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Aversive Learning Enhances Perceptual and Cortical Discrimination of Indiscriminable Odor Cues

Wen Li,1,* James D. Howard,1 Todd B. Parrish,1,2 Jay A. Gottfried1,3,4

Learning to associate sensory cues with threats is critical for minimizing aversive experience. The ecological benefit of associative learning relies on accurate perception of predictive cues, but how aversive learning enhances perceptual acuity of sensory signals, particularly in humans, is unclear. We combined multivariate functional magnetic resonance imaging with olfactory psychophysics to show that initially indistinguishable odor enantiomers (mirror-image molecules) become discriminable after aversive conditioning, paralleling the spatial divergence of ensemble activity patterns in primary olfactory (piriform) cortex. Our findings indicate that aversive learning induces piriform plasticity with corresponding gains in odor enantiomer discrimination, underscoring the capacity of fear conditioning to update perceptual representation of predictive cues, over and above its well-recognized role in the acquisition of conditioned responses. That completely indiscriminable sensations can be transformed into discriminable percepts further accentuates the potency of associative learning to enhance sensory cue perception and support adaptive behavior.

The ability to minimize contact with aversive experience is a hallmark of adaptive behavior. Via mechanisms of associative learning, organisms can use sensory information in the environment to predict impending danger and initiate flight-or-flight responses. The behavioral efficacy of associative learning thus hinges on sensitive and accurate perceptual evaluation of sensory signals. In particular, the ability to discriminate between biologically meaningful cues (e.g., smell of a 175-kg lion) and similar but irrelevant stimuli (e.g., smell of a 3-kg housecat) maximizes an organism's response sensitivity while minimizing hypervigilant and impulsive behaviors.

However, models of associative learning have traditionally focused on delineating the formation of associations between a sensory cue [the conditioned stimulus (CS)] and a biologically salient event [the unconditioned stimulus (US)] (1, 2), paying scant attention to perceptual changes in the CS itself. Several studies have considered how associative learning modifies cue-related tuning profiles in sensory cortex (3–8), although none has provided concomitant measures of sensory perception. As a consequence, direct links relating learning-induced changes in sensory cortex to perceptual gains in cue discrimination are unavailable, such that the functional importance of these neural effects on behavior remains poorly characterized. To the extent that conditioning can transform indiscriminable sensations into distinct percepts, such a mechanism would constitute a unique and potent means of optimizing adaptive behavior.

We combined functional magnetic resonance imaging (fMRI) with multivariate analytical techniques to explore the impact of aversive olfactory conditioning on perceptual and neural discrimination of predictive odor cues. The use of perceptually identical odor enantiomers (mirror-image molecules differing only in their chiral properties) enabled us to determine whether humans can acquire the ability to distinguish between odorous stimuli that initially smell the same. Twelve healthy human subjects (age range, 22 to 35 years; 8 female) were presented with four enantiomers (two different pairs), one of which (the target CS+; “tgCS+”) was repetitively paired with an electric shock (US) during a conditioning phase, whereas its chiral counterpart (“chCS+”) was not accompanied by shock (Fig. 1) (10). The second pair of odor enantiomers served as nonconditioned control stimuli (“CS−” and “chCS−”). The central prediction was that associative learning would enhance behavioral discrimination of related CS+ odorants, in parallel with reorganization of neural coding in human primary olfactory (piriform) cortex.

We first examined the behavioral effects of aversive conditioning on perceptual discrimination between the conditioned cue (tgCS+) and its related enantiomer (chCS+). We administered a triangular (triple-forced-choice) odor discrimination test (11, 12) to assess differences in perceived odor identity (i.e., the quality or character of a smell emanating from an odorous object). On each trial, subjects smelled sets of three bottles (two containing one odorant, the third containing its chiral opposite) and selected the odd stimulus. Before conditioning, discrimination accuracy was at chance (33%) for both CS+ and CS− enantiomer pairs, confirming that each pair was initially indiscriminable (Fig. 2A). After conditioning, behavioral accuracy for distinguishing between tgCS+ and chCS+ rose by more than a factor of 2, significantly exceeding both chance and pre-conditioning performance (Ps < 0.01; Wilcoxon test, two-tailed), without any improvement in distinguishing between CS− and chCS−. Subjective ratings of odor intensity, valence, or familiarity (11) did not vary across conditions (Ps > 0.4), ruling out confounds of the triangular test due to these extraneous variables and accentuating the change in perceived odor identity. Associative learning thus can enhance perceptual discriminability between initially indiscriminable odors, and these effects are specific for the CS+.

