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Histone Deacetylase Inhibition via RGFP966 Releases the Brakes on Sensory Cortical Plasticity and the Specificity of Memory Formation

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Research over the past decade indicates a novel role for epigenetic mechanisms in memory formation. Of particular interest is chromatin modification by histone deacetylases (HDACs), which, in general, negatively regulate transcription. HDAC deletion or inhibition facilitates transcription during memory consolidation and enhances long-lasting forms of synaptic plasticity and long-term memory. A key open question remains: How does blocking HDAC activity lead to memory enhancements? To address this question, we tested whether a normal function of HDACs is to gate information processing during memory formation. We used a class I HDAC inhibitor, RGFP966 (C21H19FN4O), to test the role of HDAC inhibition for information processing in an auditory memory model of learning-induced cortical plasticity. HDAC inhibition may act beyond memory enhancement per se to instead regulate information in ways that lead to encoding more vivid sensory details into memory. Indeed, we found that RGFP966 controls memory induction for acoustic details of sound-to-reward learning. Rats treated with RGFP966 while learning to associate sound with reward had stronger memory and additional information encoded into memory for highly specific features of sounds associated with reward. Moreover, behavioral effects occurred with unusually specific plasticity in primary auditory cortex (A1). Class I HDAC inhibition appears to engage A1 plasticity that enables additional acoustic features to become encoded in memory. Thus, epigenetic mechanisms act to regulate sensory cortical plasticity, which offers an information processing mechanism for gating what and how much is encoded to produce exceptionally persistent and vivid memories.

Key words: auditory cortex; chromatin modification; cortical plasticity; epigenetics; histone acetylation; memory

Significance Statement

Here we provide evidence of an epigenetic mechanism for information processing. The study reveals that a class I HDAC inhibitor (Malvaez et al., 2013; Rumbaugh et al., 2015; RGFP966, chemical formula C21H19FN4O) alters the formation of auditory memory by enabling more acoustic information to become encoded into memory. Moreover, RGFP966 appears to affect cortical plasticity: the primary auditory cortex reorganized in a manner that was unusually “tuned-in” to the specific sound cues and acoustic features that were related to reward and subsequently remembered. We propose that HDACs control “informational capture” at a systems level for what and how much information is encoded by gating sensory cortical plasticity that underlies the sensory richness of newly formed memories.

Introduction

Histone modification is dynamically regulated during memory formation (Peixoto and Abel, 2013), yet the impact of such epigenetic mechanisms on information encoding has scarcely been addressed. Stefanko et al. (2009) introduced the idea that histone
modification induced by histone deacetylase (HDAC) inhibitors can transform a learning event that would not normally lead to memory into one that does. Moreover, these memories are in a form that persists beyond the point at which natural memory fails. Effects of HDAC inhibitors thus appear to act beyond memory enhancement, which would be expected to simply “boost” the strength of memory that would have naturally formed. Instead, these findings give rise to the following question: Do HDAC-inhibitors produce a dramatic change in information processing itself to alter what and how much information becomes encoded to produce persistent and vivid memory? If so, a much more significant role for HDAC enzymes exists in learning and memory than previously realized.

Here, we focused on the class I histone deacetylase inhibitor called RGFP966 (C21H19FN4O). Based on previously published recombinant purified protein assays in vitro, RGFP966 may have selectivity for HDAC3 over HDAC1 and HDAC2 (Malvaez et al., 2013). Class I HDACs include HDAC1, HDAC2, HDAC3, and HDAC8. HDAC3 is the most highly expressed class I HDAC in the brain and a key negative regulator of gene expression required for long-term memory formation (McQuown and Wood 2011; McQuown et al., 2011).

Previous evidence suggests a role for HDAC3 in consolidation effects, since application of the selective HDAC3 inhibitor used here (RGFP966) does not itself induce synaptogenesis, nor can it rescue memory impairment in an animal model of Alzheimer’s disease (Rumbaugh et al., 2015). Rather, HDAC3 appears to require a learning event to act on memory and promote synaptic plasticity (Rumbaugh et al., 2015). Indeed, the HDAC3 inhibitor RGFP966 has been shown to act on “subthreshold” experiences that normally would not have been remembered, transforming them into incredibly robust and persistent long-term memories (McQuown et al., 2011; Malvaez et al., 2013). Although several studies have examined the role of HDAC3 and other HDACs in memory processes, none have examined the role of HDACs in regulating information processing in the auditory cortex (AC), which allows for unique approaches to understand how HDACs regulate information being encoded into long-term memory.

