Associative learning and sensory neuroplasticity: how does it happen and what is it good for?

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Historically, the body’s sensory systems have been presumed to provide the brain with raw information about the external environment, which the brain must interpret to select a behavioral response. Consequently, studies of the neurobiology of learning and memory have focused on circuitry that interfaces between sensory inputs and behavioral outputs, such as the amygdala and cerebellum. However, evidence is accumulating that some forms of learning can in fact drive stimulus-specific changes very early in sensory systems, including not only primary sensory cortices but also precortical structures and even the peripheral sensory organs themselves. This review synthesizes evidence across sensory modalities to report emerging themes, including the systems’ flexibility to emphasize different aspects of a sensory stimulus depending on its predictive features and ability of different forms of learning to produce similar plasticity in sensory structures. Potential functions of this learning-induced neuroplasticity are discussed in relation to the challenges faced by sensory systems in changing environments, and evidence for absolute changes in sensory ability is considered. We also emphasize that this plasticity may serve important nonsensory functions, including balancing metabolic load, regulating attentional focus, and facilitating downstream neuroplasticity.

Sensory systems exist to transduce elements of the external world into neural representations of the world that are actionable for other parts of the nervous system. Most neurobiological accounts of sensation model this initial transformation as a fixed process, where the ear is like a microphone and the eye like a camera, each producing an initial neural representation that principally reflects the external stimulus. A given physical stimulus thus always produces roughly the same initial neural representation, presumably allowing it to be recognized quickly and interpreted appropriately. However, in practice the value of using a fixed mapping of physical stimulus onto initial neural representation may be limited. We easily recognize a familiar object seen from a new angle or a favorite song played over a noisy background, circumstances where the peripheral neural representations of the stimulus are actually very different from our previous experience. Outside the laboratory, situations where reoccurrences of a particular stimulus evoke the same pattern of initial neural activity each time are probably extremely rare. In this light, our sensory systems seem designed not to presume that a sensory stimulus will always produce the same neural representation but rather to expect the opposite—that a given stimulus in the world may produce quite different initial neural signals each time it is encountered. Consequently, the sensory system’s principal job may be to rapidly recognize ecologically important re-occurring stimuli across these diverse initial representations. This recognition requires that the sensory system act on prior knowledge about the relationships among stimuli in the world, which can only be the result of associative learning. Learning should thus be expected to be accompanied by sensory neuroplasticity, and these changes might need to be as early as the initial neural representation of the sensory stimulus.

The influence of learning on sensation has been appreciated at least since Aristotle’s claim that sensory systems change to become like their external stimuli (Aristotle and Shiffman 2011), notably including William James’s powerful assertion that “every perception is an acquired perception” (James 1890). However, in experimental practice changes derived from sensory plasticity (e.g., the tone sounds quieter) can be challenging to discriminate from changes related to task performance (e.g., the tone sounds the same but evokes the measured behavior less strongly) or stimulus generalization (e.g., the tone sounds quieter, but the subject makes the same behavioral response). This problem is acute for conventional laboratory learning paradigms that teach a subject that an originally neutral stimulus is predictive of threat or reward because any changes in the subject’s behavioral response to the stimulus confounds possible changes in perception with anticipation of the learned outcome. Studies of the neurobiology of learning and memory have thus focused on more behaviorally tractable questions, albeit with some notable exceptions. Perhaps as a consequence, “associative learning,” where brief training can induce rapid changes in behavior without obvious changes in sensory abilities (Fanselow and Poulos 2005; Weinberger 2007, 2015; Weinberger and Bieczcad 2011), is generally considered separately from “perceptual learning,” where extended training is used to induce often modest sensory changes (Gilbert et al. 2001; Brown et al. 2004; Hawkey et al. 2004; Polley et al. 2006; Gilbert and Sigman 2007). However, evidence is accumulating that typical associative learning paradigms (and especially fear conditioning and operant conditioning paradigms) can in fact drive stimulus-specific changes very early in sensory systems, including primary sensory cortices, precortical structures, and even the peripheral sensory organs themselves. Plasticity at such an early stage suggests that the distinction between associative and perceptual learning may thus be largely semantic. To consider the potential functions of the sensory neuroplasticity resulting from these learning paradigms, we must look both at the nature of the changes and their position within the circuit.
Instincts of associative learning-induced sensory plasticity

It has been known since the early 1980s that associative learning could induce physiological changes in sensory neurons in invertebrates like Aplysia and Hermisenda, where direct connections between sensory and motor systems permit simple physical stimuli to become able to evoke reflexive behaviors (Crow and Alkon 1980; Hawkins et al. 1983; Walters and Byrne 1983). However, associative learning has now been shown to induce sophisticated stimulus-specific neuroplasticity in the mammalian auditory, visual, olfactory, somatosensory, and gustatory systems, which are only indirectly related to behavior. These effects have been observed not only in secondary sensory “association” cortices (Sacco and Sacchetti 2010) but also in primary sensory cortices (Morris et al. 1998; Öh and Scheich 2005; Polley et al. 2007; Li et al. 2008; Chen et al. 2011; Gdalyahu et al. 2012; Suga 2012; Weinberger 2015), subcortical sensory structures (Edeline and Weinberger 1991a,b; 1992; Cruikshank et al. 1992; Kay and Laurent 1999; Gao and Suga 2000; Doucette et al. 2011; Fletcher 2012), and even primary sensory neurons (Jones et al. 2008; Kass et al. 2013d; Dias and Ressler 2014).

