In vivo knockdown of GAD67 in the amygdala disrupts fear extinction and the anxiolytic-like effect of diazepam in mice

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In mammals, γ-aminobutyric acid (GABA) transmission in the amygdala is particularly important for controlling levels of fear and anxiety. Most GABA synthesis in the brain is catalyzed in inhibitory neurons from L-glutamic acid by the enzyme glutamic acid decarboxylase 67 (GAD67). In the current study, we sought to examine the acquisition and extinction of conditioned fear in mice with knocked down expression of the GABA synthesizing enzyme GAD67 in the amygdala using a lentiviral-based (LV) RNA interference strategy to locally induce loss-of-function. In vitro experiments revealed that our LV-siRNA-GAD67 construct diminished the expression of GAD67 as determined with western blot and fluorescent immunocytochemical analyses. In vivo experiments, in which male C57BL/6J mice received bilateral amygdala microinjections, revealed that LV-siRNA-GAD67 injections produce significant inhibition of endogenous GAD67 when compared with control injections. In contrast, no significant changes in GAD65 expression were detected in the amygdala, validating the specificity of LV knockdown. Behavioral experiments showed that LV knockdown of GAD67 results in a deficit in the extinction, but not the acquisition or retention, of fear as measured by conditioned freezing. GAD67 knockdown did not affect baseline locomotion or basal measures of anxiety as measured in open field apparatus. However, diminished GAD67 in the amygdala blunted the anxiolytic-like effect of diazepam (1.5 mg kg⁻¹) as measured in the elevated plus maze. Together, these studies suggest that GABAergic transmission in amygdala mediates the inhibition of conditioned fear and the anxiolytic-like effect of diazepam in adult mice.

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Introduction

Clinical evidence suggests that alterations in normal γ-aminobutyric acid (GABA) transmission might contribute to the pathophysiology of anxiety disorders in humans. For instance, various studies using nuclear imaging techniques have revealed diminished central GABA and GABA receptor levels in patients suffering from anxiety disorders, including panic disorder, generalized anxiety disorder and posttraumatic stress disorder.1–8 Blunted behavioral and central effects to benzodiazepine (BZ) administration have also been reported in patients with panic disorder.9–11 Previous work has also revealed that individuals with low GABA plasma levels at the time of trauma are reportedly more prone to develop acute posttraumatic stress disorder than other victims.12,13 The link between altered GABAergic transmission and anxiety disorders is also bolstered by the fact that patients suffering from anxiety disorders are commonly treated with BZs, which mediate their actions by augmenting the inhibitory actions of GABA.

Altered GABAergic transmission in the amygdala may be a major factor contributing to the behavioral symptoms associated with anxiety disorders. In mammals, the amygdala is rich in A-type GABA receptors (GABAARs), which are particularly important for controlling fear and anxiety.14–19 In rodents, the anxiolytic-like effects of BZs injected into the amygdala have been demonstrated on a number of animal models of anxiety, including the elevated plus maze (EPM),20–22 social interaction tests,23 conflict tests,24–26 defensive freezing27 and conditioned avoidance.28 On the other hand, microinjections of BZ antagonists and inverse agonists into the amygdala induce anxiogenic-like behaviors26,29 and prevent the anxiolytic-like effects BZs.24,30

Behavioral and physiological lines of evidence suggest that GABAergic transmission in the amygdala undergoes plastic changes that contribute to the control of fear and anxiety. For example, tone fear-conditioning results in a significant reduction of extracellular GABA levels in the amygdala during subsequent re-exposure to a tone conditioned stimulus (CS).31 Likewise, the expression of the GABA synthesizing enzyme GAD65 is reduced in the amygdala 24 h after fear-conditioning training.32 Studies investigating in vitro long-term potentiation in amygdala indicate that both excitatory synapses onto interneurons and inhibitory synapses onto pyramidal cells can undergo long-term potentiation.33,34 Both forms of GABAergic long-term potentiation induce lasting potentiation of inhibitory postsynaptic potentials recorded from amygdala pyramidal neurons and may contribute to the modification of amygdala inhibitory networks. Together, these
studies suggest that inhibitory transmission in the amygdala undergoes experience-dependent changes and that alterations in plastic changes may participate in the development and/or symptoms associated with fear and anxiety.

Although the mechanisms underlying changes in GABAergic neurotransmission are poorly understood, activity-dependent changes in the 67-kDa isofrom of glutamic acid decarboxylase (GAD67) may play a particularly important role in the alteration of inhibitory synaptic transmission. GAD67, along with a second isozyme GAD65, regulate GABA synthesis in the brain and readily influence cellular and vesicular GABA content. The evidence from knockout mice indicates that GAD67 provides the majority of basal GABA for inhibitory neurotransmission. Deletion of the GAD67 gene in mice results in >90% reduction in basal GABA levels in the brain; whereas, GAD65 deleted homozygous mice express normal levels of GABA. Levels of GAD67 are also known to change in response to functional demands of GABA. For example, activity-dependent increases in GAD67 lead to enhanced GABA synthesis, whereas activity deprivation decreases GABA levels, likely through down-regulation of GAD67. Altogether, these data indicate that changes in basal and activity-dependent levels of GAD67 can readily influence GABAergic neurotransmission and that alterations in GAD67 levels may participate in the development and/or symptoms associated with fear and anxiety.

