Medial auditory thalamic stimulation as a conditioned stimulus for eyeblink conditioning in rats

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The neural pathways that convey conditioned stimulus (CS) information to the cerebellum during eyeblink conditioning have not been fully delineated. It is well established that pontine mossy fiber inputs to the cerebellum convey CS-related stimulation for different sensory modalities (e.g., auditory, visual, tactile). Less is known about the sources of sensory input to the pons that are important for eyeblink conditioning. The first experiment of the current study was designed to determine whether electrical stimulation of the medial auditory thalamic nuclei is a sufficient CS for establishing eyeblink conditioning in rats. The second experiment used anterograde and retrograde tract tracing techniques to assess neuroanatomical connections between the medial auditory thalamus and pontine nuclei.

Stimulation of the medial auditory thalamus was a very effective CS for eyeblink conditioning in rats, and the medial auditory thalamus has direct ipsilateral projections to the pontine nuclei. The results suggest that the medial auditory thalamic nuclei and their projections to the pontine nuclei are components of the auditory CS pathway in eyeblink conditioning.

A primary emphasis in neurobiological analyses of Pavlovian eyeblink conditioning has been on identifying the anatomical sites and cellular mechanisms of memory storage (Christian and Thompson 2003; Thompson 2005). Less emphasis has been placed on identifying the neural pathways that conduct stimulation from conditioned stimuli to the sites of memory induction and retention. The cerebellum is the anatomical site of memory storage in Pavlovian eyeblink (eyelid and nictitating membrane movement) conditioning (Christian and Thompson 2003; Thompson 2005; Ohyama et al. 2006). Cerebellar function may be influenced by processes occurring within its sensory input pathways, interactions among components of the input pathways, and sources of feedback to the input pathways (Clark et al. 1997; Bao et al. 2000; Medina et al. 2002). A full characterization of the sensory input pathways to the cerebellum necessary for motor learning is, therefore, critical for developing a comprehensive understanding of cerebellar function. Identification of sensory input pathways to the cerebellum is also critical for elucidating the mechanisms underlying the ontogeny of motor learning (Freeman et al. 2005).

Several key components of the conditioned stimulus (CS) pathway in eyeblink conditioning have been identified using lesion, inactivation, unit recording, stimulation, and tract tracing techniques (Steinmetz et al. 1986, 1987, 1989; Lewis et al. 1987; Knowlton and Thompson 1988; Steinmetz 1990; Steinmetz and Sengelaub 1992; Gould et al. 1993). The monosynaptic cochlear-pontine projection may play an important role in auditory eyeblink conditioning, but a recent study found that auditory structures that receive direct or indirect input from the cochlear nuclei also play an important role in eyeblink conditioning. Halverson and Freeman (2006) found that unilateral lesions of the medial auditory thalamic nuclei (MATN), including the medial nucleus of the medial geniculate, posterior intralaminar nucleus, and suprageniculate contralateral to the conditioned eye severely impaired acquisition of eyeblink conditioning using a tone CS in rats. The findings of Halverson and Freeman (2006) suggested that the contralateral medial auditory thalamus could play an essential role in auditory eyeblink conditioning. A major unresolved issue from this study was how the medial auditory thalamus might influence the eyeblink conditioning circuitry. The most straightforward way for the MATN to play a role in the CS pathway would be through an axonal projection to the pontine mossy fiber system.

The first experiment of the current study was designed to determine whether electrical stimulation of the MATN is sufficient for establishing associative eyeblink conditioning in rats. Rats were given stimulation of the medial auditory thalamus as a CS paired with a peripheral shock unconditioned stimulus (US). A control group was given unpaired presentations of medial auditory thalamic stimulation and the US. Experiment 2 was designed to determine whether the MATN sends axonal projections to the pontine nuclei. Anterograde and retrograde tracing methods were used to assess the projections of the medial auditory thalamus in untrained rats.