This is the first study to address the function of class I HDACs in information encoding for auditory memory and learning-induced cortical plasticity. The primary auditory cortex (AI) is now appreciated for its functions in learning and memory. Plasticity in AI has become the most extensively studied of sensory areas for its function in associative memory (Weinberger, 2007; Scheich and Ohl, 2010; Pienkowski and Eggermont, 2011; Schreiner and Polley, 2014). For example, learning-induced tonotopic map expansion appears to enable both the strength of auditory memory formation and the specificity for what sounds and sound features are encoded into memory (Bakin and Weinberger 1990; Schulte et al., 2002; Bao et al., 2004; Rutkowski and Weinberger, 2005; Polley et al., 2006; Bieszczad and Weinberger 2010a, 2012; Lee and Middlebrooks, 2011), including the instantiation of sound-specific memory when tonotopic plasticity is artificially induced (Bieszczad et al., 2013; Froemke et al., 2013; Weinberger et al., 2013). Indeed, A1 plasticity might enable specific auditory features of a learning experience to become encoded for a persistent and vivid memory. We applied an auditory model to understand how an HDAC3-selective inhibitor could change information processing for what and how much becomes encoded into memory by engaging A1 plasticity.

Materials and Methods

**Animals**

A total of thirty-three adult male Sprague Dawley rats (275–350 g) were used in behavioral, electrophysiological, and/or molecular experiments. All animals were housed in an Association for Assessment and Accreditation of Laboratory Animal Care International accredited, temperature-controlled (22°C) facility on a 12 h light/dark cycle. Procedures were performed according to guidelines approved by the University of California Irvine’s Institutional Animal Care and Use Committee and in accordance with the National Institutes of Health Office of Laboratory Animal Welfare guidelines. An illustration of the experimental timeline and training procedure is provided in Figure 1.

**Behavioral training**

Water-restricted rats were trained to associate sounds with water reward (Fig. 1B, C). The protocol involved a task [developed by Bieszczad and Weinberger (2010a)] that required animals to press a bar manipulandum upon hearing a pure tone (Fig. 1B). Two sounds could be associated with reward. The first, Signal A (5.0 kHz, 70 dB SPL), was a pure tone sound that signaled the availability of reward upon the subject performing a bar-press (BP) response within 3 s of Signal A onset. The second, Signal B, was the sound of the reward delivery lever mechanism (with significant power in the 0.1 to 2.0 kHz sound frequency band; see Rutkowski and Weinberger, 2005). This signaled the reward’s immediate delivery. Acquisition was determined by performance level, calculated for each session throughout training as the percentage of bar presses within tone presentations that occurred only within the first 3 s of tone onset (rewarded BPs), scaled by the percentage of bar presses within a session that occurred at any time during tone presentations: performance = [(number rewarded BPs/number tone BPs) × (number tone BPs/number total BPs)] × 100%. All animals trained successfully learned to associate sound with reward (Fig. 1C).

**Behavioral memory test**

A memory test after training (Fig. 1B) revealed the frequency specificity of the auditory memory by determining behavioral responses to many frequencies, including that of Signal A (5.0 kHz) and Signal B (1.1 kHz). Bar-press response latencies to the onset of each test frequency (1.1, 2.4, 5.0, 10.6, and 22.4 kHz, each presented at 70 dB SPL) were determined in a single session of 200 unrewarded trials (40 trials of each test frequency, pseudorandomly intermixed). Response latency for each test tone was determined as the average latency from tone onset until a bar press for trials in which at least one bar press occurred. This latency was normalized within animal by its own average latency to press across all session trials with at least one bar press. These normalized values were used to construct a behavioral gradient to index memory specificity in the dimension of acoustic frequency for each animal, and then for the group. A distinct shorter latency response (see below, Determining behavioral frequency specificity, Individual analysis) was used as a proxy for greater frequency specificity.

**Drug administration**

The critical manipulation was pharmacological inhibition using a class I HDAC inhibitor called RGFP966. Immediately following the daily training session, a postraining systemic injection of either RGFP966 (10 mg/kg, s.c.) or vehicle (at a comparable volume to drug treatment) was delivered to each subject. The effective 10 mg/kg dose of RGFP966 (the uniquely selective HDAC3 inhibitor) was established previously in mice (Malvaez et al., 2013) and confirmed to penetrate the blood–brain barrier to rat auditory cortex with similar pharmacokinetics (Fig. 1A), which here establishes the rationale for a single-dose, single-inhibitor approach. Furthermore, the effects of postsession doses of RGFP966 for HDAC
inhibition are expected to act only on associative mechanisms of memory consolidation. Administration of RGFP966 (or vehicle) began after 4 d of initial sound-reward training to ensure all animals had initially acquired an auditory association with reward. This established an initial acquisition curve, which was also useful to determine performance-matched pairs of RGFP966- and vehicle-treated rats used in subsequent analysis of neural plasticity.