In the present study, we investigated whether altered amygdala GAD67 modulates levels of fear/anxiety and the behavioral effects of the BZ diazepam in adult mice. Specifically, we performed local intracerebral infusions of a small interfering RNA lentivirus (LV-siRNA), engineered to specifically knockdown GAD67 in the amygdala of adult mice, before behavioral testing. Since GAD67 is the rate-limiting enzyme in controlling GABA synthesis, we hypothesized that the LV-induced reduction of GAD67 would decrease GABA-mediated signaling and produce an anxiogenic-like profile in mice as reflected by increased unconditioned and conditioned fear. We also predicted that amygdala GAD67 reduction would blunt the anxiolytic-like effect to the administration of the BZ diazepam.

Materials and methods

Animals. Male C57BL/6J mice (5 weeks of age) were purchased from Jackson Laboratory and allowed to acclimate in the vivarium for 2–3 weeks before surgery. Mice were housed in standard group cages (<6 per cage) and were given ad libitum access to food and water on a 12-h light/dark cycle (light on from 0700 hours). All experiments were performed during the light cycle. This research followed the principles and standards of animal care outlined by the NIH Institutional Animal Care and Use Committee and approved by the UTHSC and Emory University Institutional Review Boards.

Test drug and dose. Diazepam, a nonselective full BZ receptor agonist (Sigma, St Louis, MO, USA) was dissolved in a 15% DMSO/saline solution and administered in a volume of 8 ml kg\(^{-1}\) at a dose of 1.5 mg kg\(^{-1}\). Dose selection was based on previous study, demonstrating that this level produces a robust anxiolytic-like behavioral effect when measure using the EPM. Animal weights were recorded before injections.

Lentivirus production. Viral vectors were derived from the HIV-based lentivirus backbone pLV-CMV–GFP–U3Nhe, which allows for virally mediated expression of green fluorescent protein (GFP) driven by a cytomegalovirus (CMV) promoter. Each small interfering RNA-expressing viral vector (LV-siRNA) was created by subcloning a H1-siRNA coding cassette into the NheI restriction enzyme site of the pLV-CMV–GFP–U3Nhe backbone. The H1-siRNA cassette was constructed by first obtaining the 60–64 bp sense and antisense oligonucleotides against GAD67 (siGAD67) (Sigma) containing the target sequence separated from its reverse complement by a 7-bp loop sequence. Scrambled-sequence oligonucleotides (SCR) were used to create control construct. The sense and antisense oligonucleotides also contained overhangs of BglII at the 5’ end and HindIII at the 3’ end downstream of a H1 pol III promoter. After annealing and phosphorylating the antisense oligonucleotides, the duplex was ligated between BglII and HindIII of the pSuper plasmid. The Hi-siRNA cassette from pSuper plasmid was PCR amplified using T3 and SK-Xba primers (T3-5’-CTCGAAAT-TAACCTCCTACTAAAGGG-3’; SK-Xba-5’-CGAAGGTCGACG GTCTAGATAAGC-3’). The resulting PCR product was digested with XbaI and subcloned into the pLV-CMV–GFP–U3Nhe backbone to create LV-siGAD67 or LV-SCR control lentiviral constructs. Active viral particles were produced by co-transfecting these lentiviral packaging constructs with plasmids coding for delta8.9 and VSV-G into HEK-293 T cells following standard methods. The packaged, unconcentrated virus was collected over a period of 5 days post-transfection, and then concentrated using ultracentrifugation and resuspended in sterile PBS/1% bovine serum albumin (BSA). The resulting titer was assessed in HEK-293 T cells and ranged from 1 × 10^6 to 1 × 10^8 infectious particles per ml.

Motor activity apparatus. Baseline motor activity and the sedative effects of diazepam were measured by examining the total ambulatory distance (in cm) during the 30-min open field test session. Activity for each mouse was measured using individual activity chambers (27.9 cm × 27.9 cm) constructed from clear polycarbonate and equipped with four 16-beam infrared arrays across the base of each chamber wall (MED Associates, St Albans, VT, USA, Model, OFA-MS). The percentage of time mice spent in the central zone of the open field was used as index of anxiety-like behavior. The central zone was defined as the central compartment of the floor located 6 cm from the perimeter of the chamber walls. Activity data were collected via computer and analyzed with the MED Associates’ Activity Monitor Data Analysis software.

EPM apparatus. The EPM consists of two open arms (50 × 6.5 cm²) and two closed arms with a wall (6.5 × 15 cm²) attached to a common central platform (6.5 × 6.5 cm²) to form a cross. The maze was elevated 65 cm above the floor. Test sessions were conducted under adjusted room lighting (100 lux) where behaviors were continuously videotaped by a video camera placed over the
apparatus. The total number of closed arm entries was used as an indicator of locomotor activity. Arm entries and latencies were analyzed and recorded via computer using automated Limelight software (Coulbourn, PA, USA).

