Spectrotemporal dynamics of auditory cortical synaptic receptive field plasticity

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The nervous system must dynamically represent sensory information in order for animals to perceive and operate within a complex, changing environment. Receptive field plasticity in the auditory cortex allows cortical networks to organize around salient features of the sensory environment during postnatal development, and then subsequently refine these representations depending on behavioral context later in life. Here we review the major features of auditory cortical receptive field plasticity in young and adult animals, focusing on modifications to frequency tuning of synaptic inputs. Alteration in the patterns of acoustic input, including sensory deprivation and tonal exposure, leads to rapid adjustments of excitatory and inhibitory strengths that collectively determine the suprathreshold tuning curves of cortical neurons. Long-term cortical plasticity also requires co-activation of subcortical neuromodulatory control nuclei such as the cholinergic nucleus basalis, particularly in adults. Regardless of developmental stage, regulation of inhibition seems to be a general mechanism by which changes in sensory experience and neuromodulatory state can remodel cortical receptive fields. We discuss recent findings suggesting that the microdynamics of synaptic receptive field plasticity unfold as a multi-phase set of distinct phenomena, initiated by disrupting the balance between excitation and inhibition, and eventually leading to wide-scale changes to many synapses throughout the cortex. These changes are coordinated to enhance the representations of newly-significant stimuli, possibly for improved signal processing and language learning in humans.

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1. Introduction

In the auditory system, neurons are tuned to various acoustic properties and parameters such as sound frequency, intensity, or repetition rate. The receptive fields and tuning preferences of auditory cells to these variables are not necessarily fixed, but can be changed depending on the forms of sensory experience, during neonatal development, throughout adulthood, and after hearing loss or other forms of auditory system pathology. As the entire receptive field of a particular neuron can be high dimensional and difficult or impossible to completely characterize in high detail, here we use shifts in excitatory and inhibitory frequency tuning in the rodent primary auditory cortex (AI) as a model for investigating the general phenomenology, mechanisms, and functional consequences of synaptic receptive field plasticity (i.e., modification of the tuning properties of synaptic inputs onto a sensory neuron).

Most previous studies of cortical receptive field organization and plasticity have relied on extracellular recordings of spike output or local field potentials. However, recent advances in understanding the organization and dynamics of cortical circuits have been obtained using intracellular techniques such as in vivo whole-cell voltage-clamp recording. With this method, it is possible to measure the excitatory and inhibitory currents activated by sensory stimulation, and use these data to compute tuning curves for tone-evoked synaptic conductances. While changes in tuning curves can be registered in terms of suprathreshold spiking activity, much of this plasticity seems to be due to adjustment of the often-subthreshold synaptic inputs that lead to spike generation; thus, examination of synaptic modifications is a natural level for investigation of cortical receptive field plasticity. Additionally, it is becoming increasingly apparent that inhibitory circuits are themselves plastic and strongly govern the modification of excitatory synapses. In many of these cases, inhibition is affected independently of excitation in a complex, dynamic manner. Therefore, in order to build predictive models and improve therapeutic treatments for auditory pathologies such as tinnitus or deafness, intracellular electrophysiological recordings are required to characterize
the changes to cortical networks after episodes of learning or injury, as this technique is currently the only available method for directly measuring inhibitory synaptic transmission.

In this review, we first describe the relation between synaptic inputs, spiking output, and the tonotopic organization of AI as a whole. We then discuss studies of changes to AI synapses that occur in response to peripheral damage both in vitro and in vivo. While hearing loss or manipulations of the sensory environment can also affect subcortical processing by upstream stations (Sanes and Constantine-Paton, 1983; Willott, 2005), and thus indirectly change the synaptic drive onto cortical neurons, a growing body of evidence indicates that modifications to cortical circuits are initiated earlier and endure for longer than changes elsewhere within the auditory pathway (Ma and Suga, 2005; Froemke et al., 2007). In the remainder of this review, we focus on developmental and adult synaptic receptive field modifications induced by patterned stimulation and sensory exposure. We hypothesize that, although there are important differences between critical period plasticity and adult plasticity, there may also be a conserved set of basic mechanisms for long-term reorganization of excitatory and inhibitory cortical inputs in the intact brain.