Distribution of RGFP966 in auditory cortex
Additional untrained animals (N = 9) were dosed with RGFP966 (10 mg/kg, s.c.) and killed at 30, 80, and 240 min after administration (N = 3 per time point). Brains were harvested and flash frozen. Samples from the auditory cortex were blotted and weighed. Water containing 0.1% formic acid at a ratio of 5:1 (v/w) was added into the auditory cortex tube. For example, 1000 μl of 0.1% formic acid was added to a 200 mg tissue. After tissue homogenization, samples were mixed with acetonitrile containing an internal standard to precipitate protein and were then centrifuged. Concentrations of RGFP966 and supernatants were determined by liquid chromatography mass spectrometry (API-4000 triple quadrupole (AB Sciex) equipped with a CTC Pal autosampler (Leap Technologies) and an Agilent 1100 high-performance liquid chromatograph). Samples were analyzed using the multiple reaction monitoring quantitation method, in which signals resulting from a specific parent–daughter ion transition, specifically 364.2–236.8 atomic mass unit for RGFP966, were measured after separation of the sample on an HPLC column (50 × 4.6 mm, 5 μm Gemini C18 column; Phenomenex) using a gradient of 0.1% formic acid in water and 0.1% formic acid in acetonitrile from 95:5 to 5:95 (v/v). Peak areas of RGFP966, normalized to an internal standard, were quantified from the regression equation (quadratic regression with 1/x weighting) obtained by quantification of standards prepared from stock solution of drug diluted in blank rat brain (Bioreclamation) and treated as described above for unknown samples. All analyses and calculations were performed using Analyst software (AB Sciex; Fig. 1A).

Electrophysiological recording
The tonotopy and borders of A1 were determined electrophysiologically in an acute “mapping” session performed at the termination of all training and testing from an anesthetized animal (sodium pentobarbital, 50 mg/kg, i.p.) that has been a standard preparation in cortical neurophysiological research (Fig. 1B). Combinations of frequency and intensity sounds (0.75–54.0 kHz in quarter-octave steps; 0–70 dB SPL in 10 dB steps; six pseudorandom repetitions) from a speaker at the entrance to the contralateral ear canal were used to evoke responses recorded from an array (1 × 4) of Parylene-coated tungsten electrodes to layers IV–V (400–600 μm depth; FHC; 1–2 MΩ). Electrodes were placed at locations that covered the surface of the auditory cortex, 250 μm apart at each
RGFP966 enables highly specific memory and A1 plasticity. A. Behavior. The latency to bar press for each tone frequency was determined in each group to indicate which frequencies best elicited reward-related behavior. Rats treated with RGFP966 (N = 6) were faster to respond than performance-matched rats treated with vehicle alone (N = 6) to frequencies associated with reward (Signal A) and reward delivery [Signal B; Z-score for paired difference in latency to respond to each test frequency; calculated as RGFP966 minus vehicle, responses to 1.1 kHz (Signal B) were on average 1.25 ± 0.75 significantly faster, p = 0.030; responses to 2.4 kHz were 2.62 ± 0.55 s faster, p = 0.280; responses to 5.0 kHz (Signal A) were 0.75 ± 0.33 significantly faster, p = 0.003; responses to 10.6 kHz were 0.75 ± 0.33 s faster, p = 0.380; responses to 22.4 kHz were 0.42 ± 0.5 s slower, p = 0.830], which indicates the formation of a more highly specific memory for acoustic frequency with RGFP966. Inset, Mean (±SE) bar-press latencies show group response gradients for the RGFP966 (N = 6) and vehicle (N = 6) groups, without respect to performance-matched pairs. B. Sound frequency representation in A1. Rats treated with RGFP966 had greater expansions of best frequency areas in A1 to overrepresent sound frequencies near the reward, Signal A (5.0 kHz; in the 4.0–6.3 kHz frequency band) and reward delivery signal, Signal B (1.1 kHz; in the 1.0–1.6 kHz frequency band). Asterisks indicate frequency bands that were significant with both paired (sign test) and unpaired (Mann–Whitney–Wilcoxon) one-sided tests after Holm–Bonferroni correction.

