Specific auditory memory induced by nucleus basalis stimulation depends on intrinsic acetylcholine

Alexandre A. Miasnikov, Jemmy C. Chen, Norman M. Weinberger *

Center for the Neurobiology of Learning and Memory, 309 Qureshey Research Laboratory, Department of Neurobiology and Behavior, University of California, Irvine, CA 92697-3800, USA

**Abstract**

Although the cholinergic system has long been implicated in the formation of memory, there had been no direct demonstration that activation of this system can actually induce specific behavioral memory. We have evaluated the "cholinergic-memory" hypothesis by pairing a tone with stimulation of the nucleus basalis (NB), which provides acetylcholine to the cerebral cortex. We found that such pairing induces behaviorally-validated auditory memory. NB-induced memory has the key features of natural memory: it is associative, highly-specific and rapidly induced. Moreover, the level of NB stimulation controls the amount of detail in memory about the tonal conditioned stimulus. While consistent with the hypothesis that properly-timed release of acetylcholine (ACh) during natural learning is sufficient to induce memory, pharmacological evidence has been lacking. This study asked whether scopolamine, a muscarinic antagonist, impairs or prevents the formation of NB-induced memory. Adult male rats were first tested for responses (disruption of ongoing respiration) to tones (1–15 kHz), constituting a pre-training behavioral frequency generalization gradient (BFGG). Then, they received a single session of 200 trials of a tone (8.00 kHz, 70 dB, 2 s) paired with electrical stimulation of the NB (100 Hz, 0.2 s). Immediately after training, they received either scopolamine (1.0 mg/kg, i.p.) or saline. Twenty-four hours later, they were tested for specific memory by obtaining post-training BFGGs. The saline group developed CS-specific memory, manifested by maximum increase in response specific to the CS frequency band. In contrast, the scopolamine group exhibited no such memory. These findings indicate that NB-induced specific associative behavioral memory requires the action of intrinsic acetylcholine at muscarinic receptors, and supports the hypothesis that natural memory formation engages the nucleus basalis and muscarinic receptors.

© 2008 Elsevier Inc. All rights reserved.

**1. Introduction**

Acetylcholine (ACh) has long been implicated in learning and memory (Deutsch, 1971; Flood, Landry, & Jarvik, 1981) and continues to be the focus of extensive research (Power, Vazdarjanova, & McGaugh, 2003). For example, pharmacological blockade of the cholinergic system impairs many forms of memory in both animals and humans (Anagnostaras, Maren, Sage, Goodrich, & Fanselow, 1999; Chudasama, Dalley, Nathwani, Bouger, & Robbins, 2004; Lozano, Armengaud, & Gauthier, 2001; Múnera, Gruart, Muñoz, & Delgado-García, 2000; Ravel, Elaagouby, & Gervais, 1994; Rudy, 1996; Schön, Atri, Hasselmo, Tricarico, LoPresti, & Stern, 2005). Cholinergic agonists and cholinesterase antagonists can facilitate memory (Introini-Collison & McGaugh, 1988; Stratton & Petrinovich, 1963), promote recovery of memory from brain damage (Russell, Escobar, Booth, & Bermudez-Rattoni, 1994) and achieve rescue from memory deficits in transgenic mice (Fisher, Brandeis, Chapman, Pittel, & Michaelson, 1998). In addition, several non-cholinergic treatments that facilitate memory, such as adrenergic agents and stress hormones, exert their effects via the cholinergic system (Salinas, Introini-Collison, Dalmaz, & McGaugh, 1997).

Such findings suggest that ACh is released during many types of learning and that its effects throughout the brain may promote the formation of memory and/or increase its strength. The nucleus basalis (NB) of the basal forebrain is a likely candidate for learning-related release of ACh to the cerebral cortex because it is the major source of cortical ACh (Bígíl, Woolf, & Butler, 1982; Johnston, McKinney, & Coyle, 1979; Luiten, Gaykema, Traber, & Spencer, 1987; Mesulam, Mufson, Wainer, & Levey, 1983; Rye, Wainer, Mesulam, Mufson, & Saper, 1984). Also, stimulation of the NB releases ACh in the cortex and produces cortical electroencephalographic (EEG) activation, the waking state accompanying most learning (Casamenti, Deffini, Abbamondi, & Pepeu, 1986; Celesia & Jasper, 1966; Détari, Juhasz, & Kokorelli, 1984; Détari, Rasmusson, & Semba, 1999; Jiménez-Capdeville, Dykes, & Myasnikov, 1997; Rasmusson, Clow, & Szerb, 1992, 1994; Rasmusson, Szerb, & Jordan, 1996).
If acetylcholine and the nucleus basalis are components of neural mechanisms that promote memory, then they might be expected to influence sensory processing, as memories are largely comprised of the record of sensory experience. Indeed, ACh and activation of the NB can modify cortical responses to and representations of sensory events in the primary auditory cortex (Ashe & Weinberger, 1991), somatosensory (Dykes, Tremblay, Warren, & Bear, 1991; Verdier & Dykes, 2001) and visual cortices (Gu, 2003). Most extensively studied in the primary auditory cortex (A1), application of ACh and anti-cholinesterase agents produces long-lasting alteration of acoustic frequency receptive fields (RFs) via muscarinic receptors (Ashe, McKenna, & Weinberger, 1989; McKenna, Ashe, & Weinberger, 1989). Stimulation of the NB produces atropine-sensitive, persistent modification of evoked responses in the auditory cortex, including facilitation of field potentials, cellular discharges and EPSPs elicited by median geniculate stimulation (Metherate & Ashe, 1991, 1993) in vitro, and facilitation of responses to tones in vivo (Edeline, Hars, Mao, & Hennevin, 1994; Edeline, Mao, Hars, & Hennevin, 1994; Hars, Mao, Edeline, & Hennevin, 1993; Hennevin, Edeline, Hars, & Mao, 1992; Hennevin, Mao, Hars, & Edeline, 1993).