2. Synaptic and spiking AI frequency tuning curves

Much has been learned about the organization and plasticity of cortical networks from extracellular recording of neuronal action potentials. These spiking receptive fields are a complex function of synaptic inputs, intrinsic ion channel activity (especially the activation threshold for Na⁺ channels), and dendritic processing (Hirsch, 2003; Huberman et al., 2008; Nowak et al., 2010). Each of these components can be regulated and modified in ways that are still being experimentally determined (Losonczy et al., 2008; Feldman, 2009; Dorrn et al., 2010), making it challenging to predict how perturbations in the patterns of sensory experience lead to changes in neural circuitry. However, to a first approximation, the organization of suprathreshold spiking tuning curves is governed by the strengths and kinetics of excitatory and inhibitory synapses, at least in young-adult and adult animals (Monier et al., 2003; Wehr and Zador, 2003; Zhang et al., 2003; Dorrn et al., 2010). For this reason, we focus here on the synaptic basis of receptive field plasticity in terms of input strength, although undoubtedly other factors that influence postsynaptic integration—directly or indirectly—also play important roles in shaping the tuning properties of cortical neurons (Häusser and Mel, 2003).

Similar to other stations along the mammalian auditory pathway, AI neurons are generally tuned to sound frequency (Fig. 1A). While many neurons have a clear preference for pure tones of a specific frequency (the ‘best frequency’), the tuning widths, best frequencies, and overall response rates depend strongly on sound level. Neurons in adult rat AI, for example, can exhibit broad sub- and suprathreshold tuning at moderate to high intensities, potentially spanning much of the total cochlear frequency range (Sally and Kelly, 1988; Zhang et al., 2003; Metherate et al., 2005). Regardless of bandwidth, the spiking tuning curve of a neuron (Fig. 1A, left) is necessarily a subset of synaptic tuning (Fig. 1A, right).

Fig. 1. Spiking and synaptic frequency tuning curves of adult rat AI. A, Example current-clamp recording of spikes and excitatory postsynaptic potentials (EPSPs) from an AI neuron. Top, representative tone-evoked responses. Bottom left, spiking tuning curve of this neuron. Bottom right, excitatory synaptic tuning curve of this cell. B, Example voltage-clamp recording of excitatory and inhibitory postsynaptic currents (EPSCs and IPSCs) from a different adult rat AI neuron. Left, synaptic frequency tuning. Right, correlation between peak excitatory and inhibitory responses across tone frequencies.

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A major feature of synaptic receptive fields in adult cat, rat, and mouse, is that the relative strengths of excitatory and inhibitory responses to brief pure tones are proportional across tone frequency (Fig. 1B). Importantly, inhibition lags excitation by several milliseconds, allowing excitatory events to evoke one or more spikes with high temporal fidelity before being terminated by a corresponding degree of co-activated inhibition (Volkov and Galazjuk, 1991; Wehr and Zador, 2003; Tan and Wehr, 2009). This phase delay for inhibition is likely due to the architecture of rodent thalamocortical circuitry, in that there are few if any direct inhibitory projections to AI from the medial geniculate body (MGB), the lemniscal auditory thalamus (Winer, 1992). It remains to be determined whether this relationship holds for integration of synaptic inputs during ongoing sounds, especially with relatively shallow modulation rates, although recent compelling data from Zador and colleagues indicate that excitatory and inhibitory conductances seem to be co-regulated over time periods of milliseconds to seconds (Wehr and Zador, 2005; Asari and Zador, 2009).

It is also unclear what specific inputs and cell types contribute to these tone-evoked responses. There is a wide diversity of cortical GABAergic interneurons, with various firing properties, dendritic projection patterns, and dynamics of short-term plasticity (Petilla Interneuron Nomenclature Group, 2008). This suggests that different inhibitory cell types might be recruited by distinct patterns of sensory stimulation, contributing differentially to inhibitory responses as measured locally in the dendrites or globally at the soma. Likewise, it is also unknown to what degree thalamic or intracortical excitatory inputs contribute to net excitation evoked by tones or other stimuli. Cortical injections of muscimol, a GABA_A receptor agonist, effectively eliminate intracortical contributions to synaptic receptive fields, sparing thalamocortical input. This method was found to reduce the bandwidth of frequency-intensity receptive fields, but spared characteristic frequency responses (Kaur et al., 2004). Thus thalamic inputs may be relatively sharply tuned, while intracortical excitation broadens tuning curves and contributes preferentially to responses away from best frequency. In contrast, Liu et al. (2007) attempted to isolate thalamic inputs using muscimol in combination with a GABA_A receptor antagonist, to prevent reduction of presynaptic transmitter release at thalamocortical afferents (e.g., by activation of GABA_A receptors on thalamic synaptic terminals) while simultaneously reducing intracortical excitation. They found that tuning curve bandwidth was left intact, suggesting that thalamic input largely determines the overall extent of subthreshold frequency tuning curves (although this then leaves the question of the functional contributions of the extensive set of intracortical connections). Regardless of the anatomical basis of synaptic receptive fields, the relative connection strengths of thalamic and intracortical inputs can be changed by various forms of experience, with intracortical synapses seemingly expressing a higher degree of plasticity than thalamic inputs. After deafferentation of the peripheral sensory apparatus, either through whisker trimming (Diamond et al., 1994) or monocular deprivation (Trachtenberg et al., 2000), changes to cortical responses and receptive fields occur first in layers 2/3 and 5 before being detected in thalamocortical layer 4. Moreover, long-term modifications to tone-evoked synaptic responses in adult rat AI can be induced by pairing sensory stimulation with neuromodulatory release; this procedure affects intracortical but not thalamocortical inputs onto cortical neurons (Froemke et al., 2007).