Associative plasticity in the auditory system

The first evidence of associative learning-related plasticity was reported in the auditory thalamus in 1956 (Galambos et al. 1956). This inspired an extensive series of investigations of learning-induced plasticity in the inferior colliculus, auditory thalamus, and auditory cortices (Öh and Scheich 2005; Scheich et al. 2011; Schreiner and Polley 2014; Weinberger 2015). In the auditory system, most individual neurons have receptive fields tuned to a single “best frequency” to which they are maximally sensitive and most strongly responsive. Within the cortex and some subcortical structures, neurons are distributed in a rough tonotopic fashion such that best frequencies are mapped onto the spatial location with a gradient from low-to-high frequency across the structure. Plasticity in the auditory system has thus historically been characterized both as changes in the receptive field of single neurons and as remapping of neural populations to alter the number of neurons preferring a given frequency.

The most straightforward studies of auditory plasticity induced by associative learning have relied on single neurons observed before and after conditioning. Early evidence of fear conditioning effects in single cortical neurons was reported in the 1980s (Kraus and Disterhoft 1982; Weinberger et al. 1984; Bakin and Weinberger 1990) when single-unit extracellular recordings of neurons in rodent area A1 often revealed dramatic shifts in their frequency tuning functions such that their best frequencies shifted in the direction of the shock-predictive tone (Fig. 1A). Follow-up experiments recording in the auditory cortex but using either auditory or visual stimuli for fear conditioning confirmed that only auditory-cued fear conditioning altered frequency coding in individual A1 neurons, thus ruling out the possibility that the observed plasticity merely indicated a change in arousal (Bakin et al. 1992), and showing that these effects persisted at least 8 wk after learning (Weinberger and Bakin 1998).

Analogous experiments trained rats to use sound intensity to localize a target location for reward and proved corresponding changes in the intensity–response functions of neurons in A1 (Polley et al. 2004), while training rats to use azimuth information (direction to the sound source) to perform a task sharpened the azimuth selectivity of A1 neurons (Zhang et al. 2013). These data demonstrate that although frequency tuning is a prominent organizational principle in the auditory system, auditory cortical plasticity permits flexibility in the representation of other stimulus characteristics such as intensity or azimuth.

Instances of associative learning-induced auditory plasticity

Changes in behavioral responses on a psychophysical detection task that indicate improved detection of a 4-kHz target stimulus at low stimulus intensities following pairing of the target stimulus with nucleus basalis (NB) stimulation. This is reflected in an increase in Hit rate at low stimulus intensities (left) and an improved d’ value (right). (E) Changes in behavioral responses on a psychophysical discrimination task indicating improved discrimination of the CS following narrowband stimulus (left) but not a control wideband stimulus pairing with nucleus basalis stimulation. (D,E are adapted from Froemke et al. 2013 and used with permission from Nature Publishing Group 2013.)
dimensions as well. They also show that both aversive and rewarding stimuli can evoke similar, though perhaps not identical, changes in the representation of associated auditory stimuli. Secondary auditory cortices are less well studied but likely exhibit similar learning-induced plasticity (Diamond and Weinberger 1984).

At the population level, pairing a neutral tone CS with shock or reward induces an expansion of the cortical subregions sensitive to the CS frequency (Fig. 1B,C; Rutkowski and Weinberger 2005; Bieszcza and Weinberger 2010a,b) or intensity (Polley et al. 2004, 2006). By varying the degree of water restriction across subjects, Rutkowski and Weinberger (2005) were able to alter the motivational value of a water reward predicted by a tone. They observed that the behaviorally inferred motivation of the rat to bar press for water reward strongly correlated \( r = 0.87 \) with the relative size of the area in A1 that responded to that tone’s frequency band, while nearby areas representing higher frequencies became proportionately smaller \( r = -0.72 \). If no further training occurs the naturally occurring expansion lasts for weeks, but extinction training, in which the subject learns that the CS no longer predicts the reward, reverses the cortical remapping (Bieszcza and Weinberger 2012). Similar work used classical conditioning in a PET scanner to train human subjects that high- or low-frequency tones predicted an aversive white noise burst and observed frequency-specific changes in auditory-evoked responses in the auditory cortex (Morris et al. 1998), though longer-term imaging experiments in humans have not been possible.