Results

Experiment I: Stimulation of the MATN as a CS Is Sufficient for eyeblink conditioning

Experiment 1 examined whether electrical stimulation of the medial auditory thalamus is sufficient for establishing associative eyeblink conditioning in rats. Rats were implanted with a bipolar stimulating electrode in the medial nucleus of the medial genicu-
late. They were then given paired or unpaired training with me-
dial auditory thalamic stimulation (300 msec, 200 Hz train; 100
µsec biphasic pulses; 30–100 µA) as the CS and a periorbital shock
US (10 msec, 1.0–2.0 mA). Electrodes placed in the medial
nucleus of the medial geniculate could possibly affect eyeblink
conditioning by stimulating nearby structures, including the an-
terior pretectal nucleus medially and the ventral nucleus of the
medial geniculate laterally, if the current spread beyond the tar-
get. A group of rats were given intentionally misplaced electrodes
in the anterior pretectal nucleus or ventral nucleus of the medial
geniculate to determine whether stimulation of these structures
produces conditioning that is as rapid as the conditioning ob-
served with electrodes placed in the medial auditory thalamus.
Rapid conditioning in the rats with misplaced electrodes would
cast doubt upon the specificity of the medial auditory thalamic
stimulation.

The MATN receive input from the ipsilateral inferior colliculus
but also send projections back to it (see experiment 2). It
would, therefore, be possible that stimulation of the medial au-
ditory thalamus has its effects on eyeblink conditioning through
activation of the inferior colliculus rather than through activa-
tion of a direct projection to the pontine nuclei. To assess the
possible contribution of the thalamic projection to the inferior
colliculus in eyeblink conditioning with medial auditory tha-
lamic stimulation as the CS, a group of rats was given electrolytic
lesions of the inferior colliculus ipsilateral to the stimulation site
1 wk before the onset of paired eyeblink conditioning.

Eyeblink conditioning
Rats in the control group that received paired training with
MATN stimulation acquired conditioning very rapidly, whereas
control rats that received unpaired training with MATN stimula-
tion did not (Fig. 1). Rats with intentionally missed electrodes
in the anterior pretectal nucleus or the ventral medial geniculate
showed only a small increase in conditioned response (CR) per-
centage during the third training session (Fig. 1). The rats given
lesions of the inferior colliculus showed conditioning that was
comparable to the paired control group (Fig. 1).

The effects described above were confirmed by an ANOVA
that revealed a significant effect of the group factor,
F(3,13) = 67.15, P < 0.001. Post hoc tests showed that the control
and lesion paired groups produced a greater CR percentage than
the groups given unpaired training and paired training with mis-
placed electrodes, P < 0.01. Two of the rats with intentionally
misplaced stimulation electrodes on the medial side of the MATN
(Fig. 2) acquired conditioning, albeit slower than rats with accu-
rately placed electrodes. The two rats with intentionally mis-
placed stimulation electrodes in the ventral medial geniculate
(Fig. 2) did not show an increase in CRs across training. These rats
received an additional session of paired training with the stimu-
lation CS followed by three sessions of paired training with a
tone CS (300 msec, 2.0 kHz, 85 dB). The rats with missed elec-
trodes that did not condition with the stimulation CS were given
training with a tone CS to determine whether damage to the
MGN caused by the electrode impaired conditioning. These rats
acquired conditioning with the tone CS during the three sessions of
training (mean CR percentage across sessions = 15.5%, 38.6%,
and 83.7%, respectively). An additional session of paired training
with stimulation was then given to assess whether any behav-
ioral savings could be observed with stimulation after learning
with a peripheral tone CS. However, these rats showed very few
CRs during this session (mean = 6.6%). The results of the follow-
up training and testing sessions indicated that the electrode
placements in the ventral medial geniculate were not effective for
producing conditioning, suggesting that the thalamo-cortical pro-
jection does not play a significant role in delay conditioning.

Conditioned response amplitude was measured on CS-alone
trials. Only the control and lesion groups given paired training
produced enough CRs during CS-alone trials for meaningful sta-
tistical analyses. An ANOVA of the CR amplitude data found no
significant group differences between the paired control and le-
sion groups. Both groups showed a progressive increase in CR
amplitude across training (mean = 3.43, 4.65, 5.44, respectively).
No significant group differences were found for the CR onset or
peak latency measures, which both decreased across sessions (on-
set mean in msec = 198.47, 128.49, 123.75; peak mean in msec
= 296.10, 255.05, 253.87).

Stimulation electrode placement
The electrodes in the paired and unpaired control groups were
consistently placed in or within 0.2 mm of the medial nucleus of
the medial geniculate (Fig. 2, left). The paired and unpaired con-
trols had equivalent electrode placements. The rats given inferior
colliculus lesions also had electrodes placed within 0.2 mm of the
medial nucleus of the medial geniculate. Intentionally missed
electrodes were placed in the ventral nucleus of the medial ge-
iculate and in the anterior pretectal nucleus (Fig. 2, right).