Of particular relevance to memory, pairing a tone with NB stimulation produces shifts in frequency tuning that are specific to (i.e., directed toward) the conditioned stimulus (CS) (Bakin & Weinberger, 1996). As in the case of memory and tuning shifts that develop during classical and instrumental conditioning, NB-induced frequency plasticity is associative, highly-specific, rapidly established, discriminative and also consolidates (i.e., becomes stronger over time) and endures (Weinberger, 1998, 2003). NB-induced CS-specific RF plasticity requires the engagement of muscarinic receptors in A1 (Miasnikov, McLin, & Weinberger, 2001). Similar cholinergically-based, muscarinic receptor-dependent CS-specific tuning shifts have been found in the bat, demonstrating species generality (Ji, Gao, & Suga, 2001; Ji & Suga, 2003; Ji, Gao, & Suga, 2005; Ma & Suga, 2005). NB-induced associative tuning shifts presumably consist of more cells becoming preferentially responsive to the CS frequency and thus to an increase in the area of CS representation might be expected, as it is in the case of learning-induced plasticity (Recanzone, Schreiner, & Merzenich, 1993; Rutkowski & Weinberger, 2005). Such specific representational expansions have been confirmed (Kilgard & Merzenich, 1998; Kilgard, Pandya, Vazquez, Ghei, Schreiner, & Merzenich, 2001).

If the NB is normally engaged during associative learning to release ACh and promote memory formation, then pairing a tone with NB stimulation, in the absence of any standard unconditioned stimulus (e.g., food or shock) might be sufficient to induce genuine memory, as behaviorally defined. That, in fact, turned out to be the case. Rats trained with one tone paired with NB stimulation were later tested for specific memory by presenting many different tones, i.e., by obtaining behavioral frequency generalization gradients. A control group received tone and NB stimulation randomly. Memory was assessed by detecting changes in tone-elicited heart rate and respiration. Specific memory for frequency would be revealed by a generalization gradient having its peak (maximal response) at the CS frequency (Mackintosh, 1974; Mostofsky, 1965; Pavlov, 1927). The paired group in these studies did in fact exhibit this CS-specific gradient, whereas the unpaired group had no differential response to any tone (McLin, Miasnikov, & Weinberger, 2002a, 2003; Miasnikov, Chen, & Weinberger, 2006). Moreover, the amount of detail in memory about tonal frequency is controlled by the degree of NB activation; low levels of activation induce associative memory that sound is important without preserving the frequency of the CS whereas higher levels of activation also induce memory of CS frequency specificity (Weinberger, Miasnikov, & Chen, 2006). The nucleus basalis appears to be operating “downstream” of motivational systems because NB stimulation that induces memory is neither rewarding nor aversive (Miasnikov, Chen, Gross, Poytress, & Weinberger, 2008).

Overall, the evidence strongly supports the hypothesis that appropriately-timed, phasic release of ACh by the cortical terminals of the nucleus basalis is sufficient for the formation of behavioral memory. However, in the absence of relevant pharmacological findings, memory induction might be attributable to some non-cholinergic effects of stimulation of the nucleus basalis. For example, the NB contains GABAergic cells that release their transmitter in the cerebral cortex, where they inhibit cortical inhibitory interneurons, i.e., promote increased cortical excitation by disinhibition (Dykes, 1997; Freund & Meskenaite, 1992; Jiménez-Capdeville et al., 1997). Therefore, we asked whether scopolamine, a muscarinic antagonist, blocks NB-induced memory. We applied scopolamine after training in order to preclude affecting performance factors during training, such as attention and arousal (McGough, 1966).

2. Materials and methods

The materials and methods were generally the same as those previously reported (Miasnikov et al., 2006, 2008; Weinberger et al., 2006), and thus will be described only briefly. All procedures were performed in accordance with the University of California Irvine Animal Research Committee and the NIH Animal Welfare guidelines. During training and testing, subjects were continuously monitored by video cameras.