Auditory cortical remapping is now sufficiently well understood to permit some fascinating experiments. Polley et al. (2006) presented rats with a series of auditory cues that varied in both frequency and intensity. For some rats, the frequency variation guided the animal to the target location for reward while in other rats the intensity variation led to the target, but all rats received the same range of auditory stimulation. Remarkably, when the rats were trained to attend to the frequency of the sounds they exhibited a remapping of best frequencies in A1 but no change in local intensity–response functions in A1. Conversely, if they were trained to respond to the sound intensity their intensity–response functions changed but not the tonotopic mapping (Polley et al. 2006). This suggests a more sophisticated plasticity than can be produced by mere temporal conjunction. Indeed, auditory cortical plasticity can selectively enhance cortical responses to tones only when they occur in a particular sequence (Kilgard and Merzenich 2002) or at a particular repetition rate (Bao et al. 2004).

Area A1 is strongly innervated by cholinergic afferents, notably from the nucleus basalis (NB), which can induce strong local plasticity via muscarinic receptors (Ji et al. 2001; Miasnikov et al. 2001; Butt et al. 2009; Froemke et al. 2013). Electrically stimulating the NB during tone presentation alters the tuning of individual neurons (Froemke et al. 2013) and remaps cortex to selectively expand the region representing the corresponding tone (Dimyan and Weinberger 1999) or other acoustic features like frequency sweeps and pulse sequences (Kilgard and Merzenich 1998a,b; Weinberger 1998; Kilgard et al. 2001). Artificially inducing this remapping by pairing NB stimulation with tone presentation results in behavioral changes where rats begin to preferentially respond to the tone whose cortical representation was expanded (Bieszcza et al. 2013) and improve their detection of near or below threshold stimuli (Fig. 1D; Froemke et al. 2013). The circuit-level basis of these population changes is beginning to be understood, with neurophysiological data showing that shock-evoked acetylcholine release can stimulate layer I interneurons in auditory cortex that in turn inhibit inhibitory interneurons in layer II/III, ultimately disinhibiting cortical pyramidal neurons required for auditory fear learning (Letzkus et al. 2011). How this mechanism translates into persistent population-level remapping of the sensory region and how it interfaces with other neuromodulators like dopamine (Bao et al. 2001; Ilango et al. 2012) and with short-term synaptic dynamics (David et al. 2009) remains to be determined.

The experiments described thus far have all provided evidence of long-term re-mapping of the cortex, with effects that can last days or weeks. However, shorter term plasticity has also been observed. Training ferrets to engage in auditory-cued selective attention tasks can produce rapid changes in the spatio-temporal receptive fields of individual neurons in A1 that persist for hours (Fritz et al. 2003). Similarly, during performance of an auditory discrimination task these neurons can be seen to temporarily alter their adaptation properties to enhance the differences in response between relevant stimuli (Atiani et al. 2014; Shamma and Fritz 2014; Yin et al. 2014). It remains unclear whether these comparatively brief changes during auditory tasks reflect longer-term plasticity that allows the circuit to operate in different task-specific “modes,” or if this short-term response plasticity is mechanistically related to that seen with longer-term remapping at all.

In addition to the remapping of the auditory cortex, associative learning can alter the tuning of neurons in subcortical auditory processing regions, including the medial geniculate body (MGB) of the thalamus. Fear conditioning shifts the best frequency of guinea pig MGB neurons toward that of a threat-predictive auditory cue (Bakin and Weinberger 1990; Cruikshank et al. 1992; Edeline and Weinberger 1992), a shift that lasts for about an hour (Edeline and Weinberger 1991b). This associative plasticity upstream of the lateral nucleus of the amygdala has been noted as potentially playing an underappreciated role in auditory-cued fear learning (Weinberger 2011). Similar changes have been observed following conditioning in the inferior colliculus of the bat (Gao and Suga 2000). To our knowledge, associative learning-induced plasticity has not been tested for in more peripheral auditory structures.

**Associative plasticity in the olfactory system**

The olfactory system has played an important role in the study of the sensory effects of associative learning, because it is the only mammalian sensory system where the initial neural representations of external stimuli are experimentally accessible in vivo. Olfactory sensory transduction occurs in olfactory sensory neurons (OSNs) in the olfactory epithelium in the nasal passages. Each OSN expresses just one out of a repertoire of hundreds of odor receptors, and as their axons course back to the brain’s olfactory bulb they segregate so that all of the OSNs expressing each individual receptor type project to only one or two specific glomeruli on the surface of the olfactory bulb (Mombaerts et al. 1996; Potter et al. 2001). Visualizing the pattern of synaptic output from OSNs into each of these glomeruli in vivo thus reveals a spatial map where the input to each glomerulus indicates the aggregate activity of OSNs expressing the corresponding receptor. Consequently, the overall pattern of input across glomeruli reliably indicates the identity of the odorant in the nose (Wachowiak and Cohen 2001; Bozza et al. 2004; Soucy et al. 2009). Moreover, the initial convergence of “bottom-up” sensory input from OSNs in the nose with “top-down” projections from sensory cortical and other brain regions occurs in these glomeruli (and not in the thalamus), permitting the effects of descending neuromodulatory projections to be readily observed. Studies of cortical function and olfactory psychophysics have provided a rich complement to this unusually accessible information on early sensory processing (Li 2014).