Analysis of A1

Evoked responses were averaged across stimulus repetitions to construct frequency response areas (FRAs) that reveal the CF, evoked threshold at CF (CF-threshold), BF (i.e., best tuning above threshold), and tuning BW (e.g., 20 dB SPL above threshold) for each A1 site. The representational area in A1 was measured using a Voronoi tessellation algorithm (based upon the voronoi algorithm in Matlab), which approximates two-dimensional areas in polygons labeled by CF (or BF) at each site that together delineate the cortical surface into regional areas of frequency representation. Areas were defined per isofrequency band, i.e., defined by the area contained by CFs or BFs within a half-octave range. Between-group Z-scores for differences from zero were used to contrast frequency specificity of A1 reorganization. Differences in the magnitude of A1 area and bandwidth changes were determined in performance-matched pairs of HDAC3 inhibition (HDAC3i)– versus vehicle-treated animals. The data shown in Figures 2 and 3 show the means of these differences between groups. Figure 3 uses a three-dimensional axis to combine the difference data presented separately for frequency and sound level in Figure 2 to give a comprehensive picture of the remodeled landscape of A1 sound representation. Tuning bandwidths were defined for each group per isofrequency band, i.e., defined by the bandwidth of averaged FRAs grouped by CFs within a half-octave range. Group mean bandwidth was determined with respect to threshold, at 10, 20, 30, and 40 dB SPL above CF threshold. Bandwidths were compared between groups at each level above threshold for each isofrequency pop-

Figure 2. RGFP966 enables highly specific memory and A1 plasticity. A. Behavior. The latency to bar press for each test tone frequency was determined in each group to indicate which frequencies best elicited reward-related behavior. Rats treated with RGFP966 (N = 6) were faster to respond than performance-matched rats treated with vehicle alone (N = 6) to frequencies associated with reward (Signal A) and reward delivery (Signal B; Z-score for paired difference in latency to respond to each test frequency; calculated as RGFP966 minus vehicle, responses to 1.1 kHz (Signal B) were on average 1.25 ± 0.75 significantly faster, p = 0.030; responses to 2.4 kHz were 2.62 ± 0.55 s faster, p = 0.280; responses to 5.0 kHz (Signal A) were 0.75 ± 0.33 significantly faster, p = 0.003; responses to 10.6 kHz were 0.75 ± 0.33 s faster, p = 0.380; responses to 22.4 kHz were 0.42 ± 0.5 s slower, p = 0.830), which indicates the formation of a more highly specific memory for acoustic frequency with RGFP966. Inset, Mean (±SE) bar-press latencies show group response gradients for the RGFP966 (N = 6) and vehicle (N = 6) groups, without respect to performance-matched pairs. B. Sound frequency representation in A1. Rats treated with RGFP966 had greater expansions of best frequency areas in A1 to overrepresent sound frequencies near the reward, Signal A (5.0 kHz; in the 4.0–6.3 kHz frequency band) and reward delivery signal, Signal B (1.1 kHz; in the 1.0–1.6 kHz frequency band). Asterisks indicate frequency bands that were significant with both paired (sign test) and unpaired (Mann–Whitney–Wilcoxon) one-sided tests after Holm–Bonferroni correction.
Figure 3. RGFP966 enables highly specific cortical remodeling that changes the landscape of sound representation in A1. A. Tonotopic plasticity in A1. Representative maps show sound frequency representation in A1 for a pair of performance-matched rats (+ VEH, treated with vehicle during training; + RGFP966, treated with RGFP966 during training). Maps were constructed using Voronoi tessellation algorithms that circumscribe and denote CF tuning by colored polygons (cool colors show low CFs, whereas warmer colors show higher CFs). Note the general progression of CFs from posterior to anterior sites across the cortical surface. Solid lines outline cortical areas that represent a quarter-octave range around the Signal A frequency. Likewise, stippled lines outline cortical areas that represent a quarter-octave range around Signal B. Ruler bar shows 1.0 mm across the AP and DV surfaces, as indicated. B. Signal-specific map reorganization. Cortical remodeling induced by RGFP966 in A1 enables expansions (shown in red shading) in the representation of reward-predicting auditory cues, here shown as the difference in the amount of percentage increase in A1 tonotopic representation across different frequency ranges (y-axis) at different sound levels (x-axis). Rats treated with RGFP966 show enhanced specificity of A1 reorganization relative to vehicle-treated rats (increases greater than vehicle controls indicated by red shading and decreases less than controls by blue). Maximal expansion in A1 representation occurs for the identity of specific reward signals, e.g., sound Signal A (5 kHz, 70 dB, shown by the solid circle) and Signal B (1–2 kHz, indicated by the thick solid line). Note that this figure combines and expands the data shown in Figure 2.