2.1. Subjects and surgery

The subjects were 16 adult male Sprague–Dawley rats (104 ± 17 days of age, 446 ± 68 g, mean ± SD), housed individually with ad libitum food and water, on a 12/12 h light–dark cycle (lights on at 7:15 AM). Following several days of adaptation to the vivarium, animals were handled and learned to sit calmly during attachment of a thermistor assembly and a cable to their skull pedestal. Under general anesthesia (sodium pentobarbital, 40 mg/kg i.p., Abbott Laboratories, North Chicago, IL), an 0.8-mm diameter stainless steel recording epidural screw electrode was inserted into the right primary auditory cortex at the locus showing the largest amplitude evoked potential (200–400 μV) to a contralateral noise burst. Two screws over the frontal sinus served as reference electrodes. A concentric bipolar stainless steel stimulating electrode (#SNEX-100x13, David Kopf Instruments, Tujunga, CA) was implanted through the contralateral (left) hemisphere (45° angle in the frontal plane at AP = −2.2, L 3.2; Paxinos & Watson, 1997) into the right nucleus basalis. The final locus was determined by obtaining 1–5 s of auditory cortical EEG activation to stimulation (200–500 μA, pairs of 0.2 ms opposite polarity pulses, 100 Hz, 200–300 ms trains; S88 stimulator, PSIU6 isolation units, Grass Instrument Co., Quincy, MA). A dental, acrylic pedestal was built with two aluminum hex-threaded standoffs embedded therein, and all leads connected to a miniature socket that could be led to a commutator via a multi-conductor cable. Subjects were allowed 1–2 weeks to recover from surgery.

2.2. Stimuli, recording, and data analyses

Training and testing took place while each subject was in a box (23 x 23 x 31 cm) supplied with fresh bedding and lined inside with acoustic-damping tile, contained in a double-walled acoustic chamber (Industrial Acoustics Company, Bronx, NY). Acoustic stimuli were pure tones (1.0–15.0 kHz, 2 s duration, cosine 10 ms rise/fall time [10–90%], 70 dB SPL) produced by TDT System 3 components (Tucker–Davis Technologies [TDT], Alachua, FL) and delivered to two loudspeakers calibrated for low (electrodynamic, #40-1421, RadioShack, Fort Worth, TX) and high (electrostatic, #ES-1, TDT) frequencies positioned 35 cm above the floor of the box. NBstim current used during training was several times weaker than that used during surgery because the absence of anesthesia greatly reduces the threshold for EEG activation, that was determined as described below.

To assess the induction of memory, we measured changes in the disruption of the ongoing respiration pattern to various tones before and following training (below). Respiration was detected as breathing-related thermal fluctuations with a glass-encapsulated thermistor attached to a lightweight pedestal-mounted assembly positioned in front of a naris. The details of this device, as well as those for the amplification of respiration signal were previously described (Miasnikov et al., 2006).

The output signal was fed to differential band-pass amplifiers (1–100 Hz), digitized by two A-D modules, one for on-line calculation of the autocorrelation function (ACF) [100 samples/s], the other for off-line analysis (2000 samples/s). The highest possible ACF value is 1.0, i.e., a perfect sinusoid. The ACF was used to present tones only when the subject was in a state of quiet waking. This corresponded to ACF values of 0.80–0.95 over a 4 s pre-tone period. Respiration patterns are characteristic of different behavioral states (Fig. 1). An ACF > 0.95 occurs during slow...
wave sleep. ACFs – 0.75 or lower occur during REM and during active exploration, both states characterized by irregular respiration. We chose the lower criterion of ACF = 0.80 as a conservative means of avoiding any contamination from more activated states, including grooming or the initiation of searching behavior while subjects were still recumbent. These states could have ACF values –0.40–0.75.

Trials meeting the criterion of baseline ACF = 0.80–0.95 for over 4 s were presented if the scheduled inter-trial interval period had passed (30–180 s). This state control was employed to avoid giving stimuli when very high levels of ACh were being released in the cortex, as during exploration or REM sleep (Giovannini, Rakovsky, Benton, Pazzaglia, Bianchi, & Peppe, 2001; Jasper & Tessier, 1971; Kametani & Kawamura, 1990; Marroso, Portas, Mascia, Casu, Fà, Giagheddu, Imperato, Gessa, 1995) to prevent a ceiling effect, thus promoting a physiologically-effective release of ACh by NBstm.

Offline analysis consisted of the calculation of Fast Fourier Transform (FFT) functions for a period of 4 s preceding a trial (Pre) and 24 s following tone onset (Post). Major changes in respiration occurred within 0.5–12.5 s after tone onset. During the quiescent state, the respiration signal is almost completely contained within the bandwidth of 0.975–2.925 Hz. The FFT data were used to calculate a “Respiration Change Index” (RCI), on a second-by-second basis. The index was sensitive to increases and decreases of both frequency and amplitude. RCIs were calculated as: $\text{RCI} = \frac{\text{ACm} - \text{AC0}}{\text{AC0}}$, where $\text{ACm}$ is the 4 s epoch of cortical activation (decrease in low frequency activity often accompanied by increase in gamma activity). The current levels used in subsequent training with NB stimulation did not elicit body movements.

To induce and subsequently evaluate stimulus-specific memory, we used the approach of acquiring behavioral baseline responses to many frequencies, then training with one frequency and testing the training effects with many frequencies.

2.3. Experimental design

The subjects were assigned to two groups, saline (n = 8) and scopolamine (n = 8). NB stm recovery from surgery was observed. NB stm thresholds were determined while subjects were in a quiet waking state. NB stm was delivered every few minutes at increasing levels starting at ~50 μA (100 Hz bipolar, 200 ms train) until stimulation reliably elicited 3–8 s epoch of cortical activation (decrease in low frequency activity often accompanied by increase in gamma activity). The current levels used in subsequent training with NB stimulation did not elicit body movements.