In a seminal paper, Jones et al. (2008) used a line of genetically targeted mice where all of the OSNs expressing the M71 odor
receptor could be identified in histological sections of the olfactory epithelium to test the hypothesis that M71 receptor expression in the olfactory epithelium could be altered by extended fear conditioning. In these experiments, they found that fear conditioning these mice with an odorant known to activate M71-expressing neurons greatly increased the number of OSNs expressing the M71 receptor within the epithelium 3 wk later (Fig. 2A–C), while mere exposure to that odorant or fear conditioning with a different odorant that does not activate the M71 receptor had no effect. This constituted the first evidence that primary sensory neurons could be affected by associative learning in mammals. Subsequent work from this laboratory has not only replicated this finding but discovered that the descendants of these fear-conditioned mice also exhibited elevated numbers of M71 receptor-expressing OSNs and higher behavioral sensitivity to the M71-activating odorant (Dias and Ressler 2014). The epigenetic mechanism of this heritability is incompletely understood but appears to involve changes in DNA methylation in the gene encoding the M71 receptor.

This structural work has been complemented by neurophysiological evidence that the synaptic output of OSN populations exhibits associative, odor-specific enhancement in vivo. In mice expressing the fluorescent exocytosis indicator synaptopHluorin in all mature OSNs (Bozza et al. 2004), odorant-evoked neurotransmitter release from OSN presynaptic terminals was visualized both before and after discriminative olfactory fear conditioning in the same individual mice (Kass et al. 2013d). The behavior of OSN populations that responded selectively to the CS− odorant, which was presented but not paired with shock, was unchanged by 3 d of discriminative fear conditioning, but OSN populations that selectively responded to the CS+ odorant, which did predict an impending footshock, released much more neurotransmitter during CS+ presentations after fear conditioning than at preconditioning baseline (Fig. 2D,E). This facilitated increased the sensitivity of these OSNs across odor concentrations such that fear conditioning’s effect on the OSN output was comparable with that of a fourfold increase in odorant concentration in naïve animals (Fig. 2F). Surprisingly, OSN populations that responded to both the CS+ and CS− odorants (which shared some molecular features and thus both bound to some of the same odor receptors) were only potentiated when responding to the CS+, suggesting that somehow information about the significance of the global representation of the odorant is available even in these specific subpopulations of “primary” sensory neurons. The differential facilitation of single OSN populations when responding to two different odorants and the comparatively brief 3-d interval between the first odorant-shock pairing and the observed plasticity suggests that these physiological changes in OSN activity were not caused by structural changes in the numbers of OSNs. Similarly, facilitated responses have been observed using intrinsic signal imaging (which reflects some combination of presynaptic, postsynaptic, and glial cell metabolism) in olfactory bulb glomeruli

**Figure 2.** Learning-induced olfactory plasticity in olfactory sensory neurons. (A) Increased numbers of M71 receptor-expressing olfactory sensory neurons in the olfactory epithelium following fear conditioning with the M71-exciting odorant acetophenone (upper) compared with untrained controls (lower). (B) Increased number of axons from M71-expressing olfactory sensory neurons (blue) projecting to the corresponding dorsal and medial glomeruli on the surface of the olfactory bulb following fear conditioning with acetophenone (left) compared with untrained controls (right). (C) Increased area of the M71-associated glomerulus following odor-cued fear conditioning (upper) and odor-cued, cocaine-induced, conditioned place preference (lower). (A–C are adapted from Jones et al. 2008 and used with permission from the Society for Neuroscience 2008.) (D) Mice receiving fear conditioning show enhanced odor-evoked neurotransmitter release from OSNs activated by the shock-predictive (CS+) odor methyl valerate (MV) after fear conditioning (POST) compared with before fear conditioning (PRE). No change was observed for the shock nonpredictive (CS−) odor butyl acetate (BA) or the unexposed control odor isomyl acetate (IAA). RLIs indicate the resting light image of the olfactory bulb. (E) Summary of overall changes in odor-evoked neurotransmitter release across glomeruli and experimental groups. (F) Enhanced neurotransmitter release across concentrations of the CS+ odor, including those above and below the trained concentration (green bar), indicate a general increase in neural sensitivity to the CS+ odorant. (D–F are adapted from Kass et al. 2013d and used with permission from the American Association for the Advancement of Science 2013.)
following odor-cued operant conditioning (Abraham et al. 2014). These physiological changes thus appear to be evoked by both emotionally positive and negative associations and operate in parallel to the changes in OSN numbers observed by Jones et al. (2008) and Dias and Ressler (2014).