Figure 1. Medial auditory thalamic stimulation as a conditioned stimu-
lus supports associative eyeblink conditioning. Mean conditioned re-
sponse (CR) percentage for rats with accurately placed electrodes given
paired (black circles) or unpaired training (white circles), rats given in-
tentionally misplaced electrodes (black triangles), and rats given inferior
colliculus lesions (gray squares) across 30 10-trial blocks of trials. The
vertical dashed lines indicate breaks between daily training sessions.

Figure 2. Electrode placements. The tips of stimulation electrodes in the
control rats given paired (black circles) or unpaired (gray circles)
training were in or within 0.2 mm of the medial division of the medial
geniculate (left). Rats given intentionally misplaced electrodes had tip
placements 1.0 mm medial and lateral to the medial division of the
medial geniculate (right). Numbers indicate the anterior-posterior stere-
taxic coordinates of the sections relative to bregma (This figure was re-
produced with permission from Elsevier © 1998, Paxinos and Watson
1998).
Inferior colliculus lesions

The inferior colliculus contralateral to the conditioned eye was damaged extensively. Figure 3 depicts the largest and smallest inferior colliculus lesions. The lesions were large but restricted to the inferior colliculus. The most extensive damage was in anterior nuclei, which is where most of the labeled axon terminals were found following biotinylated dextran amine (BDA) injections into the MATN (see Experiment 2, below).

Experiment 2: MATN project to the pontine nuclei

The results of Experiment 1 indicate that stimulation of the MATN can serve as a sufficient CS for eyelid conditioning. If the MATN are part of the auditory CS pathway, they should send axonal projections to the pontine nuclei either directly or indirectly. Anterograde and retrograde tract tracing were used in Experiment 2 to determine whether there are direct projections from the MATN to the pontine nuclei. BDA was used as an anterograde tracer to determine whether the MATN send axon terminals to the pontine nuclei. Fluorogold (FG) injected into the basal pontine nuclei was used as a retrograde tracer.

The injection sites for BDA included the medial nucleus of the medial geniculate, dorsal posterior intralaminar nucleus, and suprageniculate nucleus (Fig. 4A). Some of the BDA was taken up by cells in the visual nuclei that are dorsal to the medial geniculate. The injection sites for FG included the dorsolateral, lateral, and ventral pontine nuclei (Fig. 5A).

BDA-labeled axon terminals were evident in the dorsolateral, lateral, medial, and ventral pontine nuclei ipsilateral to the injection site (Fig. 4C, D). No terminals were observed in the contralateral pontine nuclei. The terminals appeared to have arisen from descending fiber projections through the area of the ipsilateral lateral lemniscus and the dorsal pons. The same pattern of terminal labeling in the pontine nuclei was seen in all of the rats given BDA infusions.

Retrogradely labeled neurons were seen in the medial division of the medial geniculate, the most dorsal part of the posteriorntralaminar nucleus, posterior limitans nucleus, and throughout the suprageniculate nucleus in rats given infusions of FG into the pontine nuclei (Fig. 5B, C). The thalamic retrograde labeling was exclusively ipsilateral to the infusion site.

Labeled neurons and terminals were observed throughout the ipsilateral inferior colliculus (Figs. 4B, 6B). Labeling was denser in the anterior external inferior colliculus relative to the central nucleus. Labeled neurons were seen in the inferior colliculus with BDA (Fig. 4B) and FG (Fig. 6B). As with the pontine nuclei, the terminal labeling with BDA was ipsilateral in the inferior colliculus.

The cochlear nuclei exhibited retrogradely labeled neurons bilaterally, primarily in the ventral nucleus. The density of labeled neurons appeared to be greater in the contralateral cochlear nuclei, as seen in rabbits (Steinmetz et al. 1987) (Fig. 6A).

Extensive terminal labeling and retrograde labeling were observed in the intermediate and deep layers of the superior colliculus and the anterior pretectal nucleus (Fig. 7). A particularly dense area of labeled neurons was seen in the deep superior colliculus just dorsal to the PAG in the rats given infusions of FG into the pontine nuclei (Fig. 7A). The dorsal and lateral PAG were also strongly labeled. The PAG had a high density of retrogradely labeled cells and terminals. The PAG labeling was exclusively ipsilateral to the injection sites.