**Fear-conditioning apparatus.** Mice were given fear conditioning in standard rodent modular test chambers (ENV-008-VP; MED Associates, St Albans, VT, USA) equipped with high-frequency speakers mounted inside each chamber. The tone CS was a 30-s, 70-dB SPL, 12-kHz tone delivered by GW Instek function generator (GF6-1003, Instek, Chino, CA, USA). The US (unconditioned stimulus) was scrambled foot shock delivered by current generator (Model ENV-414, MED Associates, St Albans, VT, USA) to a removable grid floor that consisted of 36 stainless steel rods (3.2 mm) placed 7.9 mm apart. The footshock US intensity was 0.4 mA verified by using a 0.5-kOhm resistor across the bars of the shock grids and measuring the voltage drop between the bars to calculate the constant current. Conditioned freezing responses were recorded with video cameras mounted on top of each conditioning apparatus. Footshock stimuli and freezing responses were controlled, collected and analyzed via computer using automated FreezeFrame2 software (Coulbourn, Whitehall, PA, USA).

**Cell culture experiments.** Primary cultures of hippocampal neurons from C57BL/6J mice (P10) were prepared using a modified protocol based on Brewer. Briefly, the hippocampi were dissected on ice and dissociated in Hibernate A medium (BrainBits, Springfield, IL, USA). Neurons were plated onto poly-lysine-pre-coated 24-well plates at a density of 2.5 x 10^5 cells cm^-2 in Neurobasal A media (Invitrogen, Carlsbad, CA, USA). The cultures were kept in a humidified incubator at 37°C and 5% CO2. Two weeks after plating, each well received either 1 μl of LV-siGAD67, LV-SCR, or was left untreated followed by 3 days of incubation at 37°C and 5% CO2. Neurons were then fixed with 4% paraformaldehyde in PBS and blocked from nonspecific binding in PBS containing 1% BSA and 3% normal goat serum. GAD67 protein was stained with rabbit anti-GAD67 antibody (MAB5406; 1:500; Millipore, Billerica, MA, USA) and goat anti-rabbit Alexa Fluor 568 (1:1000; Invitrogen). After PBS rinse, cells were treated with Hoechst 33342 stain (10 μg ml^-1) for 10 min to identify nuclei of both neuronal and glial cells. Wells were viewed using a Leica DMRA microscope (Nussloch, Germany) and random images from LV-SCR, LV-siGAD67, and untreated wells (n = 8, area = 0.175 cm^2 per each) were captured using a Nikon Digital Sight DS-U1 camera system and NIS Elements BR2.30 Software. Hoechst, GAD67 and GFP-positive neurons from the same area were counted, and the level of infection was calculated as a percentage of the total Hoechst-stained nuclei.

**Surgery and testing.** Mice received bilateral microinjections of either the LV-siGAD67 or the control LV-SCR into the lateral and basolateral amygdala (BLA). After anesthetizing by injections of Ketamine-Metomidine (80 mg kg^-1, 1.0 mg kg^-1, i.p.), mice were mounted in a stereolocic apparatus, and small holes were drilled in the skull above the injection site. A 30-gauge Hamilton microsyringe was lowered to the following coordinates from bregma based on the mouse brain atlas of Paxinos and Franklin: AP = −1.4, ML = ±3.3, DV = −5.0. The microsyringe was left in place 10 min before and after each injection, and a total volume of 1.0 μl LV-siGAD67 or LV-SCR was administered at a rate of 0.05 μl per min at each site. The incision was closed with Cyanocrylate glue after surgery. Each mouse was placed on a heated pad after a postsurgical i.p. injection of Antisedan (4.0 mg kg^-1) and the narcotic analgesic buprenorphine (0.05 mg kg^-1, s.c.) was administered to mice after recovery from anesthesia. Postsurgical monitoring was performed before testing. Forty-four mice received LV injections (LV-siGAD67 = 22, LV-SCR = 22), three mice died during surgery or recovery period.

Prior testing can alter performance on subsequent tests. To reduce this impact, the test order of behavioral tests was arranged so that mice were evaluated on a tests viewed to be least invasive before more invasive tests. Two weeks after surgery, the effect of GAD67 knockdown on motor activity was analyzed. Three days later, LV-siGAD67 and LV-SCR mice (n = 20, each group) were tested on the EPM. Half of each surgery group (n = 10) received 1.5 mg kg^-1 per i.p. of diazepam 30 min before testing. The second half of each group (n = 10) received vehicle. Studies of concentration-time profiles in adult mice indicate the half-lives of diazepam and metabolites are ~8 h after a single dose. Thus, to limit the effects of prior diazepam treatment on subsequent testing, fear conditioning was conducted following a 3-day drug washout period after EPM testing. The testing order for animals was counterbalanced for treatment condition, and experimenters were blind to the treatment condition of mice.

**Motor and EPM testing.** Baseline motor activity was measured by examining the total ambulatory distance (in cm) during the 30-min open field test session. For EPM testing, each animal was weighed and injected with diazepam (1.5 mg kg^-1) or vehicle (15% DMSO/saline). At the start of each session, a mouse was placed in the central ‘hub’ of the maze with its head facing an open arm and was allowed to freely explore for 5 min. The percentage of open arm entries ([open arm/open + closed arm] entries) x 100) and percentage time in open arms ([time in open arms/time in open + closed arms]) x 100) were computed. Both of these parameters are indicators of an anxiolytic-like effect. The total number of closed arm entries was used as an indicator of locomotor activity. Arm entry was considered complete if all four paws entered a closed or open arm from the central platform.