Given the plasticity in the initial sensory input to the brain, it is perhaps not surprising that effects of associative learning can be observed at later stages of the olfactory system as well. Within glomeruli, OSNs make synapses onto mitral cells, which serve as the principal output neurons from the olfactory bulb to the piriform cortex. Extracellular recordings from mitral cells in rats during the performance of an operant task have revealed that cellular activity is strongly influenced by learned expectations about whether an odor predicts a reward rather than the mere identity of the odor in the nose (Kay and Laurent 1999; Doucette et al. 2011). Olfactory fear conditioning selectively increases the odor-evoked activity of mitral cells receiving input from OSNs driven by the CS (Fletcher 2012). Similar behavior continues downstream, where olfactory fear conditioning alters the odor response patterns of single neurons in the anterior piriform cortex such that discriminative conditioning between odor mixtures narrows the response tuning of pyramidal neurons while simple conditioning of an odor broadened their tuning (Chen et al. 2011).

In humans, the physiological response to odors is not experimentally accessible in the OSNs or olfactory bulb, but functional magnetic resonance imaging (fMRI) can reveal odor-specific patterns of activity in the piriform cortex. In a fascinating experiment, Li et al. (2008) combined fMRI with psychophysical testing to observe the effects of fear conditioning on pairs of enantiomers, odors that are chemical mirror images of each other and thus usually smell very similar. Prior to aversive conditioning (the human analog of fear conditioning) the two enantiomers in each pair were indiscriminable to subjects and evoked very similar patterns of activity in the piriform cortex. Aversive conditioning was then performed using one odor from one pair of enantiomers as the CS+ and a second odor from a different pair of enantiomers as the CS−. After conditioning, the odor used as the CS+ not only evoked patterns of neural activity in piriform cortex that were now discriminable from its enantiomer counterpart, but the CS+ and its counterpart became perceptually discriminable to the subjects. No changes were observed for the pair used as the CS−. This shows some of the strongest evidence that associative learning induces sensory neuroplasticity in humans with corresponding effects on sensory abilities. Purely psychophysical experiments have also demonstrated that odor-cued aversive conditioning in humans can increase the subject’s sensitivity to the shock-predictive odorant (Åhs et al. 2013) and that this facilitation can last at least a few days (Parma et al. 2015).

**Figure 3.** Learning-induced sensory plasticity in the somatosensory and gustatory systems. Whisker-shock fear conditioning reduces the number of individual neurons in the barrel cortex that respond to the shock-predictive whisker movement compared with unpaired controls (A, B), but induces a substantial increase in the size of the whisker-evoked responses in the neurons that do respond (C). (A–C are adapted from Gdalyahu et al. 2012 and used with permission from Elsevier 2012.) (D) Three examples of individual neurons in the gustatory cortex (GC) whose tantant-evoked responses changed in magnitude and time course following conditioned taste aversion (CTA) learning. The tantast was presented at time zero, and the horizontal bar indicates the time period of statistically significant change. (E) Approximately 30% of recorded gustatory cortical neurons were changed by conditioned taste aversion learning, while few were changed by a control paradigm. (D,E are adapted from Grossman et al. 2008 and used with permission from the Society for Neuroscience 2008.)

**Associative plasticity in other sensory systems**

Though the most extensive work on associative learning-induced sensory plasticity has been done in the auditory and olfactory systems, mounting evidence suggests that it may be ubiquitous across sensory modalities. In the mouse somatosensory cortex, whisker manipulation typically evokes firing across a broad range of cortical neurons within the corresponding cortical barrel. Training the mouse that a particular whisker manipulation predicts an impending shock causes a notable sparsening of these responses, such that fewer neurons respond to the whisker stimulation (Fig. 3A, B) while those that continue to respond (Fig. 3C) exhibit much larger increases in activity (Gdalyahu et al. 2012).