Immunofluorescence
Slides were fixed in 4% paraformaldehyde for 10 min at room temperature, blocked in 8% normal goat serum (Jackson Immunoresearch Laboratories) with 0.3% Triton X-100 in PBS, and incubated overnight at 4°C with primary antibody in block solution, followed by a 2 h incubation at room temperature with secondary antibody. Primary antibodies used were acetyl-histone-H4K8 (1:1000 dilution; Cell Signaling Technology), acetyl-histone-H3K14 (1:1200; Millipore), and acetyl-histone-H2BK12 (1:1000; Abcam). Adjacent sections were labeled using primary antibody against NeuN (1:100; Millipore). The secondary antibody was goat anti-rabbit IgG-FITC (1:1,000; Millipore Bioscience Research Reagents) or goat anti-mouse IgG-Cy3 (1:200; Millipore). All sections were counterstained with DAPI (1:25,000; Invitrogen). Immunolabeling was quantified by measuring the optical density from comparable 4 μm thick images from each animal using ImageJ software. The primary auditory cortex was defined by the area corresponding to area “Au1” as defined by Paxinos and Watson (1998) [AP, −6.5 to −3.5 mm; dorsoventral (DV), −3.38 to −5.60 mm] and by cell morphology and density using definitions for A1 and cortical layers as is typically described: a decreased cell density in layer V of primary cortex, where layer V is defined as ∼51–77% of the distance through the cortical thickness from the pia surface (Games and Winer, 1988; Weedman and Ryugo, 1996; Hefti and Smith, 2000). Hippocampus (HIPP) was defined by the area corresponding to the approximate location of CA1/CA2 (AP, −6.8 to −5.0 mm) in the same dorsal–ventral sections as were collected for AC analysis for valid comparison.
individual behavioral gradients, which made it possible, e.g., to detect memory for either Signal A or Signal B, or for both signals.

**Group analysis.** To determine the effects of HDAC3i on specific memory encoding, the individual normalized behavioral gradient for an animal treated with vehicle was subtracted from the normalized behavioral gradient of a performance-matched HDAC3i-treated animal (i.e., pairs of HDAC3i minus vehicle animals). There were six pairs of performance-matched animals (total, N = 12), whose difference gradients were averaged to find peaks in the mean difference gradient. A peak in the difference gradient was defined as described above for individual animal analyses to indicate which frequencies HDAC3i-treated animals had better remembered relative to vehicle-treated animals. Figure 2 (inset) also shows normalized behavioral gradients that were averaged within a group, without respect to performance-matched pairs (HDAC3i group, N = 6; vehicle group, N = 6), to show actual bar-press latencies for each group.

**Statistics.** A paired-sample t test or Z test and nonparametric sign test (for performance-matched paired data) or Mann–Whitney–Wilcoxon (for unpaired data) tests were used to determine significant differences in group data with α levels held at 0.05. Nonparametric statistics were used for small-sample behavioral analysis, as described in Results. Pearson’s correlation coefficient was used for brain versus behavior analyses. The Holm–Bonferroni method appropriately adjusted for multiple comparisons across sound frequencies. Group sizes were chosen to establish power at 0.80 (α = 0.05).

**Results**

**HDAC inhibitor RGFP966 alters the behavioral specificity of memory formation**

To test the hypothesis that HDAC inhibition has a key function in specific information encoding, we used the class I HDAC inhibitor called RGFP966. In a previous study, we established that RGFP966 crosses the blood–brain barrier with a brain/plasma ratio of 0.45 (Malvaez et al., 2013). In a substrate-dependent biochemical assay using purified recombinant HDACs, RGFP966 was found to be specific for HDAC3, with an IC50 value of 0.08 μM and no effective inhibition of any other HDAC at concentrations of up to 15 μM (Malvaez et al., 2013). Malvaez et al., (2013) found that a dose of 10 mg/kg resulted in concentrations above the IC50 value between 30 min and 1 h after delivery (yet 10-fold lower than the 15 μM concentration at which HDAC1 and HDAC2 may become engaged), indicating this dose is ideal for studying posttraining effects of RGFP966 on memory formation. Here, we confirmed that RGFP966 at a 10 mg/kg dose penetrates the blood–brain barrier into rat auditory cortex with typical pharmacokinetics, which together establish feasibility for the modulation of A1 plasticity due to action in the auditory cortex (Fig. 1A).