To induce and subsequently evaluate stimulus-specific memory, we used the approach of acquiring behavioral baseline responses to many frequencies, then training with one frequency and testing the training effects with many frequencies.

2.4. Effectiveness of NB stimulation on the EEG of the auditory cortex

To ensure that any differences of NB-induced memory were not due to differential effectiveness of NB stm, all subjects received tone and NB stm unpaired within a few days after completion of the post-training assessment (Day 4) of memory induction. This was necessary as it is not possible to determine the effects of NB stm itself during training because it is inevitably preceded by the CS tone which itself acquires the ability to alter the ongoing EEG (McLIn et al., 2003). Tones (8.00 kHz) were included to simulate as closely as possible the training conditions without the confound of tone preceding NB stm. Presentation of tones may formally have constituted an extinction session, the subject of which is beyond the scope of the present report.

We quantified changes induced by NB stm by calculating the power spectra of electroencephalographic (EEG) recordings obtained from the auditory cortical electrode. The epidural A1 signal was amplified and filtered (1000–, band-pass 1–1000 Hz), digitized at 500 samples/s and processed off-line. The FFT Power was calculated at a frequency resolution of 0.575 Hz for frequencies up to 59.965 Hz. The FFT data were used to calculate an EEG “Power Change Index” (EEG PCI), on a second-by-second basis for each EEG frequency band separately as follows: delta, 0.59.8–2.92; theta, 2.93–8.78; alpha, 8.79–14.62; beta1, 14.63–20.47; beta2, 20.48–33.15; gamma, 33.16–59.97. EEG PCIs were calculated on each trial as follows: EEG PCI = [(Post – Pre)/(Post + Pre)], where “Pre” was the first 2 s out of four immediately preceding a trial. The index was sensitive to both increases and decreases in EEG Power. A negative value indicated a decline and a positive value indicated a rise in power within a specified frequency band relative to its baseline (McLIn, Masni-kov, & Weinberger, 2002b). In addition, as the major EEG changes were a decrease in alpha power and an increase in gamma power, we calculated an “EEG Activation Index [EAI] to directly compare the EEG effectiveness of the groups: EAI = (Gamma PCI) + [Alpha PCI] × (–1.0). This yielded generally positive values that gave equal weighting to changes in both alpha and gamma power.

2.5. Location of electrodes

Following the completion of the experiment, an electrolytic lesion (4 ms pulses at 100 Hz, 500 μA for 20–60 s) was made with bipolar current through the stimulating electrode while the animal was under sodium pentobarbital anesthesia. The animal was then given an overdose of sodium pentobarbital and perfused through the heart with 0.1 M phosphate buffer (pH 7.3). The brain was removed and the Anterior-to-Posterior (AP) and Medialto-Lateral (ML) coordinates of the recording electrode relative to Bregma and midline, respectively, were precisely measured on the skull from the interior of

---

**Fig. 1. Behavioral state control. Examples of measures of respiration corresponding to four major behavioral states: exploration, quiet waking, slow-wave sleep, paradoxical sleep. The pattern of respiration is characteristic for each state (Weinberger et al., 2006).**

(A) When respiration is maximal (Autocorrelation function calculated for the 4-s epoch of respiration record exceeds 0.95), the animal responds poorly to tones and its EEG is typical of slow-wave sleep. (B) When respiration is somewhat less regular (AC = 0.80–0.95), the animal generally sits calmly with its eyes opened, is quite responsive to tones, and its EEG is somewhat less regular. (C) When respiration is chaotic, with many high amplitude transients, the EEG is low voltage-fast, typical of REM sleep. (D) When respiration is chaotic, with many high-amplitude transients, the EEG is low voltage-fast and the animal is exploring or grooming. The respiration AC range was set at 0.80–0.95 to maximize the probability of presenting training trials when subjects were in the quiet waking state. The relative regularity of respiration and sufficient reactivity to tones during this quiescent state provides an optimal baseline for the detection of tone-elicited disruptions.
the calvaria with a caliper at 0.1 mm resolution. The auditory cortex recording site, which had been determined during surgery by noise burst-induced local field potentials, was estimated by projecting the location of the epidural recording electrode onto the underlying cortical surface. The recording site location was subsequently verified by plotting the obtained coordinates of the mentioned cortical projection onto a stereotaxic map of the auditory and surrounding areas of cortex derived from the Paxinos and Watson (1997) atlas. The AP and ML data collected from individual subjects were combined with respect to experimental protocol, and the groups (saline vs. scopolamine) were compared using \( t \)-test to determine whether the location of the recording sites differed.

Following several days of post-fixation in paraformaldehyde solution with 0.8 M sucrose added for subsequent tissue cryoprotection, the brain was blocked and sectioned at 50 μm with a freezing microtome. The sections were mounted onto gelatin-coated slides, dried and stained for Nissl substance to recover the electrocorticographic lesion sites and thus to determine the actual locus of intracranial stimulation. The location of the stimulation site was then projected onto the closest outline of frontal (coronal) sections taken from the Paxinos and Watson (1997) atlas. Based on subcortical structure outlines, the AP, ML, and Dorsal-to-Ventral (DV) coordinates of the tip of the stimulating electrode were determined and converted into standardized atlas dimensions. The AP, ML, and DV data collected from individual subjects were combined with respect to experimental protocol, and the groups (saline vs. scopolamine) were compared using \( t \)-test to determine whether the location of the sites of stimulation differed.