In the rat gustatory cortex (GC), the tantant-evoked spiking behavior of ensembles of cortical neurons changes during conditioned taste aversion learning (Grossman et al. 2008; Moran and Katz 2014). In one study ~about 30% of the neurons altered the temporal features of their response to the CS+ tantast (Grossman et al. 2008). Interestingly, these cortical changes were principally observed later in the tantant-evoked response (i.e., after 1 sec; see Fig. 3D, E), while such plasticity was evoked throughout the response in simultaneous recordings from the basolateral amygdala (Grossman et al. 2008), suggesting an active shaping of gustatory response during memory retrieval. Neurons exhibiting plasticity in the GC during taste aversion learning partially revert to their original patterns during extinction learning (Moran and Katz 2014). Importantly, gustatory cortical neurons can also exhibit anticipatory responses preceding the unavoidable delivery of a pleasant or unpleasant tantast following classical conditioning (Gardner and Fontanini 2014). The circuit mechanisms underlying this physiological plasticity remain poorly understood, but cholinergic projections to the GC are necessary for the taste aversion learning to occur (Gutierrez et al. 2003; Parkes et al. 2014). As in the auditory system, there is growing evidence that the gustatory thalamus also exhibits associative plasticity and may mediate the learned responses to taste-predictive auditory cues in the GC (Samuelsen et al. 2013).
Visually cued fear conditioning in humans results in increased CS+ evoked BOLD signal not only in the amygdala but also in the extrastriate visual cortex (Knight et al. 1999; Morris et al. 2001). Steady-state visual evoked potentials in visual cortex are selectively amplified for the CS+ following visually cued aversive conditioning in humans (Song and Keil 2014). In rats trained to use visual stimuli to navigate to a hidden platform in a water maze, thalamocortical projections exhibited facilitated long-term potentiation and an increased visual evoked potential was observed in primary visual cortex compared with rats that were exposed to the same visual stimuli when they did not predict the location of the platform (Hager and Dringenberg 2010). In another interesting experiment, rats were fitted with goggles that presented monocular visual cues indicating the effective latency of water reward presentation in an operant task while the activity of single neurons in the deep layers of V1 was recorded (Shuler and Bear 2006). Early in conditioning this activity principally reflected the physical features of the stimulus, but after training many cortical neurons exhibited more complex patterns of activity (including both increase and decreases in firing rate) that persisted after visual stimulus offset and reflected the timing of the anticipated reward. Perhaps most remarkably, in the rat visual system simply associating an auditory cue with a visual cue (in the absence of any explicit reinforcement) is sufficient for the auditory cue to begin to evoke responses in primary visual cortex when the visual cue is later paired with footshock (Headley and Weinberger 2015).

Potential functions
Despite the robust physiological evidence of sensory plasticity and burgeoning understanding of its circuit-level mechanisms, we still have a limited understanding of how this plasticity is beneficial to the organism. Improved ability to detect stimuli with learned ecological importance would be obviously useful, but in most experiments such improvements are difficult to decouple from purely behavioral responses to the test stimuli (especially when using fear conditioning). Moreover, many potential changes in sensory processing need not actually result in a perceptual change—a tone could produce a larger neural response in A1 without necessarily being perceived as louder, for instance. However, even without outright changes in sensory abilities, sensory plasticity could be useful for directing attention, optimizing memory encoding and retrieval, metabolic savings, or other functions. Some of these functions could also be served by non-associative plasticity evoked by mere exposure to a particular sensory environment (Pienkowski and Eggermont 2009, 2010; Jakkamsetti et al. 2012; Kass et al. 2013b,c; Cadiou et al. 2014), and could potentially use overlapping mechanisms.

Sensory functions
A common question in interpreting these sensory plasticity studies is why a system capable of improved sensory function would not actually use that potential all the time or for all stimuli. The answer lies in the fact that sensory systems are constantly making tradeoffs. Increased sensitivity risks saturation should the stimulus become stronger, more rigorous discrimination of similar stimuli can impair recognition of the same stimulus on a noisier background, faster discrimination speed often comes at the expense of decreased accuracy, and so forth. Consequently, improvements in sensory function can be surprisingly difficult to define because “better” may depend on the organism’s goals or circumstances at that moment. Changes in the neural processing of sensory stimuli could of course reflect absolute improvement in detection or discrimination abilities (Recanzone et al. 1993; Li et al. 2008; Åhs et al. 2013; Froemke et al. 2013), but these changes may instead reflect retuning of the sensory system’s internal parameters to favor discrimination over generalization or sensitivity over dynamic range rather than an expansion in overall physiological abilities. Direct evidence of these tradeoffs after associative learning has not been experimentally accessible, but early auditory exposure to a pure tone that produces a similar expansion of the cortical representation of the corresponding frequency is accompanied by both impaired discrimination of that tone from others and improved discrimination among neighboring tones (Han et al. 2007). Perhaps because of these tradeoffs, there is a growing body of evidence that early neural processing of sensory inputs is actively shaped by task demands (Kay and Laurent 1999; Doucette et al. 2011) and expectations (Headley and Weinberger 2015; Czarnecki et al. in revision), reflecting a plastic, highly flexible processing system.

One of the most straightforward demonstrations of improved sensory function is increased sensitivity to the target stimulus. In a pair of recent odor-cued aversive conditioning studies in human subjects, Lundstrom and colleagues demonstrated a significant reduction in olfactory detection threshold that was specific to the shock-predictive odorant and could last at least several days (Åhs et al. 2013; Parma et al. 2015). Importantly, these findings included clear evidence that the increased sensitivity persisted even after the CS+ evoked skin conductance responses habituated, suggesting that the improved sensory performance was not being confounded by an anticipatory reaction to the shock. This study also found no change in the perceived intensity of the CS+ odorant, suggesting that higher brain regions somehow took the increased sensitivity into account in shaping the ultimate percept. This increased absolute sensitivity is consistent with the neurobiological evidence of increased numbers of CS+ responsive OSNs (Jones et al. 2008) and hyperresponsiveness of those OSNs to the CS+ odor (Kass et al. 2013d). Similar improvements in auditory detection have been observed following the induction of A1 plasticity (Fig. 1D) by pairing quiet tones with NB stimulation in rodents performing an auditory-cued operant task (Froemke et al. 2013).