RGFP966 or vehicle was injected into rats learning to associate specific sound frequencies with reward in an auditory instrumental conditioning paradigm. There were two auditory signals associated with reward (Fig. 1B). The first sound, Signal A (a 5.0 kHz pure tone presented at 70 dB), signaled reward delivery contingent on a bar-press response. The second sound, Signal B (generated by a mechanical lever that delivered the reward through a port), signaled immediate availability of reward. Prior studies using this training protocol found that learning can produce strong memories and A1 plasticity for the 5.0 kHz pure tone, i.e., Signal A (Bieszcad and Weinberger, 2010a,b). Furthermore, HDAC inhibition by RGFP966 may promote acetylation during consolidation of auditory memory. Lysine 8 on histone 4 (H4K8) and lysine 14 on histone 3 (H3K14) are known targets of HDAC3, whereas lysine 12 on histone 2B (H2BK12) is not a known target. Immunofluorescence analyses did not reveal statistically significant increases in acetylation at two potential acetylation sites (1.2 ± 1.6% relative decrease of H3K14-Ac in primary auditory cortex, t test, p = 0.78; 3.1 ± 3.5% relative increase in CA1 of the hippocampus, t test, p = 0.19; 8.0 ± 6.6% relative decrease of H2BK12-Ac in A1, t test, p = 0.89; 2.7 ± 7.5% relative decrease in HIPP, t test, p = 0.64; N = 6 rats per group; N = 3 replicates per rat). However, there was evidence of tendency for increased acetylation at H4K8 in A1 (but not HIPP) in RGFP966-treated animals relative to vehicle-treated controls (4.9 ± 3.6% relative increase of H4K8-Ac in A1, t test, p = 0.09; 0.4 ± 4.3% increase in HIPP, t test, p = 0.46), suggesting that the locus of effect might be confined to a more specific cortical region within A1.

After 2 weeks of daily training and RGFP966 administration, we found that HDAC inhibition by RGFP966 did not affect acquisition or final performance level relative to vehicle-treated controls (Fig. 1C). However, a behavioral test for the sound specificity of the auditory memory formed revealed that the group
treated with RGFP966 remembered the signal sounds with higher specificity for acoustic frequency. Relative to a performance-matched vehicle-treated group, the behavioral gradient for frequency specificity at memory test showed that RGFP966 induced more specific memory for Signal A (Fig. 2A). In addition, we discovered that RGFP966 also permitted the incorporation of additional information for a second frequency, here called Signal B. In contrast, no vehicle-treated animals developed memory for both frequency Signals A and B. Therefore, RGFP966 enabled the formation of a more specific auditory memory as well as memory that incorporated information for not just one, but two sound frequencies that were associated with reward.

**RGFP966 alters signal-specific primary auditory cortical plasticity**

Evidence for the neural mechanism of increased behavioral specificity was found in A1. We discovered that RGFP966 enhanced the cortical representations of the two reward-related signal sounds. Indeed, the tonotopic areas of both the Signal A and Signal B frequencies increased in RGFP966-treated animals relative to vehicle-treated controls (Fig. 2B). Furthermore, in addition to a specific effect on sound frequency map expansion in A1, the RGFP966 effect extended to an additional acoustic parameter: sound loudness. This was detectable because Signal A was unique in its sound level during training, i.e., it was always presented at 70 dB. A1 neurophysiology was determined at multiple sound levels, including 70 dB, thereby enabling analysis of neural specificity for sound loudness. Remarkably, RGFP966 enhanced the expansion for the 5.0 kHz (Signal A) frequency only in its representation in A1 near the 70 dB SPL (Fig. 2C). The high signal specificity for both the frequency and loudness of Signal A was in contrast to Signal B, which was found to have a specific effect on A1 sound frequency representation, but a general effect on the representation of sound level. The specific frequency representation of Signal B (near 1.1 kHz) was significantly greater across all sound levels in A1, i.e., the effect generalized across sound loudness (Fig. 2C). A potential explanation for the lack of level-specific effects for Signal B is that it was not learned at a unique sound level during training. Indeed, the general effect on A1 level representation for Signal B need not be a failure of plasticity, but rather an induction of plasticity that accurately reflects the characteristic sensory features of Signal B (e.g., as a single specific frequency across a broad sound level). Together, these neurophysiological findings suggest that the effect of RGFP966 on A1 was to enable plasticity that accurately and specifically captured many acoustic characteristics of the cues associated with reward.