### 3. Results

#### 3.1. Effect of post-training scopolamine on specific memory induced by NB stimulation

Behavioral frequency generalization gradients (BFGG) were obtained before and after pairing the CS and NBstm. The scopolamine and saline groups exhibited the same respiratory responses to tones of 1–15 kHz before training. In contrast, 24 h after training, they exhibited differential responses despite the fact that they had undergone identical pairing of an 8.00 kHz tone with NBstm. Fig. 4 provides examples of respiratory records for a single presentation of three tones for a saline animal and a scopolamine animal. In this example, both subjects exhibited little response to the CS tone (8.00 kHz), a lower (2.75 kHz) or higher (15.00 kHz) tone prior to training. However, after training, the saline animal displayed a large response to the CS frequency, but still no responses to the other tones. This pattern of response change is typical of NB-induced behavioral memory (McLin et al., 2002a; Miasnikov et al., 2006; Weinberger et al., 2006). In contrast, the scopolamine subject displayed neither post-training response to the CS frequency, nor to the other frequencies.

Fig. 5 summarizes the group data. Over the 200 presentations of the test tones before pairing, there was substantial response, particularly for the CS frequency band (6.25–9.75 kHz) and the high frequency band (11.5–15.0 kHz). This profile is generally consistent with the audiogram of the rat (Heffner, Heffner, Contos, & Ott, 1994). Although, as mentioned, animals were differentially sensitive to different frequencies (two-way ANOVA, frequency factor: \( F_{(2,41613)} = 50.21, p < .0001 \), the difference between the scopolamine and saline groups was not significant \( F_{(1,41613)} = 3.44, p > .06 \), and the interaction was not significant \( F_{(2,41613)} = 1.54, p > .20 \) (Fig. 5A). Conversely, the groups did differ after pairing (two-way ANOVA, group: \( F_{(1,41600)} = 25.37, p < .0001 \); both frequency factor \( F_{(2,41600)} = 37.85, p < .0001 \) and the group x frequency interaction \( F_{(2,41600)} = 3.44, p < .05 \) differed significantly, as well. Post-hoc tests revealed that this difference was limited to the CS frequency band, at which the saline group exhibited a larger response than the scopolamine group (Tukey's test: \( p < .00002 \)). The groups did not differ in response to either the low \( p > .10 \) or high \( p > .75 \) frequency bands (Fig. 5B). These findings indicate that the saline group had acquired a CS-specific memory following CS–NBstm pairing that could be expressed 24 h after training, whereas the scopolamine group did not. To better understand the nature of the differences between the groups, within-group analyses were also conducted. They revealed that the saline group had developed an absolute increase in responses to all three frequency bands, although only the increase at the CS frequency
band was statistically significant (Tukey: CS band, \( p < .02 \); low band, \( p > .25 \); high band, \( p > .45 \)). The scopolamine animals also exhibited some changes in responses to test frequencies, but none were significant (Tukey: CS band, \( p > .95 \); low band, \( p > .30 \); high band, \( p > .95 \)).

### 3.2. Location of electrodes

A summary of the location of the electrodes is presented in Fig. 6. The cortical sites of recording are summarized in Fig. 6A. All of the electrodes were located above the primary auditory cortex. The recording sites of the saline and scopolamine groups were intermingled and did not differ either in the AP (\( t_{(14)} = 0.59, p > .55 \), two-tailed \( t \)-test) or the ML (\( t_{(14)} = 1.38, p > .15 \)) dimensions. The center of the location area (grey oval) corresponds to the X–Y coordinates of the mean for the group on the flattened standardized cortical surface, and the horizontal and vertical edges of the oval correspond to the ranges from mean – SD to mean + SD for the AP (–4.81 ± 0.91 mm, mean ± SD) and the ML (9.26 ± 0.30 mm) coordinates, respectively.

Fig. 6B shows the implantation path of the stimulating electrode within the right hemisphere (the 45° angled light electrode trace from the upper left towards the lower right). The stimulation region is delineated with a dashed square around the lesion site found within the ventral portion of internal capsule. The NB stimulation sites from all subjects are shown in Fig. 6C. The stimulation sites of the saline and scopolamine groups were intermingled and did not differ either in the AP (\( t_{(14)} = 0.23, p > .80 \), two-tailed \( t \)-test), the ML (\( t_{(14)} = 1.26, p > .20 \)), or the DV (\( t_{(14)} = 1.13, p > .25 \)) dimensions. In general, the coordinates of the area of stimulation, as referenced to the coronal plane, were as follows: AP, 2.14 ± 0.25 mm (mean ± SD); the ML, 3.33 ± 0.40 mm; and the DV, 7.63 ± 0.22 mm. All the located stimulation sites were in the basal forebrain within structures containing corticopetal cholinergic cells, including those that project to the auditory cortex (Bigl et al., 1982; Johnston et al., 1979; Luiten et al., 1987; Mesulam et al., 1983; Rye et al., 1984).