In contrast to a detection task, the discrimination of a target stimulus from similar stimuli may be beneficial or not (Wisniewski et al. 2014), and the stimulus features that guide discrimination and generalization are seemingly determined by the challenges presented during the learning process (Guttman and Kalish 1956; Wisniewski et al. 2009). In effect, setting the correct balance between discrimination and generalization is the key to recognizing a familiar stimulus when encountered from a new perspective or background. Consequently, the mere fact that a stimulus representation has expanded to include more cortical area does not necessarily imply an improvement in discrimination (Talwar and Gerstein 2001; Ohl and Scheich 2005). The studies of Åhs et al. and Li et al. used an explicit discrimination paradigm, where more than one odor was presented during the conditioning session but only one was paired with shock. This did produce a new ability to discriminate between the shock-predictive odor and its enantiomer (even though that enantiomer did produce a new ability to discriminate between the shock-cueing odorant, suggesting that higher brain regions somehow took the increased sensitivity into account in shaping the ultimate percept. This increased absolute sensitivity is consistent with the neurobiological evidence of increased numbers of CS+ responsive OSNs (Jones et al. 2008) and hyperresponsiveness of those OSNs to the CS+ odor (Kass et al. 2013d). Similar improvements in auditory detection have been observed following the induction of A1 plasticity (Fig. 1D) by pairing quiet tones with NB stimulation in rodents performing an auditory-cued operant task (Froemke et al. 2013).

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conditioning that used a single odor during fear conditioning was found to broaden olfactory tuning curves in the piriform cortex, consistent with generalization rather than discrimination, while discriminative conditioning using multiple odors instead narrowed tuning curves, suggesting enhanced discrimination (Chen et al. 2011). This distinction makes ecological sense—in simple conditioning the CS shared many nonolfactory features with the test odor, including being an unnatural odor, being presented at high concentration, and being encountered exclusively in an experimental context, while in discriminative conditioning the inclusion of a CS—odor demonstrated that those nonolfactory features were less predictive of shock than the chemical identity of the odor itself. It remains unknown whether human subjects are equally sensitive to this distinction, but these results suggest that associative learning-induced sensory plasticity may enhance discrimination or reduce it depending on the circumstances.

Nonsensory functions
As our physiological methods become more capable of observing small neuronal networks (Gdalyahu et al. 2012; Kato et al. 2012; Moran and Katz 2014) and molecularly or anatomically defined cell types (Kass et al. 2013a; Wachowiak et al. 2013), it is growing increasingly clear that “neuroplasticity” can encompass a stunningly broad range of changes in neuronal spiking activity, synaptic connectivity, and neurochemistry, even in the same brain region at the same time. In sensory systems, this realization is reopening classical questions about whether and how specific patterns of neural activity relate to the features of the corresponding physical stimulus as opposed to the predictive utility of that stimulus, its behavioral implications, and other contextual or even cognitive factors (Shuler and Bear 2006; Kass et al. 2013d; Headley and Weinberger 2015). The key questions of sensory neuroplasticity consequently extend beyond the traditional “what new information has been encoded about the stimulus to envision not just changing neural representations but dynamic neural codes, where the same external stimulus is represented by different firing patterns or even different neurons depending on higher order features like potential meaning, the need for attentional and memory storage resources, and expectation or surprise. Such ideas bring with them homeostatic constraints, like the need to limit the brain’s metabolic load, and temporal constraints, where a perception achieved too late will not be actionable for the organism. Learning-induced changes in sensory systems may very well serve these goals rather than traditional sensory functions.

Neural computation is metabolically expensive, with action potential firing and especially synaptic signaling consuming the largest part of the brain’s energy budget, including ATP and oxygen consumption (Harris and Attwell 2012). Reductions in firing rate or in the number of neurons responding to a stimulus (Gdalyahu et al. 2012; Kass et al. 2013b) may thus reflect an optimization of stimulus representations according to metabolic constraints, while changes in spike timing or synchrony (Bao et al. 2004; Weinberger et al. 2013; Kay 2015) or increases in neural response to one stimulus balanced by decreased response to other stimuli (Froemke et al. 2013) present potentially metabolically neutral mechanisms for encoding stimulus importance. In this light the large increases in stimulus-evoked synaptic activity (Kass et al. 2013d) and action potential firing rates (Bakin and Weinberger 1990; Polley et al. 2004) after conditioning constitute an investment in significant resources that presumably serve an important purpose.