The ipsilateral cerebral cortex showed extensive retrograde labeling, as seen in previous anatomical studies (Legg et al. 1989). Labeled neurons were mostly seen in layer 5 (Fig. 6C). Sensory and motor areas were labeled heavily relative to cingulate, perirhinal, and entorhinal areas.

Discussion

Rats given electrical stimulation of the medial auditory thalamus as a CS paired with a peripheral US showed rapid acquisition of
eyeblink conditioning. Rats given unpaired presentations of the stimulation CS and the peripheral US showed no increase in eyeblink responses across training trials; indicating that acquisition of eyeblink conditioning in the paired group was due to associative learning. Conditioned responses produced by the paired group increased in amplitude across training trials, and the peak amplitude occurred just prior to the onset of the US, as seen with conditioning in rats using peripheral CSs (Nolan and Freeman 2006). Rats with electrodes intentionally misplaced 1.0 mm lateral to the medial nucleus of the medial geniculate showed no conditioning. Rats with electrodes placed 1.0 mm medial to the medial nucleus of the medial geniculate showed acquisition of eyeblink conditioning but at a substantially lower rate than rats with accurately placed electrodes. Acquisition of eyeblink conditioning with auditory thalamic stimulation as the CS was not impaired by lesions of the inferior colliculus, suggesting that the stimulation probably did not produce conditioning through thalamic projections to the inferior colliculus.

The conditioning seen in rats given paired presentations of auditory thalamic stimulation and a peripheral US indicates that activation of the medial auditory thalamus is sufficient for establishing associative eyeblink conditioning in rats. Moreover, rats in the paired group showed a rate of acquisition that was substantially faster than seen with tone or light CSs in our laboratory, reaching an asymptotic level of conditioning within the first 100-trial training session. Asymptotic conditioning using a peripheral CS is typically seen after 200 training trials. The rapid conditioning seen in the paired group suggests that the projection from the medial auditory thalamus to the pons is very effective and sufficient to serve as a CS when other sensory projections to the pons are not activated.

Stimulation of the ventral medial geniculate as a CS was ineffective for establishing eyeblink conditioning in the current study. The ventral nucleus of the medial geniculate provides a large proportion of the thalamic input to the primary auditory cortex (Winer 1992), which then projects directly to the pontine nuclei. Stimulation of the ventral medial geniculate or auditory cortex has been used as a CS to establish eyeblink conditioning in rabbits (Knowlton and Thompson 1992; Nowak et al. 1999). Auditory cortex stimulation as a CS resulted in robust eyeblink conditioning that was cerebellum-dependent, even though the cerebral cortex is not essential for eyeblink conditioning (Knowlton and Thompson 1992). Stimulation of the ventral medial geniculate was moderately effective as a CS when relatively strong stimulation parameters were used (150 µA, 200 Hz, 1.2 msec) (Nowak et al. 1999). It is possible that the stimulation parameters used in the current study (30–100 µA, 200 Hz, 0.1 msec) were
simply insufficient for activating the thalamo-cortical auditory system. On the other hand, the parameters used by Nowak et al. (1999) may have produced conditioning through current spread to the neighboring MATN. The failure to obtain conditioning in the rats given stimulation of the ventral medial geniculate may, therefore, be consistent with the findings of the previous studies.

In contrast to the ventral medial geniculate, stimulation of the anterior pretectal nucleus as a CS produced a small amount of eyeblink conditioning. The anterior pretectal nucleus has been implicated as part of the visual CS pathway for eyeblink conditioning in rabbits (Koualtidis et al. 1988) and projects directly to the pontine nuclei (Fig. 7B). However, there are no published reports describing the use of pretectal stimulation as a CS in eyeblink conditioning. The modest conditioning seen in the rats with electrodes placed in the anterior pretectal nucleus may have been due to stimulation of its projection to the pontine nuclei. The relative insufficiency of pretectal and ventral medial geniculate stimulation as a CS in the current study suggests that the medial auditory thalamic stimulation did not produce conditioning by stimulating neighboring nuclei.