### 3.3. Verification of same effectiveness of NB stimulation

It might be thought that the failure of the scopolamine group to display NB-induced memory was due to the use of a less effective NBstm. We compared levels of current used during pairing and found no significant difference: saline = 68 ± 9 \( \mu \)A; scopolamine = 67 ± 8 \( \mu \)A (mean ± SD); \( t_{(14)} = 0.349, p > .70 \), two-tailed. Also, the groups differed neither in weight (\( t_{(14)} = 1.710, p > .10 \)), nor age (\( t_{(14)} = 1.264, p > .20 \)). However, equal stimulating currents do not

---

**Fig. 3.** Experimental design. (A) The four main stages of the experiment used to obtain pre-training and post-training behavioral frequency generalization gradients (BFGG) for the scopolamine group. (The control ["saline"] group received identical treatment except for the injection of saline.) A fifth stage was used to quantify the effect on the EEG of NBstm alone; such quantification could not be obtained during the pairing session because NBstm was necessarily preceded by the CS tone which itself elicited conditioned EEG changes (McLin et al., 2003). The asterisk indicates that the effects of NBstm were determined within a few days after Day 4, when the last BFGG’s were obtained. (B) Detailed temporal relationships of stimuli for the various phases of the experiment: delivery of test tones (Days 1, 2 and 4), tone–NBstm pairing (Day 3) and unpaired NBstm (*)
guarantee equal physiological effectiveness. Therefore, we quantified the amount of change in the EEG induced by Nbstm.

Fig. 7A and B shows the quantified changes in power for all EEG frequency bands: (delta, theta, alpha, beta1, beta2 and gamma) for the saline and scopolamine groups, respectively. Band-wise statistical comparison at points of maximal change from baseline (EEG PCI levels immediately preceding Nbstm) showed that the effectiveness of Nbstm in the saline and scopolamine groups did not differ. The EEG power, expressed as EEG PCI, generally declined at alpha (saline, $0.301 \pm 0.013$ vs. scopolamine, $0.308 \pm 0.011$; mean ± SE, measured at 1.8 s; between-group difference [BGD]: $t(3039) = 0.385, p > .65$); beta1 (BGD 1.8 s; $p > .70$), and theta (BGD 1.8 s; $p > .95$) frequency bands. The power generally increased at beta2 (BGD 0.8 s; $p > .75$), and gamma bands (BGD 0.8 s; $p > .90$). The changes within the delta band were most consistent at a longer latency (3.8 s) but the difference between groups also was not significant (BGD $p > .65$).

With subjects in a state of quiet waking, the major elicited EEG changes are a decrease in the power of alpha activity and a concomitant increase in the power of gamma activity (McLin et al., 2002b; Miasnikov et al., 2008). Taken together, the two bands form an envelope that represents cortical activation. To determine whether this measure of activation was equal in both groups, we used an EEG Activation Index that combined data from both bands (see Section 2). These data are shown in Fig. 7C, which also gives an example of EEG response to Nbstm (“all bands”, “alpha” and “gamma”). The EAI exhibited essentially identical dynamics in the saline and scopolamine groups. The peaks of the functions were subjected to a statistical comparison and were found not to differ statistically ($t(2) = 0.271, p > .80$). Therefore, we conclude that the failure of the scopolamine group to exhibit memory is not attributable to its Nbstm effect being functionally weaker than in the saline group.

4. Discussion

4.1. Summary and validity of the findings

The induction of specific behavioral memory by direct intervention in the brain is unusual, if not unique. The present experiment is a logical extension of research on the putative release of acetylcholine from the nucleus basalis during natural learning. It is a con-
sequence of a model of learning-induced associative plasticity in the auditory cortex (Weinberger, 1995, 1998; Weinberger, Ashe, Metherate, Mckenna, Diamond, & Bakin, 1990). The model stipulates that as auditory thalamic neurons develop associative increases in discharge to conditioned stimuli, they activate the NB via the amygdala on training trials. This appropriately-timed activation of the NB is hypothesized to release ACh and promote the long-term storage of specific associative memory traces in the primary auditory cortex. We refer to this part of the model as the “NB/ACh Memory Induction Hypothesis”. Our experimental strategy has been to mimic natural associative learning by stimulation of the NB following onset of a tone. Prior findings have revealed the fact of the induction of specific, associative memory for acoustic frequency (McLin et al., 2002a, 2003; Miasnikov et al., 2006). Moreover, they have shown that the level of NB activation can control the degree of auditory frequency detail that is incorporated into memory: low levels of activation induce the memory that “sound is important”, without preserving the information on the frequency of the CS, whereas higher levels of activation also allow formation of the memory for the importance of the CS frequency (Weinberger et al., 2006).