**Conditioned fear training, retention and extinction.** During training, mice were placed in chambers, and after 5 min, received the first of 5 tone + shock trials at a 2-min variable intertrial interval (range, 1–3 min). One day after training, animals were given a retention test session containing 15 CS-alone test trials (intertrial interval, 2 min). The extinction training consisted of six test sessions. On each of six consecutive days, mice were given a test session consisting of 15 CS-alone test trials (tone-no shock). CS-alone test trials were presented at an intertrial interval of 2 min.

**Behavioral data analyses.** Levene’s and Bartlett’s tests conducted on data sets showed no strong departures from
normality and homogeneity, respectively. In no cases were both the normality and homogeneity of variance assumptions violated, thus parametric tests were used to analyze data sets.

EPM-dependent measures, including percent of open arm entries and percent of open arms time, were analyzed separately by means of a 2 × 2 ANOVA (analysis of variance) with surgery group (LV-siGAD67, LV-SCR and drug level (Nondrug, Drug) as the between-subjects factors. The 30-min open field test session was evaluated with surgery group as a between-subjects factor and blocks of 10-min intervals (blocks 1–3) as a within-subjects factor. The acquisition of conditioned freezing was analyzed by means of a repeated-measure ANOVA with surgery group as a between-subjects factor and training trial (trials 1–5) as a within-subjects factor. The retention of fear was analyzed by comparing group mean freezing responses (session 1) using a t-test. The acquisition of extinction was analyzed by means of a repeated-measure ANOVA with surgery group as a between-subjects factor and test session (sessions 1–6) as a within-subjects factor.

**Histological analyses.** The effectiveness and consequences of LV-siGAD67-induced gene knockdown in the amygdala were assessed by quantifying GAD67 protein levels and mRNA levels of GAD67 and GAD65. Protein levels were assessed by via western immunoblotting and mRNA levels via *in situ* hybridization.

**Western immunoblots.** The brains were removed and blocked rapidly over ice into 2-mm-thick coronal sections. Amygdala punches were obtained with a 1-mm brain punch tool, and punches from each side were homogenized on ice with a sonicator in 25 μl homogenized lysis buffer (5 mM HEPES, 1 μM EDTA, protease inhibitors) with a protease inhibitor cocktail kit (Roche, Indianapolis, IN, USA). These samples were kept frozen at −80°C until western blot assay. Protein concentrations were measured using 2.5 μl samples of homogenates with BCA protein assay reagent and BSA as the reference standard (Pierce, Rockford, IL, USA). After boiling for 5 min, equal amounts of protein (20 μg) were subjected to polyacrylamide gel electrophoresis using 7.5% polyacrylamide gels (Bio-Rad Laboratories, Hercules, CA, USA) to separate proteins. After electrophoresis, samples were transferred to nitrocellulose membrane (0.45 μm, Bio-Rad Laboratories) at 30 mA for 2 h at 4°C (25 mM Tris–HCl, 190 mM glycine, 20% methanol). Membranes were washed three times for 5 min with blocking buffer (2% nonfat dry milk, 0.1% Tween 20, 50 mM NaCl, 10 mM HEPES, pH 7.4), then incubated with a primary antibody at 4°C overnight. Following three 5-min washes with blocking buffer, membranes were incubated with a corresponding HRP-coupled secondary antibody (1:4000 dilution) at room temperature for 60 min, subsequently washed with water to remove the substrate solution, and detected with SuperSignal West chemiluminescence substrate (Pierce). Protein bands were visualized and analyzed using an Alpha Innotech Fluorchem imaging system (Alpha Innotech, San Leandro, CA, USA).

Monoclonal mouse anti-glyceraldehyde-3-phosphate dehydrogenase antibody (GAPDH; 1:3000) was obtained from Fitzgerald Industries (Concord, MA, USA). Total blotted protein levels were normalized to levels of GAPDH to control for variations in protein loading. As such, the relative values were quantified as the protein of interest divided by the loading control. Statistical analyses were performed with independent samples t-tests (LSD). For ease of presentation, figures are presented as percentages of control group (LV-SCR).

**In situ hybridization.** Mice were anesthetized with isoflurane and the brains were rapidly dissected and frozen on crushed dry ice. Coronal sections (20 μm) of the brains were cut on a cryostat (Leica) at −20°C, mounted on gelatin-coated slides, and stored at −80°C until processed for histochemistry. For each brain, sections were placed on consecutive sets of slides such that each set contained similar sequential sections of the brain. Sections were collected from areas before the amygdala to anterior areas of the prefrontal cortex and sections were stained with Nissl stain.

In *in situ* hybridization and analyses were performed to examine the expression of GAD67 mRNA as previously described and validated. Briefly, hybridization riboprobes were prepared from linearized clones using appropriate RNA polymerase and radioactive 35 S-UTP in the polymerase reaction. Radiolabeled antisense RNA strands were base hydrolyzed and isolated using a sephadex spin columns (Roche). Probes were diluted to a concentration of 100 000 c.p.m. per μl in buffer, and sections were incubated overnight in humid chambers at 52°C with 75 μl of probe-buffer solution covered with a Parafilm coverslip. Slides were then washed, air dried and placed against Biomax MR autoradiography film (Eastman Kodak, New York, NY, USA) for 2 days. Next, film was scanned (3000 dpi) and saved in JPEG format. To estimate the levels of mRNA, the total area of each target site subregion was outlined with reference to Nissl-stained sections and the atlas of Paxinos and Franklin. Nissl-stained sections were also used for the identification of injection sites.