We next determined whether behavioral frequency specificity was linked to the magnitude of frequency-specific A1 area gain. Indeed, there was a significant positive brain–behavior correlation: the greater the A1 expansion, the more specific the auditory memory. A positive curvilinear relationship indicated that the stronger the specificity of signal frequency memory (revealed as the peak in behavioral response to Signal B at 1.1 kHz), the greater the specific frequency representation in A1 (Pearson’s correlation coefficient, $R^2 = 0.6775; p < 0.001$). Interestingly, this relationship was strongest at the level of CF, which describes A1 frequency representation at evoked threshold, rather than representation at BF ($R^2 = 0.3196; p = 0.055$), which predicted the observed nonsignificant correlation for the other, Signal A frequency ($R^2 = 0.2647; p = 0.44$), for which A1 had reorganized only at BF (i.e., near 70 dB). These data support the hypothesis that RGFP966 mediates a transformation of learning experiences into highly specific long-term memories via modulation of the sensory cortical plasticity of sound representation in A1.

Collectively, it appears that RGFP966 enables A1 remodeling in a multidimensional way to increase cortical representations of multiple sounds and sound features that are associated with reward (Fig. 3). To determine the extent of this apparent effect of RGFP966 to regulate A1 plasticity, we studied another form of representational plasticity in A1: breadth of frequency-tuning bandwidth. The different forms of representational plasticity (map vs BW changes) are important to distinguish since they may have distinct functional importance for the detection and discrimination of behaviorally relevant cues, respectively (Scheich et al., 2011). Thus, differences in receptive field bandwidth between RGFP966- and vehicle-treated animals were also examined. Here, we found that A1 sites with CF tuning to within half an octave around Signal A (5.0 kHz) had decreased breadth of tuning bandwidth with RGFP966 treatment (relative to vehicle controls). Signal B fell on the low-frequency edge of detection for calculating BW, so we were unable to determine whether similar decreases in bandwidth were present for the Signal B sound frequency (1.1 kHz). Nevertheless, the effect of RGFP966 on representation of Signal A was to sharpen A1 frequency tuning by as much as an octave (Fig. 4). We conclude from these data that RGFP966 affected cortical remodeling of (at least) two forms: signal-specific representational expansion and decreased frequency-tuning bandwidth.

**Discussion**

**HDAC inhibition enhances memory and cortical plasticity with added auditory detail**

Animals treated with RGFP966 after learning an association between sound and reward remembered more signals related to reward and remembered highly specific auditory features of those signals. Evidence for the role of RGFP966 to enhance the encoding of sensory information emerged in two ways. First, RGFP966 enabled memory that was stronger, as revealed by shortened latency behavioral responses to signal sounds. Second, RGFP966 induced memory that was more specific. Behavioral responses indicated memory for not just one signal sound (as in the vehicle-treated condition), but two signal sounds associated with reward. Furthermore, this effect of RGFP966 was not evident in acquisition or absolute levels of achieved performance in the task. Rather the effect of RGFP966 was on the specificity of auditory information encoded, which was revealed in a subsequent memory test for the frequency specificity of long-term memory. Thus, RGFP966 appears to alter information encoding (what and how much is encoded as a signal) without an effect on associative mechanisms (those which link the signal with reward) per se.

At the neural level, the effect of RGFP966 to alter frequency specificity was recapitulated in the apparent frequency-specific tonotopic plasticity for the remembered sound signals. Signals A and B both had enlarged cortical representations. Tonotopic plasticity was highly specific for the signals’ sound frequencies, and it was also highly specific for the signals’ sound levels. Future experiments are necessary to determine whether the specificity of A1 plasticity for sound level is behaviorally relevant. Learning-induced A1 plasticity can be highly specific for a variety of features of sound, including sound level (Polley et al., 2006), timing (Bao et al., 2004), and even location (Lee and Middlebrooks, 2011), among others (Scheich et al., 2011). Overall, the results suggest that HDAC inhibition opens the gates to neural processes enabling information encoding of highly specific sensory features of signals associated with reward. A general prediction ex-
tending from these findings is that HDACs might regulate what and how much information of any type will become encoded into memory.