Anterograde and retrograde neural tract tracers provided evidence for direct projections from the MATN to the pontine nuclei. The projections from the medial auditory thalamus to the basolateral pons were distributed across the dorsolateral, lateral, medial, and ventral nuclei. It is not clear which pontine nuclei are essential for auditory CS input to the cerebellum in rats. In rabbits, the dorsolateral and lateral nuclei have been implicated as the essential nuclei for an auditory CS. Lesions of the lateral and dorsolateral pontine nuclei abolish nictitating membrane CRs to auditory but not visual CSs (Steinmetz et al. 1987). Selective lesions of different pontine nuclei have not been performed in rat eyeblink conditioning studies, but a unit recording study of auditory CS-elicited neuronal activity in the pontine nuclei in infant rats revealed auditory responses in the dorsolateral, lateral, medial, and ventral nuclei (Freeman and Muckler 2003). The rodent auditory CS pathway may be more distributed among the pontine nuclei relative to rabbits, suggesting that each of the auditory thalamo-pontine projections observed in the current study could play a role in transmission of CS stimulation to the cerebellum.

Precise infusions of anterograde tracers into the subnuclei of the medial auditory thalamus are needed to determine whether each thalamic nucleus projects to a different pontine nucleus or if the thalamic nuclei have overlapping projections. The possibility of overlapping projections is suggested by the findings of a previous study that found that lesions of the auditory thalamus prevented acquisition of eyeblink conditioning only with complete destruction of the medial division of the medial geniculate, posterior intralaminar nucleus, suprageniculate, and the posterior limitans nucleus (Halverson and Freeman 2006). Lesions that spared any of the nuclei had very little effect on eyeblink conditioning. An implication of this finding is that the thalamic nuclei provide parallel input that overlaps in the region(s) of the pons that is essential for auditory eyeblink conditioning.

As noted in previous studies, a projection from the inferior colliculus to the MATN, as well as a feedback connection from the medial auditory thalamus to the inferior colliculus, was observed (LeDoux et al. 1987; Winer 1992). The inferior colliculus in turn projects to the basolateral pontine nuclei (Fig. 6B; Kawamura 1975). It was therefore possible that stimulation of the medial auditory thalamus produced conditioning by activating the inferior colliculus and its projection to the pons. This possibility was partially ruled out by the demonstration of unimpaired acquisition in the group given large electrolytic lesions of the inferior colliculus. The results of the current study generally suggest that auditory CS signals are conveyed to the cerebellum via a serial pathway from the inferior colliculus to the medial auditory thalamus and then to the basolateral pontine nuclei.

The current findings combined with the results of a previous study (Halverson and Freeman 2006) indicate that the medial auditory thalamic contribution to the CS pathway is unilateral. Lesions of the medial auditory thalamus that were contralateral to the conditioned eye severely impaired auditory eyeblink conditioning (Halverson and Freeman 2006). In the current study, stimulation of the contralateral medial auditory thalamus was sufficient for eyeblink conditioning, and the projections from the medial auditory thalamus to the pontine nuclei were exclusively ipsilateral. The projections from the inferior colliculus to the medial thalamus were also ipsilateral. These findings suggest that although there are multiple crossed projections within the mammalian auditory system, the auditory CS pathway in eyeblink conditioning includes a unilateral projection from the inferior colliculus to the medial thalamus and then to the pons. It will be important in future studies to determine whether the auditory CS pathway component of the pontine projection to the cerebellum is also unilateral.

Projections from the PAG to the pontine nuclei are an unexpected finding of the current study. The PAG is well known as an important component in the behavioral organization of fear (LeDoux et al. 1988; Fanselow 1994), but its role in eyeblink conditioning is not well characterized. Recordings of unit activity from the PAG during eyeblink conditioning in rabbits revealed stimulus-elicited and response-related activity (McCormick et al. 1983). The effects of PAG lesions or inactivation during eyeblink conditioning have not been reported. However, lesions of the amygdala slow acquisition of eyeblink conditioning (Lee and Kim 2004; Blankenship et al. 2005). It is possible that amygdala modulation of the acquisition of eyeblink conditioning is mediated through its projection to the PAG and the PAG projection to the pons.