We next clarified the neural mechanisms underlying learning-induced perceptual enhancement of the predictive cue (11). Because neural representations of odor identity are maintained in posterior piriform cortex (12–14), and given the highly distributed spatial organization of afferent projections into the piriform region (15–17), we used multivariate fMRI (18, 19) to test the hypothesis that spatially distributed patterns of neural activity in piriform cortex evoked by tgCS+ and chCS+ would be reorganized as a consequence of associative learning (fig. S1). By extracting the raw fMRI signal intensity from every activated piriform voxel (fig. S2), we found that the spatial activity patterns in posterior piriform cortex strongly correlated for the CS+ pair (tgCS+; chCS+) and the control pair (CS−; chCS−) before odor-shock learning (Fig. 2B), corresponding to the high perceived similarity within each pair. However, after conditioning, these spatial correlations declined for the CS+ pair, particularly in...
comparison to the CS− pair (P < 0.05; Wilcoxon test, two-tailed) (Fig. 2B), in line with the learning-induced behavioral enhancement in odor discrimination between tgCS+ and chCS+. Additional analysis showed that within-pair correlations were significantly higher than across-pair correlations at preconditioning (P < 0.005, f = 0.66), which suggests that our multivariate technique has satisfactory discriminant validity for distinguishing between odor classes (II). Finally, the absence of respiratory differences across conditions (II) suggests that the imaging effects were not due to sniff-related confounds.

Condition-specific odor maps from one subject (Fig. 3) illustrate how spatial patterns of piriform activity were selectively reorganized from pre- to postlearning for the CS+ pair, whereas the patterns for the CS− pair remained highly coupled. Spatial “difference” maps (Fig. 3, right column) further show that there was minimal signal deviation (light-colored voxels) between CS− and chCS− at both pre- and postconditioning, whereas substantial pattern variation (dark-colored voxels) emerged betweentgCS+ and chCS+ postconditioning. Moreover, the response in any given piriform voxel could change in either direction (activation or deactivation); this exemplifies the sensitivity of multivariate pattern-based fMRI approaches to characterizing neural information in human sensory cortex (20).

The postlearning changes described above were paralleled by robust evidence for aversive conditioning. First, online physiological measurements (II) of the odor-evoked skin conductance response (SCR) revealed significant enhancement to tgCS+ at post- versus preconditioning when compared to the CS− odorants (Fig. 4A). These SCR changes, however, were not selective for the
tgCS+, because tgCS+ and chCS+ changes did not significantly differ ($P > 0.2$). In fact, there was a small but nonsignificant SCR increase to chCS+ relative to CS– odors ($P > 0.4$). Second, learning-induced changes in amygdala and orbitofrontal cortex (OFC) (analyzed using conventional fMRI approaches) (II) paralleled the SCR effects. A condition × time interaction (II) demonstrated progressive decreases in amygdala activity evoked by tgCS+ (versus CS– odors) as learning proceeded (Fig. 4B and table S1), consistent with prior studies of aversive learning that used visual CS+ stimuli (21, 22). Moreover, comparison of post- to preconditioning revealed increased mean responses to tgCS+ (versus CS– odors) in the OFC bilaterally (Fig. 4, C and D), another region implicated in associative learning (23, 24). Interestingly, fear conditioning partially generalized to the chCS+ odorant, which at a reduced threshold ($P < 0.005$ uncorrected) showed similar profiles in amygdala and OFC (table S2). These findings validate the efficacy of our paradigm to induce aversive olfactory learning, thereby supporting the idea that the perceptual and neural changes in sensory discriminability were a consequence of associative learning. The results further show that fear conditioning to an odor cue recruits many of the same regions involved in conditioning to visual and auditory cues, emphasizing the multimodal versatility of these learning networks.

The shock-dependent spatial modifications in posterior piriform cortex were seen in the absence of changes in the magnitude of mean activation. Although the conventional (univariate) fMRI analysis revealed increased mean activity in OFC (Fig. 4C), there was no evidence for similar mean changes in posterior piriform cortex, even at reduced threshold ($P < 0.01$ uncorrected). At the same time, fMRI multivariate (pattern) analysis of OFC (Fig. 2C, left) showed no evidence for enhanced spatial discrimination between the CS+ odors, but rather suggested further loss of coding specificity (more highly correlated patterns). In fact, the spatial correlation changes for the CS+ pair significantly differed between posterior piriform cortex and OFC ($P = 0.01$; Wilcoxon test), highlighting a regional specificity for the ensemble learning effect. This anatomical-functional double dissociation suggests that fear conditioning recruits functionally distinct networks acting in concert to maximize adaptive behavior: an emotion system (e.g., amygdala and OFC) optimized to detect threat signals with high sensitivity, and a perceptual system (e.g., posterior piriform cortex) optimized to encode signal specificity.

