similar downregulation of GluR1 was observed in the BLA suggests that this explanation is unlikely, because BLA-dependent tasks were unaffected. However, it should be noted that the overall decrease in levels of GluR1 in the CeA was significantly greater than that seen in the BLA; therefore, we cannot rule out the possibility that the behavioral deficits observed were caused by altered GluR1 expression.

Although our findings with the gria2 KO mice parallel behavioral deficits seen in rats with CeA lesions, it cannot be concluded from the present data that the deficits we see are attributable to impaired transmission in the CeA. The CeA receives inputs from a large number of brain regions, including cortical and thalamic sensory areas, and other amygdala nuclei. It is likely that neurotransmission in these pathways is also affected by the gria2 deletion. Moreover, there are substantial connections from the CeA to the dopaminergic neurons of the ventral tegmental area (VTA) (Fudge and Haber, 2000) and it has been suggested that it is this projection that plays an important role in conditioned approach (Everitt et al., 2000). Because up to 84% of VTA dopamine neurons carry GluR2 subunits (Chen et al., 2001), it is possible that our results reflect disruption of this pathway rather than disruption of transmission within the CeA itself.

In summary, gria2 KO mice display deficits in both conditioned approach and PIT. Although responding for CR is enhanced in gria2 KO mice, the rate-enhancing effects of amphetamine on responding for CR are absent. The results reported here have interesting implications for understanding the mechanisms

![Figure 5. Immunohistochemical analysis of GluR1 in WT and gria2 KO mice.](image)

**Table 1. Comparison of the behavioral consequences of gria1 or gria2 deletion with lesions of the basolateral or central regions of the amygdala on stimulus–reward learning tasks and control measures**

<table>
<thead>
<tr>
<th>Behavior</th>
<th>gria1 KOa</th>
<th>BLA lesionsb</th>
<th>gria2 KO</th>
<th>CeA lesionsb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pavlovian conditioning</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>Conditioned approach</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>Pavlovian-to-instrumental transfer</td>
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<td>Normal</td>
<td>Impaired</td>
<td>Enhanced</td>
</tr>
<tr>
<td>CR</td>
<td>Normal</td>
<td>Normal</td>
<td>Impaired</td>
<td>Impaired</td>
</tr>
<tr>
<td>Primary reinforcement</td>
<td>Impaired</td>
<td>Enhanced</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>(instrumental)</td>
<td>Enhanced</td>
<td>Normalc</td>
<td>Normal</td>
<td>Normald</td>
</tr>
<tr>
<td>Amphetamine-potentiation of CR</td>
<td>Not tested</td>
<td>Normalc</td>
<td>Impaired</td>
<td>Impairedd</td>
</tr>
<tr>
<td>Second-order schedules of reinforcement</td>
<td>Impairedd</td>
<td>Not tested</td>
<td>Normald</td>
<td>Normald</td>
</tr>
</tbody>
</table>

The effects of gria1 deletion resemble BLA lesions, whereas gria2 deletion resembles lesion of the CeA. Pavlovian conditioning is defined as the ability to use a cue as a signal for reward availability. Responding for primary reinforcement as assessed using a VI schedule of reinforcement for food reward.

aMead and Stephens, 2003.
bEveritt et al., 2000.
dBurns et al., 1993.
underlying relapse in drug abuse. By dissociating the effects of reward cues on behavior at the level of receptor subunits, in addition to the previously demonstrated anatomical dissociation, the ability to prevent cues from influencing certain aspects of behavior without interfering with all stimulus-controlled behaviors becomes more realistic. The ability of reward-paired cues to elicit enhanced pursuit of the associated reward (PIT) has been cited by some as a critical element of the relapse process (Wyvell and Berridge, 2001), and as such, GluR2-containing AMPA receptors may be a suitable target for antagonism when developing treatments for relapse prevention.

References