A critical function of early sensory systems is the allocation of limited attentional resources, as reflected in both neural processing and the selection of stimulus sampling behaviors. Some of the earliest work on classical conditioning-induced changes in sensory processing noted that even in an organism as simple as Aplysia, heightened sensory responses to a conditioned stimulus could facilitate attention and the selection of an ecologically appropriate response (Walters and Byrne 1983). In mammals, attentional systems exhibit an almost paradoxical system of preattentive processing, where many stimuli are superficially evaluated in parallel and across sensory modalities to select the next focus of attention for in-depth serial evaluation (Treisman 2006). The purpose of the usually heightened neural response to threat- or reward-predictive stimuli may not be to improve the sensory processing of these stimuli per se but to increase the likelihood that those stimuli attract top-down attention when they do occur. While selective attention has proved difficult to study in animal models, there is considerable recent evidence that aversive conditioning in humans does indeed produce an attentional bias such that threat-predictive stimuli tend to attract and hold attention (Croy et al. 2010; Felmingham et al. 2011). This bias improves pre-attentive processing by enhancing detection of masked target stimuli (Armony and Dolan 2002; Kleim et al. 2012), possibly through the conditioned facilitation of activity in sensory regions. Computational efforts to reconstruct the sensory stimulus from the activity it elicits in sensory cortex have demonstrated that changes in attentional and motivational state can notably influence the balance among sensory features in its neural representation (Mesgarani et al. 2009).

Finally, the usually increased neural response to threat-predictive stimuli may serve to facilitate learning and memory itself. The greater activity levels associated with threat- or reward-predictive stimuli are often similar to those evoked by higher intensity, more salient versions of the same sensory stimulus (e.g., the larger odor-evoked input to the brain after olfactory fear conditioning is similar to the input evoked by a fourfold stronger presentation of the same odor; see Kass et al. 2013d), which generally are learned about more quickly than weaker stimuli. Many forms of synaptic plasticity occur more strongly when presynaptic neurons are firing more quickly or more synchronously (Caporale and Dan 2008; Edeline 2012), suggesting that heightened sensory responses could directly increase the rate of learning about the stimulus in a virtuous cycle. Though the details may depend on the location and stimulation paradigm, there is evidence that some forms of sensory plasticity emerge and then revert, either over time (Edeline and Weinberger 1999) or with continued training on an associative learning task (Takahashi et al. 2011). This plasticity may thus facilitate the transfer of information about important stimuli to downstream regions (Jeanne et al. 2013) or potentially expand the range of local stimulus processing algorithms, of which the most effective subset are selected for retention (Takahashi et al. 2013). Following learning, the larger stimulus response could also facilitate the retrieval of stimulus-associated memories and selection of stimulus-appropriate behaviors (Znamenskiy and Zador 2013).

Conclusions
Associative learning can induce dramatic stimulus-specific plasticity in both the behavior of individual sensory neurons and the macroscopic remapping of sensory regions, including auditory, olfactory, somatosensory, gustatory, and visual systems. These effects are best understood in primary sensory cortices but also occur in pre cortical structures and even as early as the primary sensory neurons themselves. Mechanisms of this plasticity remain poorly understood, but clearly include a role for neuromodulators (notably acetylcholine) as a trigger to evoke plasticity during or after learning. The function of these forms of plasticity may be to improve detection or discrimination of critical stimuli or to
enable these stimuli to better engage attentional and learning circuitry.

In the auditory system, there has been considerable debate about the best way to conceptualize associative learning-induced cortical and subcortical plasticity. This debate has evolved along with our changing understanding of the auditory cortex’s function, initially emphasizing changes in frequency coding (Bakin and Weinberger 1990), expanding to emphasize spectrotemporal and spatial elements of the auditory stimuli (Polley et al. 2006; Fromke et al. 2013; Zhang et al. 2013; Schreiner and Polley 2014), and more recently emphasizing more cognitive models of cortical function like categorization of stimuli (Scheich et al. 2011) and dynamic, task-dependent reshaping of representations (Shamma and Fritz 2014; Yin et al. 2014). The more recent observations of learning-related plasticity in the olfactory, gustatory, and somatosensory systems have similarly been interpreted in light of the current understanding of each sensory structure’s purpose. The theme of this evolution is ultimately teleological: as we develop new ideas about what each system is for, we find corresponding explanations for why associative plasticity might be useful. However, regardless of the distinct neurobiological details associated with transducing and analyzing the physical stimulus for each modality, these diverse sensory systems share the challenge of detecting, recognizing, and cueing the organism’s response to a changing stimulus in a changing world. By comparing the effects of associative learning across sensory systems we may find thus insight not only into shared neuroplasticity motifs but also into the basic functions of the sensory systems themselves.

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References


Edeline JM. 2012. Beyond traditional approaches to understanding the functional role of neuromodulators in sensory cortices. Front Behav Neurosci 6: 45.


Weinberger NM, Miasnikov AA, Biesczad KM, Chen JC. 2013. \( \gamma \) Band plasticity in sensory cortex is a signature of the strongest memory rather than memory of the training stimulus. *Neurobiol Learn Mem* **104**:49–63.


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