The hybridization signal intensities of the brain regions were calculated on the basis of gray values between 0 and 255. Hybridization signals were determined for the regions of interest, as well as an adjacent background area with little or no hybridization signal.

**Results**

**In vitro efficiency of GAD67 protein knockdown in primary cell cultures.** The lentivirus designed to knockdown expression of GAD67 was initially tested *in vitro* by examining its efficiency in knocking down the expression of GAD67 primary cultures of hippocampal neurons from C57BL/6J mice (P10). For this experiment, 24-well plate cell cultures were treated with either 1 μl of LV-siGAD67, 1 μl of LV-SCR or left untreated. Three days later, plates were processed for either immunocytochemistry (ICC) or western blotting. In general, ICC images revealed pronounced GFP expression in lentiviral-infected wells. As seen in Figure 1a,
wells infected with LV-SCR and stained with anti-GAD67 (red) showed prevalent expression of GAD67. Antibody-stained wells of LV-siGAD67 revealed considerably less GAD67 staining (Figure 1b). A total of 764 cells were examined for GAD67 and/or GFP fluorescent expression (siGAD67 = 512, SCR = 252), of which ≈32% expressed GFP. When compared with LV-SCR wells, the percent of GFP-positive cells that also expressed GAD67 was significantly less in LV-siGAD67-infected cells (Figure 1c). Western blot processing of cells harvested from plate cultures revealed that LV-siGAD67-infected cells qualitatively expressed significantly less GAD67 protein than control-infected cells (Figures 1d and e).

Efficiency of GAD67 mRNA knockdown within the BLA. After the conclusion of the behavioral testing, the levels of protein or mRNA expression were examined in LV-infected mice to assess whether viral infections successfully induced GAD67 knockdown in vivo. In situ hybridization analyses for GAD65 and GAD67 mRNA were conducted on coronal brains sections obtained from half of the experimental animals (n = 10 per group). Figure 2a illustrates the qualitative decrease of GAD67 mRNA expression within the amygdala following LV infection. Nissl-stained sections revealed that injection sites were centered in the BLA with an anterior–posterior range of ~−1.2 to −2.0 mm with reference to bregma. Stained sections also exhibited intact cellular and regional morphology at the site of injection, with no clear evidence of cell death or degradation. Quantitative evaluation of GAD67 mRNA levels in LV-siGAD67 and LV-SCR-infected mice indicated that LV-siGAD67 mice expressed significantly less mRNA in the BLA when compared with LV-SCR controls (Figure 2b). LV-siGAD67 mice also showed less mRNA expression in the central nucleus of the amygdala (CE), but the group difference was not reliably significant.

Quantitative evaluation of GAD67 mRNA levels in LV-siGAD67 and LV-SCR-infected mice indicated that LV-siGAD67 mice expressed significantly less mRNA in the BLA when compared with LV-SCR controls (Figure 2b). LV-siGAD67 mice also showed less mRNA expression in the central nucleus of the amygdala (CE), but the group difference was not reliably significant. With reference to LV-SCR control animals, the relative expression levels of GAD67 mRNA within the BLA and CE of LV-siGAD67-infected animals were 65 and 87%, respectively. GAD65 mRNA in the BLA was 126% in LV-siGAD67 mice with reference to controls. To examine protein levels, amygdala brain punches were obtained in remaining half of the experimental animals (n = 10 per group) and homogenized samples were subjected to western immunoblot analysis. As seen in Figure 2c, immunoblot analysis indicated that LV-siGAD67 effectively knocked down targeted protein in vivo, relative to LV-SCR t(18) = 2.67, P < 0.05.

GAD67 knockdown does not affect open field motor activity. Two weeks after stereotaxic infection of lentivirus targeting the bilateral amygdala, mice were subjected to a behavioral battery. Before fear conditioning, LV-infected
mice were tested in the open field maze to determine whether GAD67 knockdown affected baseline motor activity. As seen in figure 3a, both LV-siGAD67 and SCR mice showed similar levels of activity when measured for 30 min. Overall, both groups showed a decrease in activity during the 30-min session, $F(1,39) = 82.58, P < 0.01$; however, neither the main effect of group, $F(1,39) = 1.16, P > 0.05$, or group × treatment interaction, $F(1,39) = 0.32, P > 0.05$, were significant. An independent t-test revealed no group differences in the time mice spent in the central zone of the open field, $t(39) = 0.89, P > 0.05$, suggesting no differences in basal anxiety levels. The mean percentages of time in the central zone (with standard error of the mean) for LV-siGAD67 and LV-SCR groups were 43.5% (4.5) and 38.6% (3.4), respectively.

GAD67 knockdown impairs the anxiolytic-like effects of diazepam in the EPM test. To examine whether GAD67 knockdown affected the anxiolytic-like effects of diazepam, both LV-siGAD67 and LV-SCR animals were divided into two groups that received either diazepam (1.5 mg kg$^{-1}$) or vehicle 30 min before EPM testing ($n = 10$ per group). As seen in Figures 3b and c, comparisons of vehicle- and diazepam-treated animals revealed a number of significant effects. An examination of the percentage of open arm time detected significant effects of group, treatment and group × treatment interaction. Overall, the percentage in open arm time was greater for diazepam-treated animals, $F(1,37) = 11.0, P < 0.05$, revealing the general anxiolytic-like effects of diazepam. In general, the percentage in open arm time was greater in LV-SCR compared with LV-siGAD67 mice, $F(1,37) = 4.7, P < 0.05$. Analyses of the group × treatment interaction, $F(1,37) = 4.3, P < 0.05$, indicated that vehicle-treated LV-siGAD67 and SCR animals showed similar percentages in open arm entries, $t(39) = 0.15, P > 0.05$. In diazepam-treated animals, the percentage in open arm time exhibited by LV-SCR mice was significantly greater than LV-siGAD67 mice, $t(39) = 2.30, P < 0.05$, indicating that diazepam induced a greater anxiolytic effect in SCR compared with LV-siGAD67 mice.