**Informational capture by HDAC inhibition**

The current report of behavioral and neural effects on auditory memory formation and A1 plasticity suggest that HDACs can regulate information encoding. Indeed, sensory cortical plasticity and its epigenetic regulation suggest key roles for both mechanisms in the systems-level formation and consolidation of information-rich memory. An HDAC-mediated role in the specificity of auditory memory and A1 plasticity could exemplify a fundamental principle that explains reported effects of HDAC3 in hippocampus-dependent memory. Synthesis of prior findings with this report introduces a new hypothesis: HDACs regulate informational capture. Here we define informational capture as the transformation of a sensory "snapshot" from the immediate short-term memory of a current experience into a persistent long-term memory that retains the perceptual vividness of the original experience.

A general function of HDACs might be to regulate the scope of capture from perception and short-term memory into long-term memory. Thus, HDAC inhibitors would contribute to concurrently enhancing and inducing memory for multiple and highly specific sensory features of behaviorally relevant signals. Furthermore, we assign the predicted influence of HDACs to associative processes per se, as opposed to attention-related or perceptual processes active during the task (i.e., because RGFP966 is administered after training and therefore not present in the system during learning sessions). The effect of HDAC inhibition to enhance memory specificity could explain other reports showing long-term memory enabled by RGFP966 during subthreshold learning for object location (Malvaez et al., 2013, their supplemental Fig. 2). Having memory for these precise stimuli that are rich in detail and multisensory information would enable their correct recognition and discrimination at testing. Our interpretation of such data is that RGFP966 facilitates task performance by enabling memory to form and consolidate with exceptional sensory detail.

**Epigenetic mechanisms regulate information encoding**

Epigenetic regulation of neural plasticity in adult A1 opens questions of how chromatin modification mechanisms such as histone acetylation might act in the cortex as a whole to altogether contribute to information processing and depth of encoding in memory formation. Future experiments could reveal that memory for a specific frequency facilitates learning of a two-tone auditory discrimination in which one tone signals reward ("tone +" cue) and the other the unavailability of reward (a "tone −" cue). In this case, acquisition and achieved performance levels might also be facilitated with RGFP966 because specific information about frequency is necessary to perform correct frequency discriminations.

Although identifying molecular mechanisms was beyond the scope of the present study, that epigenetic regulation per se is key for controlling the specificity of learned information is supported by complementary evidence that DNA methylation can alter auditory discrimination behavior in rats (Day et al., 2013) and olfactory discrimination behavior in honeybees (Biergans et al., 2012). Subsequent analyses could determine the key genes regulated by a family of epigenetic players, including HDAC3 (considering that RGFP966 may preferentially affect HDAC3 activity), that ultimately affect the specificity of memory. The auditory model used here is unique in that it can be used to identify an epigenetic mechanism for memory formation at the level of cortical plasticity, which offers an approach in future studies to detect critical changes at an underlying level in gene expression profiles by targeting those neurons in A1 identified by shifts in receptive field properties within regions of cortical over-representation for reward-related sounds. Tools are becoming available to target the critical cell types, cortical columns, and layers of A1 tissue that have undergone plasticity, so future studies can begin to establish the molecular and genetic mechanisms for learning-induced sensory cortical plasticity that appears to be regulated by HDACs in the formation of sensory-specific memory. Such studies will be necessary to prove that HDAC3 is indeed key to information processing in the auditory cortex, as suggested by results generated from the use of the HDAC3 selective inhibitor RGFP966 in this study.

**Conclusions**

Overall, the findings show that an inhibitor selective for HDAC3 can be used to enable representational plasticity in sensory cortex that underlies memory for highly specific auditory information. As such, HDAC3 inhibitors could be key target drugs for altering the threshold of memory induction in the transformation from immediate perception to enduring memory. We introduce the hypothesis of “informational capture,” whereby currently available perceptual information is transformed by HDAC-dependent mechanisms from short- to long-term memory. It is possible that HDAC3 (or other HDACs of the class I type) regulates the amount of information entered into memory during initial memory encoding, memory consolidation, or both. The proposed function of these HDACs in sensory system plasticity and consolidation is particularly relevant to clinical conditions that suffer from failures in the specificity of encoded information. For example, combined behavioral and pharmacological approaches are provocative for their prospective ability to rescue or enable auditory learning, comprehension, or language acquisition, which demand auditory signal specificity to successfully attribute significance to particular sounds and their meaning. Benefit could be gained from behavioral therapies with concurrent use of class I HDAC-targeting drugs. We propose here that HDACs, and perhaps HDAC3 specifically, will gate memory for informative cues and their complex sensory features in any associative learning experience.

**References**


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