Despite the fact that the NB is the major source of cortical acetylcholine, only indirect evidence has supported the hypothesis that the release and actions of ACh have a causal role in NB-induced memory. Support has been based on established relationships between the NB, ACh and the EEG. First, cholinolytic atropine blocks EEG activation and reduces cortical levels of ACh (Phillis & York, 1968; Szerb, 1964). Second, stimulation of the basal forebrain causes both the release of acetylcholine in the cortex and cortical EEG activation (shift from higher voltage slower waves to lower voltage faster waves) (e.g., Cape & Jones, 1998; Cape, Manns, Alonso, Beaudet, & Jones, 2000; Casamenti et al., 1986; Cesolia & Jasper, 1966; Détári, Juhász, & Kukorelli, 1983; Rasmusson et al., 1992, 1994, 1996) and corresponding neural activation (Jiménez-Capdeville et al., 1997). Third, specific neurotoxic lesions of NB cholinergic corticopetal neurons deplete the cortex of ACh and impair EEG activation, i.e., increase slow wave activity and decrease fast (e.g., gamma) waves (Bernston, Shafi, & Sarter, 2002; Wenk, Stoehr, Quintana, Mobley, & Wiley, 1994). Fourth, cholinergic corticopetal neurons in the NB exhibit a strong correlation with the EEG; increased rates of discharge occur with increased activation and vice versa (Chernysh & Weinberger, 1998; Duque, Balatoni, Détári, & Zaborszky, 2000; Détári et al., 1984; Détári et al., 1999; Manns, Alonso, & Jones, 2000a, 2000b; Szymusiak & McInty, 1986). Together, these findings support the “NB/ACh Memory Induction” hypothesis and are also consistent with the view that release of cortical ACh is at least part of the mechanism underlying the activation state of the EEG (Détári et al., 1999; Metherate & Ashe, 1992).

The present study has now provided direct evidence to support this hypothesis. The formation of specific associative memory by tone paired with stimulation of the nucleus basalis is impaired or blocked by scopolamine (Figs. 4 and 5). The most parsimonious interpretation of the findings is that NB-induced memory formation requires the release of acetylcholine in the brain and is not mediated by putative non-cholinergic effects of NB stimulation. However, there are alternative interpretations of the findings.

First, it might be argued that NB stimulation in this study did not induce genuine associative memory because the design did not include a non-associative control group. Such a control group was not employed because previous research has consistently shown that subjects receiving unpaired or random tone and NB stimulation do not form CS-specific memory. Instead they exhibit flat behavioral (respiratory and cardiac) generalization gradients and may even exhibit decreased responses to the “CS” attributable to tone repetition lacking consequences (McLin et al., 2002a, 2003; Miasnikov et al., 2006; Weinberger et al., 2006).

Second, memory induction could have failed because the scopolamine group received less effective NB stimulation. However, there was no significant difference in the level of NB stimulation and NB stimulation produced the same type and magnitude of EEG activation in both groups. Quantitative analyses revealed significant differences neither in comparisons of individual EEG bands nor in the integral EEG Activation Index that combined data on dominant representative EEG bands, i.e., alpha and gamma (Fig. 7). Thus, differences in the expression of memory 24 h after training were not the result of a lesser effectiveness of NBstim in the scopolamine group.
Third, it might be thought that scopolamine could have interfered with non-mnemonic processes. However, any scopolamine-induced effects on putative performance factors, such as attention and arousal changes during tone–NBstm pairing, can be ruled out because scopolamine was administered post-training. Also, testing for memory was delayed for 24 h, so that the drug was not present at the time of behavioral assessment of memory.

Fourth, given that scopolamine was administered systemically, it was possible that the failure of this group to exhibit specific memory was caused by peripheral effects of the drug. However, an extensive literature, covering numerous tasks and several species including the rat, has established that systemically-administered scopolamine hydrobromide trihydrate affects learning and memory via the brain. These findings include tone and context fear...
conditioning (rat) (Anagnostaras et al., 1999), avoidance conditioning (mouse) (Flexner, Flexner, & Church, 1991), inhibitory avoidance (rat) (Goto, Kuzuya, Endo, Tajima, & Ikari, 1990), negative patterning (rat) (Moran, 1992), auditory and visual discrimination (rat) (van Haaren & van Hest, 1989), odor discrimination (rat) (De Rosa & Hasselmo, 2000), conditional discrimination (rat) (Wan, Pang, & Olton, 1997), conditioned taste aversion (rat) (Ramírez-Lugo, Miranda, Escobar, Espinosa, & Bermúdez-Rattoni, 2003), 5-choice serial reaction time (monkey) (Spinelli, Ballard, Feldon, Higgins, & Pryce, 2006), operant specific number (rat) (van Haaren & van Hest, 1989), spatial discrimination (rat) (Steckler & Holsboer, 2001), spatial memory (rat) (Pitsikas, 2007), working memory (rat) (Shannon & Yang, 2007), reference memory (rat) (Miyagawa, Honma, & Sato, 1995), recognition memory (rat) (Woolley, Marsden, Sleight, & Fone, 2003), paired associates (human) (Atri et al., 2004) and free recall (human) (Danion et al., 1990). Thus, it seems highly likely that the impairing effects of scopolamine on NB-induced specific memory were caused by the block of muscarinic receptors in the brain. The present findings are agnostic regarding any possible critical loci of muscarinic action within the brain. To address these issues, future studies will need to employ locus-specific and receptor-specific manipulation of cholinergic neurotransmission.