Also consistent with the anxiolytic-like effects of diazepam, the percentage of open arm entries was greater in diazepam-
treated animals versus animals that received vehicle, as denoted by a significant treatment effect. The main effect of group across treatments was marginally nonsignificant, $F(1,37) = 3.28, P = 0.08$, and the group × treatment interaction was statistically nonsignificant, $F(1,37) = 1.39, P > 0.05$.

Assessment of total closed arm entries indicated significant differences in locomotor activity among treatments, $F(3,30) = 8.33, P < 0.01$. Diazepam-treated animals made significantly more closed arm entries than the vehicle group, indicating a drug-induced increase in exploratory/motor-stimulant activity. The mean closed arm entries (with standard error of the mean in parentheses) for vehicle- and diazepam-treated animals were 6.7 (0.87) and 22.6 (0.85), respectively. No differences in the number of closed arm entries were observed between groups at each level of treatment ($P$ values $> 0.05$). Thus, observed group differences in the percentage of open arm time cannot be attributed to differences in locomotor activity.

**GAD67 knockdown does not affect the acquisition or retention of conditioned freezing.** We next examined whether GAD67 knockdown in amygdala would affect acquisition or expression of conditioned fear. Fear training to an auditory cue was performed in a Med Associates conditioning chamber and fear testing was performed in a modified context to reduce contextual fear. During the conditioning phase of the session, freezing scores similarly increased from the first to the last CS presentation in both groups as shown in Figure 4a. The ANOVA showed a significant main effect of trial, $F(4,39) = 32.4, P < 0.01$, but no group effect or group × trial interaction, $F$-values $< 1.0, P$-values $> 0.05$. When tested 24 h after conditioning, both LV-siGAD67 and LV-SCR mice displayed similar levels of tone CS freezing to the unreinforced CS, $t(39) = 0.78, P > 0.05$, signifying that the retention of fear was similar in both groups (Figure 4b).

**GAD67 knockdown impairs extinction of conditioned fear.** Following the first cued fear retention test, mice were repeatedly tested for five additional test sessions with the unreinforced auditory CS to examine the extinction of fear. As illustrated in Figure 4c, LV-siGAD67 mice showed higher levels of fear during repeated testing when compared with controls. Repeated-measure analysis of groups across test sessions exposed significant effects of group and session, $F$-values $< 1.0, P$-values $> 0.05$. Although there were no differences between initial levels of conditioned fear on test 1, there were significant differences between the levels of conditioned freezing on test 2, test 3 and test 4 of extinction sessions, $t$s(39) $> 2.48, P$-values $< 0.02$. No differences in fear levels were observed on extinction tests 5 or 6, $t$s(39) $< 1.30, P$-values $> 0.05$.

**Discussion**

The present study first sought to determine whether the knockdown of GAD67 in the amygdala would modulate the acquisition and extinction of conditioned fear. Both LV-siGAD67 and LV-SCR groups showed similar increases in conditioned fear during training; thus, the reduction of amygdala GAD67 produced no apparent changes in the acquisition of fear as measured by conditioned freezing. Following cued fear acquisition, mice were repeatedly tested over 5 days with the unreinforced auditory cue to examine...
subsequent retention and extinction of fear to the conditioned stimuli. Groups showed no differences in fear on the initial test, suggesting similar levels of fear retention. However, there were significant differences between the levels of conditioned fear on succeeding tests. Overall, mice with targeted amygdala GAD67 knockdown showed higher levels of fear on subsequent extinction sessions. Together, these data suggest that although there does not appear to be a direct effect of GAD67 knockdown on acquisition or retention of conditioned fear, there is an impairment of extinction of conditioned fear measured with freezing.

It is important to note that GAD67 is also expressed in glial cells as well as neurons. Furthermore, VSV-G-pseudotyped lentiviral vectors carrying a CMV promoter can express transgenes in both cell subsets. Thus, it is possible that decreased GAD67 levels in glial cells may have also contributed to our results. However, past studies have revealed that the percentage of glial cells transduced by lentiviruses is relatively low compared with neurons. Future research using lentiviral vectors with neuron-specific promoters that induce GAD67 knockdown in different interneuron classes (for example, somatostatin, cholecystokinin, parvalbumin) may help in determining the distinctive roles of GAD67 in specific interneuron types and glial cells.