The current findings and those of a previous study (Halverson and Freeman 2006) demonstrated that the MATN play an important role in auditory eyeblink conditioning. The medial auditory thalamus could serve as part of the sensory input that conveys tone information to the cerebellum. Another possibility is that the thalamic nuclei mediate the short latency pathway from the cochlear nuclei to the pons, providing gain to boost the auditory signal en route to the cerebellum. It is also possible that conditioning-specific modifications of thalamic activity (Disterhoft and Olds 1972; Gabriel et al. 1975; Ryu and Weinberger 1978; Birt and Olds 1981; Supple and Kapp 1989; Edeline and Weinberger 1992; Lennartz and Weinberger 1992; McClellon et al. 1995; O’Connor et al. 1997; Maren et al. 2001; Poremba and Gabriel 2001) are necessary for this hypothetical modulatory boost of auditory input to the pons.
Medial auditory thalamus and eyelink conditioning

Materials and Methods

Experiment I

Subjects

Subjects were 17 adult Long Evens male rats (250–300 g). –150 d of age at the beginning of the experiment. The rats were housed in Spence Laboratories of the Psychology at the University of Iowa with a 12-h light-dark cycle, with light onset at 7:00 am.

Groups

Rats were assigned to groups that received paired (n = 13) or unpaired (n = 4) training. Four of the rats in the paired group were given intentionally misplaced electrodes (two in the anterior pretectal nucleus, two in the ventral nucleus of the medial geniculate), and three were given inferior colliculus lesions.

Surgery

One week prior to training, rats were removed from their home cage and anesthetized by an i.p. injection of sodium pentobarbital (80 mg/kg). An i.p. injection of atropine sulfate (0.45 mg/kg) was administered to reduce respiratory tract secretions. After onset of anesthesia, a bipolar stimulating electrode (Plastics One, Inc.) was implanted in the right medial geniculate. The stereotaxic coordinates were −5.7 mm posterior to bregma and −2.8 mm lateral to the midline. The bipolar stimulating electrode was lowered −6.1 mm below the surface of the skull and secured to the skull with dental acrylic. Four rats were implanted with intentionally misplaced bipolar stimulating electrodes at 1.6 mm (n = 2) or 3.6 mm (n = 2) lateral to the midline. A subset of the rats given paired training was given electrolytic lesions of the inferior colliculus ipsilateral to the stimulation electrode (n = 3). The lesions were produced by passing 1.0 mA of DC current for 10 sec at three sites via an insect pin insulated with epoxyite. The stereotaxic coordinates were 8.3, 8.8, and 9.2 mm posterior to bregma; 2.2, 2.0, and 1.9 mm lateral to midline; and 4.2, 4.6, and 4.3 mm below the skull surface. Skull holes were sealed with bone wax.

All of the rats were implanted with differential electromyographic (EMG) electrodes in the left upper eyelid muscle (orbicularis oculi), and a ground electrode was attached to a stainless steel skull screw. The EMG electrode leads terminated in gold pins held in a plastic connector, which was secured to the skull with dental acrylic. A bipolar stimulating electrode (Plastics One, Inc.) for delivering the shock US was implanted in a plastic connector immediately caudal to the left eye. The bipolar electrode terminated in a plastic connector that was secured to the skull with dental acrylic.

Conditioning apparatus

The conditioning apparatus consisted of a small-animal sound attenuation chamber (BRS/LVE) with a small-animal operant chamber (BRS/LVE) contained inside. The rats were kept in the operant chamber during conditioning. Cables with connectors for the EMG, US, and CS electrodes were attached to a commutator. The electrode leads from the rat’s head stage were connected to peripheral equipment and a desktop computer. Computer software controlled the delivery of stimuli and the recording of eyelid EMG activity (JSA Designs). The US was delivered through a stimulus isolator (World Precision Instruments). EMG activity was recorded differentially, filtered (500–5000 Hz), amplified (2000 ×), and integrated (time constant = 20 msec). MGN stimulation was triggered through a programmable stimulator (Master 8, A.M.P.L.), which controlled signal input to the stimulus isolator (World Precision Instruments) that delivered the electrical stimulation.

MATN stimulation

Electrical stimulation of the medial auditory thalamus functional as the CS, which was administered in a 200-Hz train of 0.1-msec biphasic pulses for 300 msec. The stimulation threshold for the CS was found before training by setting the stimulating current to elicit an observable behavioral response and then decreasing the current in 5-µA increments until no movement was detected. Observable movements included, but were not limited to, eye blinks, orienting responses, ear movements, and head movements. A maximum stimulation threshold was set at 100 µA (range = 30–100 µA). The most typical behavioral response elicited by stimulation was a discrete movement of the left ear.