4.2. Characteristics of nucleus basalis-induced memory

Specific memory induction by appropriately-timed, paired electrical stimulation of the cholinergic nucleus basalis is noteworthy for several reasons. First, NB stimulation sufficient to induce memory is neither rewarding nor punishing (Miasnikov et al., 2008). This contrasts with memory induction that might be induced by brain stimulation that is motivationally significant, such as rewarding stimulation of the ventral tegmentum (Schultz, 2001) and the nucleus accumbens (Day & Carelli, 2007), or aversive stim-

Fig. 7. Effectiveness of NBstm (200 presentations) on the auditory cortex (levels of EEG activation) after completion of the training/test protocol in the saline and scopolamine groups. (A and B) Group mean EEG spectral changes. For each frequency band, changes were computed as the EEG Power Change Index: EEG PCI = (Post - Pre)/(Post + Pre) where the “Pre” period was the mean of the first 2 s out of four immediately preceding tone onset and post measures were calculated for consecutive periods of 1 s. Note major effects are an increase in gamma (closed circles) and a decrease in alpha (closed triangles) power. The EEG changes are comparable in the two groups. (C) Direct comparison of the levels of EEG activation produced by NBstm in saline (black circles) and scopolamine (open squares) groups (mean ± SE). The graph shows the mean “EEG Activation Index” (EAI) for NBstm alone (see Section 2). Note that the responses to NBstm were robust in both groups, and there was no significant difference between the peak magnitudes of the EAI values (p > .80 for designated data points). Insert: Example of auditory cortical EEG activation by NBstm (200 ms, 100 Hz, 64 μA bipolar stimulation) observed for “All Bands”, and the two frequency bands that exhibited the largest changes in power (“Alpha” and “Gamma”) in an animal from scopolamine group. “All Bands”: original records obtained with band-pass filters set at 1–1000 Hz. “Alpha” and “Gamma”: corresponding records band-passed with digital filters set at 8.8–14.6 Hz to emphasize alpha and 33–59 Hz to emphasize gamma bands, respectively. An arrow shows the moment of NB electrical stimulation. Note the EEG activation, including a distinct decrease in higher voltage, slower waves (“Alpha”) and increase in lower voltage faster waves (“Gamma”).
ulation of the periaqueductal gray (Behbehani, 1995), intralaminar thalamic nuclei or anterior cingulate cortex (Seward & Seward, 2002). The effects of substitution of a brain reward or punishment for an environmental reward or punishment, while interesting, would hardly be surprising. Thus, it seems likely that the memory-inducing function of the NB is “downstream” of motivational detection and the assignment of positive or negative valence. This location in the information processing stream should enable the NB to promote memory across a variety of situations and types of motivation.

Second, the present findings provide, in part, an anatomical–physiological basis for the actions of cholinergic agents on memory. Certainly, other groups of cholinergic cells are involved in learning and memory. Recent studies of the magnitude and release of ACh have found locus-dependent competition (McIntyre, Pal, Marriott, & Gold, 2002) or cooperation between different structures (McIntyre, Marriott, & Gold, 2003). The roles of various cholinergic cells in the numerous processes underlying learning and memory might be clarified by using the strategy of natural learning mimicry, i.e., by determining the extent to which their activation might induce specific memory. Similar considerations apply to other neuromodulatory systems, such as noradrenergic and serotonergic. Thus, pairing a sensory stimulus with activation of the locus coeruleus and raphe, respectively, could be undertaken given the present state of knowledge of the nucleus basal and memory induction.

Third, the cholinergic induction of specific behavioral memory raises the issue of what sort of memory has been induced. In the absence of an external biologically-significant reinforcer, what is the nature of an association with the tonic conditioned stimulus? We suggest that the subjects learned that “8.00 kHz is important”, nothing more and nothing less. In short, they would now associate the CS frequency with behavioral relevance, although having no information on why the CS tone has become relevant. Such a memory would increase the salience of 8.00 kHz compared to other tones, but by itself would not predict any sensory or motivationally-significant reinforcer. This hypothesis can be tested by determining if this “information” can be transferred to a new situation. For example, increased salience to the CS frequency might interfere with an ongoing task or even retard new learning. Even more interesting, subjects might be able to use this information to facilitate the solution of a novel problem.

Beyond such transfer studies, NB-induced specific memory accomplishes the task of a “dissection” of memory. Workers have long recognized that associative memories have several components, including sensory content, associations between discrete stimuli, between a stimulus and a context, hedonic content, and even the outcome of greater understanding of the “causal fabric” of the environment. NB-induced memory may provide for an elemental memory in which the sensory content of the signal stimulus is encoded and stored without many of the usual associations or memory components. Such artificially-induced memory should be subject to a simpler and more elemental analysis than natural memories. Already, there is some characterization of the neural underpinning of NB-induced memory. It is accompanied by a retuning of the primary auditory cortex to favor processing of signal frequencies (reviewed in Weinberger, 2007). Such an increase in cells responding preferentially to signal stimuli of increased salience might constitute a neurobiological substrate of a “memory code” for the increased behavioral importance of environmental events (Rutkowski and Weinberger, 2005). Given the feasibility of inducing specific, behaviorally-validated memories and the findings to date, this approach to the neurobiology of learning and memory holds considerable promise.

Acknowledgments

We thank Nataliya Gross, Gabriel K. Hui, Julia Martinson and Jacque Weinberger for assistance. This study was funded by the NIDCD/NIMH, DC-02938.

References


Activation to acquisition: Functional aspects of the basal forebrain cholinergic system (pp. 325–345). Boston: Birkhäuser.