**Amygdala GAD67 and fear conditioning.** Although GABAergic mechanisms have been implicated in numerous amygdala-dependent functions including learning, memory and anxiety, studies have not yet definitively demonstrated that the reduction of endogenous GAD activity within the amygdala is causal relative to the behaviors being tested. Knockout mice lacking GAD65 display altered conditioned fear behavior; however, these mutants display decreased GAD65 throughout the brain and developmental alteration in cortical plasticity. Pavlovian conditioning appears to alter both isoforms of GAD as well as GABA. For example, tone fear conditioning results in a significant reduction of extracellular GABA levels in the amygdala during subsequent re-exposure to the tone CS. Likewise, the expression of GAD65 is significantly reduced in the amygdala after fear-conditioning training. The inhibition of CS-elicited conditioned responses associated with extinction are in part thought to be mediated by circuits activating or strengthening GABAergic transmission in the amygdala; thus, the increase in GAD activity may represent a mechanism that enables enhanced inhibitory control of fear responses. The above findings are also particularly relevant in the light of past studies showing pharmacological treatments that increase the excitability of amygdala by decreasing GABA transmission facilitate conditioned fear responses. We have previously reported that levels of GAD67 mRNA in the amygdala also decrease after the acquisition Pavlovian fear; whereas, fear extinction produces a training-induced increase in GAD67 mRNA transcript levels. Unpredictably, the reduction of GAD67 mRNA and protein did not significantly influence the acquisition of conditioned fear in the current study. It is possible that freezing levels in Figure 4a reflect a behavioral ‘ceiling effect’ that thwarted detection of group differences during acquisition. Accordingly, we cannot completely rule out the possibility that levels of GAD67 mRNA and/or protein significantly affect fear acquisition. In such case, group differences in levels of freezing during extinction sessions may be due to different pre-extinction fear levels. Future research using lentiviral vectors with neuron-specific promoters that induce GAD67 knockdown in different interneuron classes (for example, somatostatin, cholecystokinin, parvalbumin) may help in determining the distinctive roles of GAD67 in specific interneuron types and glial cells.

**Figure 4** Conditioned fear and extinction in mice with amygdala GAD67 knockdown. (a) Acquisition of conditioned fear to auditory conditioned stimulus (CS) + shock pairing as measured by freezing. There was no difference in acquisition of fear in animals receiving bilateral amygdala injections of LV-SCR or LV-siGAD67. (b) Animals were tested 24 h after training to examine the retention of conditioned fear (session 1 extinction). There was no difference between the groups on level of conditioned fear. (c) Percent conditioned freezing is graphed for sessions 1–6 of extinction tests. Mice infected with LV-SCR demonstrate significant greater decreases in their level of conditioned fear, compared with LV-siGAD67-infected mice. Impaired extinction of fear, measured by percentage of freezing, is evident across multiple testing sessions. Error bars denote 1 standard error of the mean (s.e.m.). Star indicates that the difference between virus groups was statistically significant ($P<0.05$).
experiments including lower shock US levels, fewer training trials or different indices of fear (for example, fear-potentiated startle) may evaluate this possibility. It must also be noted that although alterations in mRNA concentrations are widely used as a surrogate for changes in protein, in vivo mRNA levels obviously do not necessarily correlate with protein levels. Numerous posttranscriptional mechanisms regulate the rate, timing and amount of the encoded protein (see below).

**Amygdala GAD67 and anxiety.** Our study also examined whether a reduction of amygdala GAD67 would result in an increased anxiety-like behavior and decrease the anxiolytic-like effect of diazepam as measured in the open field and EPM. Because a decrease in GABAergic transmission is associated with an anxiogenic-like profile in rodents, we hypothesized that LV-induced silencing of amygdala GAD67 would result in an anxiogenic-like profile. However, in the absence of diazepam, we found that LV-siGAD67 and LV-SCR groups showed similar EPM and open field performance. These findings contrast with previous data, suggesting that a reduction of central GAD produces an anxiogenic-like phenotype in mice. For example, GAD65-deficient mice exhibit increased anxiety-like behaviors, which are attributed to reduced GABAergic neurotransmission in the amygdala and hippocampus. However, these mutants also demonstrate increases in spontaneous seizures and are sensitive to drug-induced seizures, suggesting that the GAD deficits in other brain regions may also contribute to anxiety-like profile. A decrease in amygdala GAD67 may be compensated for by an increased extrasynaptic GABAAR function, as proposed for deletion of Gad65. On the other hand, mice selectively bred to display high levels of anxiety-related behavior show marked increases in the expression of GAD65 and GAD67 mRNAs and GABA. The increased expression of GADs may be a developmental mechanism to compensate for persistent activation of the amygdala in high-anxiety mice.

**Amygdala GAD67 and behavioral sensitivity to diazepam.** While amygdala down-regulation of GAD67 produced no measureable changes in anxiety-like behavior, it did reduce the behavioral sensitivity for diazepam as assessed in the EPM. These results are consistent with past studies that show GAD65 KO mice display diminished behavioral responses to BZs and barbiturates. Human studies have revealed that patients suffering from anxiety disorders have blunted behavioral responses to BZ administration and have diminished central BZ binding. GAD67 provides the magnitude of basal GABA for inhibitory neurotransmission, thus, a reduction in the occupancy of GABAARs by GABA is a simple explanation for the diminished diazepam potency in LV-siGAD67 mice. Changes in the allosteric interactions between the GABA and BZ binding sites may also account for the reduced behavioral sensitivity to diazepam. In vivo changes in GABA concentrations and receptor modulators are known to induce homologous uncoupling. A variety of transcriptional mechanisms, including alteration in the transcription of GABAAR subunit genes, play a role in neuro-adaptive processes underlying uncoupling. Thus, it is conceivable that GAD67 down-regulation leads to altered subtype expression or posttranslational receptor modifications that reduce agonist effects of diazepam.