Paired training

Rats were given three paired training sessions. The paired training sessions consisted of 100 trials; each with 90 trials of the stimulation CS paired with the shock US (10 msec, 1.0–2.0 mA) and 10 stimulation CS-alone trials, occurring on every 10th trial. The CS-alone trials were included in order to assess behavioral responses (integrated EMG activity) uncontaminated by unconditioned responses (URs). The interstimulus interval for paired trials was 290 msec. Trials were separated by an intertrial interval (ITI) that averaged 30 sec. Behavioral data were examined from computer records of EMG responses. CRs were defined as responses that crossed a threshold of 0.4 V above the baseline activity during the CS period, but at least 80 msec after CS onset, to avoid contamination of the CR measures by the startle (alpha) response. The amplitude, onset latency, and peak latency of the CR were measured on CS-alone trials in which a CR occurred.

Unpaired training

Rats in the unpaired control group were given three 200-trial sessions of explicitly unpaired presentations of the CS and US. The same time durations for the CS and US were used as in the paired procedure. The ITI was set to average 15 sec to match the total time spent in the conditioning chamber and the temporal distribution of CS and US presentations with the paired groups. The method for defining CRs was the same as that used in the paired procedure.

Histology

After training was completed, rats were euthanized with a lethal injection of sodium pentobarbital (90 mg/kg) and transcardially perfused with 0.1 M buffered physiological saline, followed by a 10% formalin solution. The brains were post-fixed in formalin for 2 d and then put in a solution of 30% sucrose in PBS before sectioning. The brains were sectioned at 50 µm with a sliding microtome. Sections were then stained with thionin. The bipolar stimulating electrode location and extent of the inferior colliculus lesions was determined by examining consecutive sections.

Experiment 2

Subjects

Subjects were seven adult Long Evens male rats (250–300 g). –150 d old at the beginning of the experiment. The rats were housed in Spence Laboratories of the Psychology at the University of Iowa with a 12-h light-dark cycle, with light onset at 7:00 am.

Surgery

The rats were removed from their home cage and anesthetized by an i.p. injection of sodium pentobarbital (80 mg/kg). An i.p. injection of atropine sulfate (0.45 mg/kg) was administered to reduce respiratory tract secretions. After onset of anesthesia, pressure injections of 0.05 µL of 4% FG were made into the lateral pontine nuclei (n = 4) or 0.05 µL of a 10% BDA solution (Molecular Probes) into the medial division of the medial geniculate (n = 3). BDA and two of the FG infusions were delivered at a rate of 1.0 µL/h through a 30-gauge cannula. Two of the FG injections were made through a glass pipette with a tip diameter of 70 µm. The cannula or glass pipette remained in position for 15 min before removal. The post-injection survival period was 14 d.

Anterograde labeling with BDA

Two weeks following BDA infusion, the rats were perfused with 0.1 M PBS followed by 4% paraformaldehyde. After a post-fixation period of 24 h, the brains were sectioned on a freezing
microtome at 50 μm. Visualization of BDA was accomplished by using the standard avidin–biotin complex (ABC) method (Vectorstain ABC Elite Kit, Vector Laboratories). The tissue slices were incubated in the ABC solution for 60 min at room temperature. Following three washes in phosphate buffer, the tissue was incubated in 0.05% diaminobenzidine in phosphate buffer with hydrogen peroxide (0.01%) added to the solution. The tissue was incubated until a clear reaction developed (~15 min). The labeled sections were examined using a light microscope.

**Retrograde labeling with FG**

At the end of the survival period, the rats were perfused with 0.1 M PBS followed by 4% paraformaldehyde. After a post-fixation period of 24 h, the brains were sectioned on a freezing microtome at 50 μm. Visualization of FG was accomplished by immunohistochemistry with free floating sections. The brain sections were pretreated with H2O2 to block endogenous peroxidase, followed by incubation in normal goat serum. They were then incubated in a rabbit primary antibody for 72 h at 4°C followed by incubation in the biotinylated secondary antibody. The tissue slices were incubated in the ABC solution for 60 min at room temperature. Following three washes in phosphate buffer, the tissue was incubated in 0.05% diaminobenzidine in phosphate buffer with hydrogen peroxide (0.01%) added to the solution. The tissue was incubated until a clear reaction developed (~5 min). The labeled sections were examined using a light microscope.

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