We considered that aversive conditioning could have heightened attention (or arousal) to tgCS+, evoking response changes in olfactory cortex. However, anterior piriform cortex, the purported target of human olfactory attention (25), was not modulated in response to associative learning, either in univariate (at $P < 0.01$ uncorrected) or multivariate (Fig. 2C, right) analysis. By comparison, the idea that aversive learning updates odor quality representations in posterior piriform cortex would closely accord with its role in coding odor identity in both animal (13, 26) and human (12, 14) models of olfactory processing. It is therefore unlikely that attention or arousal directly modulates odor coding in posterior piriform, although it remains possible that these mechanisms could mediate olfactory perceptual plasticity indirectly.

Aversive conditioning therefore has a direct influence on how perceptual information about a CS+ is updated in sensory-specific cortex, providing a potent neural substrate to guide the behavioral discrimination of predictive cues. The spatial reorganization of sensory coding in piriform cortex may reflect changes in olfactory receptive-field tuning, leading to improved perception of odor cues, such that unique or “tagged” piriform representations might gain privileged access to critical nodes underlying aversion-minimizing behaviors.

Knowing what to avert presents behavioral challenges that an organism must solve to survive. Prior work on fear conditioning has focused on how the CS comes to produce behavioral (i.e., conditioned) responses, rather than how conditioning alters sensory processing of the CS itself, resulting in perceptual learning and enhanced discrimination. We hypothesize that the substantial effect of emotional experience on perceptual processing in sensory cortices should have a vital impact on adaptive behavior and should thus be considered an indispensable component for models of learning and decision-making (4, 27, 28). Clinically, our data raise the intriguing possibility that neurobiological derangements in the ability to distinguish between salient cues and perceptually related inconsequential stimuli may underlie the emergence of anxiety disorders characterized by exaggerated sensory sensitivity and hypervigilance. This may provide a unique mechanistic framework for the development of new therapeutic interventions.

References and Notes
Excitatory Transmission

Sergiy Sylantyev,1 Leonid P. Savtchenko,1,2 Yin-Ping Niu,3 Anton I. Ivanov,4 Thomas P. Jensen,1,2 Dimitri M. Kullmann,3 Min-Yi Xiao,3 Dmitri A. Rusakov1

The synaptic response waveform, which determines signal integration properties in the brain, depends on the spatiotemporal profile of neurotransmitter in the synaptic cleft. Here, we show that electrophoretic interactions between AMPA receptor–mediated excitatory currents and negatively charged glutamate molecules accelerate the clearance of glutamate from the synaptic cleft, speeding up synaptic responses. This phenomenon is reversed upon depolarization and diminished when intracleft electric fields are weakened through a decrease in the AMPA receptor density. In contrast, the kinetics of receptor-mediated currents evoked by direct application of glutamate are voltage-independent, as are synaptic currents mediated by the electrically neutral neurotransmitter GABA. Voltage-dependent temporal tuning of excitatory synaptic responses may thus contribute to signal integration in neural circuits.

Although ion currents through postsynaptic AMPA receptors are small (~10^{-11} A), they can exert a lateral voltage gradient (electric field) of ~10^4 V/m inside the synaptic cleft (1, 2), which raises the possibility that they can affect the dwell time of electrically charged neurotransmitters (3). Does electrodiffusion, therefore, play any role in synaptic transmission?

The excitatory neurotransmitter glutamate is negatively charged at physiological pH (pK = 4.4), which implies that postsynaptic depolarization should, in principle, retard its escape from the synaptic cleft (Fig. 1A). AMPA receptor–mediated excitatory postsynaptic currents (AMPA EPSCs) decay more slowly at positive than at negative holding voltages in dentate basket cell dendritic AMPAR EPSCs in CA1 pyramidal cells (4) and in cerebellar granule cells (5). However, this has not been reported for AMPAR EPSCs generated at perisomatic synapses on CA1 or CA3 pyramidal cells (6–8). We evoked dendritic AMPAR EPSCs in CA1 pyramidal cells by stimulating Schaffer collaterals; the EPSC decay time τ (defined here as the area/peak ratio) increased monotonically with depolarization (Fig. 1B). The ratio between τ recorded at +40 mV and at −70 mV (τ_{+40}/τ_{-70}) was consistently above one [average = 2.17 ± 0.09, n = 49, P < 0.001; (fig. S1A)]. This asymmetry...