**The influence of GAD67 inhibition on GABA release.** A reduction in GAD67 may decrease inhibitory transmission by influencing vesicular as well as nonvesicular release of GABA. The majority of GAD67 in the brain is present in the constitutively active state and is targeted to synapses, axons and dendrites. GAD67 is recognized for fulfilling metabolic demands of neurons and the synthesis of GABA for nonvesicular release through GABA transporters, which can be induced to operate in a reverse direction. The role of GAD67 in the synthesis and transport of GABA into synaptic vesicles is unclear. Past studies have recognized that the structural and functional coupling between GAD65 and vesicular GABA transporter mediates the synthesis and transport of GABA into synaptic vesicles. However, recent evidence suggests that GAD67, like GAD65, forms large clusters in presynaptic termini of inhibitory interneuron and may also synthesize GABA from glutamate and transport this newly made transmitter into synaptic vesicles. Consistent with this, subcellular fractionation of brains from GAD65 KO mice reveals that GAD67 is localized in presynaptic clusters in the absence of GAD65. Thus, the reduction in GAD67 in the current study may decrease vesicular along with nonvesicular release of GABA.

**Amygdala GABAergic transmission.** In the amygdala, synaptic release of GABA plays a crucial role in controlling pyramidal neuron excitability and responsiveness through feedforward, feedback and tonic inhibition. Local inhibitory interneurons in the BLA predominantly terminate on α1-, α2- and α3-containing GABAARs located on pyramidal projection neurons. Pyramidal cells also contain extrasynaptic α4-GABAARs, which are tonically activated by ambient GABA and are important for controlling basal neuronal excitability. Recent evidence indicates that a reduction of α1-GABAARs in the BLA does not affect baseline anxiety-like behavior or the anxiolytic-like activity of diazepam as assessed using the EPM model. Thus, the reduction of GABA-mediated activation of α2- and α3-GABAARs could explain the attenuated behavioral sensitivity to diazepam seen in the current study. Consistent with this hypothesis, experimental data from genetically altered mice and subtype-selective ligands indicate that the α1-GABAAR subtype is involved in sedative and amnesic processes, whereas the α2- and α3-GABAARs are specifically involved in anxiety. It should also be mentioned that the release of GABA in the BLA can suppress glutamatergic afferents via presynaptic GABABRs.

Although GAD67 mRNA levels in the central nucleus of the amygdala (CeA) were not significantly reduced, it is unclear whether LV-siGAD67 injections reduced mRNA levels in intercalated (ITC) amygdala neurons that are located between the BLA and the CeA. ITC cells of the amygdala are densely packed inhibitory neurons that send efferent projections to CeA that contains α2 and α3-GABAAR subtypes. These neurons are innervated by excitatory inputs from the
BLA and infralambic cortex, which have both been implicated in mediating the extinction of conditioned fear. A reduction of GABA in these ITC neurons could explain the extinction deficit seen in the current study. Our analyses have also not examined whether the GAD67 reduction reflects an overall reduction in interneurons or a selective reduction in one or more interneuron subtypes. Important tasks of future studies therefore will be to examine these possible influences on our current findings.

Alterations in GAD67 and the regulation of synaptic plasticity in GABAergic transmission. The mechanisms that regulate the development and structural plasticity of GABAergic neurotransmission remain poorly understood. Many studies indicate that activity-dependent alterations in GAD67 may play a role in the plasticity of inhibitory synaptic connections. The expression of GAD mRNAs is known to be regulated by neuronal activity, so that tonically firing GABAergic neurons express high amounts of GAD and particularly GAD67 mRNA and protein. In hippocampal GABAergic neurons, activity-dependent upregulations of GAD65 and GAD67 mRNA and protein are observed following kainate- or picrotoxin-induced seizures. This seizure-induced plasticity may reflect a compensatory upregulation in the subpopulation of surviving GABA neurons to control the increased excitability of the region; similarly, direct in vitro kindling of the dentate gyrus results in the emergence of fast GABAergic transmission that coincides with the expression of GAD67 and vesicular GABA transporter mRNA. Conversely, withdrawal of cortical circuits from their normal excitatory inputs results in the downregulation of GAD67 mRNA and GABA levels. Evidence suggests that the efficacy of GABA-mediated mIPSCs can be up- and down-regulated in response to altered levels of activity in neuronal networks. Together, these studies indicate that activity-dependent and bidirectional alterations in GAD67 and GABA metabolism provide a mechanism for activity-dependant regulation of inhibitory synaptic plasticity.

In summary, this study demonstrates that the reduction of GAD67 in the amygdala of adult mice results in a deficit in the extinction, but not the acquisition or retention, of conditioned fear. Furthermore, amygdala GAD67 reduction blunts the anxiolytic-like effects of diazepam without affecting basal anxiety levels. Studies performed in humans and animal models have implicated alterations in GABAergic transmission may participate in anxiety disorders. Our findings suggest that changes in basal and/or activity-dependent levels of amygdala GAD67 may be important in mediating some of the symptoms associated with anxiety disorders.

Conflict of interest
The authors declare no conflict of interest.

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In vivo knockdown of GAD67 in the amygdala

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