3. Sensory deprivation modifies AI synapses

Experience-dependent changes to AI synaptic circuitry have principally been studied in two main ways: after sensory deprivation and in response to patterned stimulation. In each case, cortical synapses are most susceptible to manipulations or loss of sensory input during developmental critical periods (Katz and Shatz, 1996; Buonomano and Merzenich, 1998; Hensch, 2005). Auditory cortical critical periods usually last for a few days or weeks, beginning with hearing onset, and can occur at distinct ages for different receptive field properties or across cortical sectors (Chang et al., 2005; de Villers-Sidani et al., 2007; Razak and Fuzessery, 2007; de Villers-Sidani et al., 2007; Insanally et al., 2009; Sanes and Bao, 2009; Popescu and Polley, 2010). In humans, deafness during this time, even if ameliorated with devices such as hearing aids or cochlear implants later in life, can lead to profound impairments in speech and language comprehension (Eisenberg, 2007; Zeitler et al., 2008).

Developmental hearing loss, either sensorineural or conductive, has a number of effects on synaptic transmission throughout the auditory system (Takesian et al., 2009). Studies in brain slices have revealed that auditory neurons become more excitable after bilateral hearing loss (Fig. 2A). For example, when the cochlea is surgically ablated during early postnatal development, the mean strengths of excitatory synapses in the gerbil auditory midbrain and cortex are increased (Vale and Sanes, 2002; Kotak et al., 2005). Conversely, the amplitudes of inhibitory events are decreased in both cortex and inferior colliculus after deafening (Takesian et al., 2009), possibly due to a loss of presynaptic GABAergic terminals and a reduction in the number of postsynaptic GABA receptors (Sarro et al., 2008). These synergistic changes in excitation and inhibition after loss of afferent input presumably increase the overall excitability along the central auditory pathway, in a manner reminiscent of the synaptic adjustments described in the visual cortex after neonatal monocular deprivation (Maffei et al., 2004). Such plasticity might also account for the lower activation and perceptual thresholds for cochlear implant use in deafened animals (Snyder et al., 1990; Raggio and Schreiner, 1999), and could be related to the etiology of tinnitus after noise exposure (Eggermont and Roberts, 2004).

The effects of hearing loss have also been studied on shorter time scales. A technically-impressive recent study characterized the changes to excitatory and inhibitory inputs in young adult rat AI in vivo immediately after a brief episode of high-intensity noise exposure (Scholl and Wehr, 2008). Ten minutes of acoustic trauma with continuous tonal exposure at 110–120 dB sound pressure level (SPL) led to an increase in thresholds of auditory brainstem responses by almost 50 dB, indicating that significant hearing loss had been induced by this procedure. This induced a set of long-lasting synaptic modifications distributed across frequency tuning curves, as measured with whole-cell voltage–clamp recordings from AI neurons in anesthetized animals. Changes to tone-evoked excitatory and inhibitory synaptic strengths were rapidly expressed (within a few minutes) and endured for the duration of the recordings. These synaptic adjustments led to shifts in the best frequencies of excitation and inhibition, a disruption of excitatory–inhibitory balance (Fig. 2B), and prolonged the time course of membrane potential responses. As a consequence, synaptic frequency tuning curves became broader and the temporal precision of AI responses was degraded.

Although these changes were complex, Scholl and Wehr (2008) were able to identify several consistent features of synaptic receptive field modification induced by acoustic trauma. First, relative to the frequency of the traumatic tone, inhibitory responses evoked by lower frequency tones (2–3 octaves away) were reduced. However, within an octave or so of the traumatic tone frequency, inhibitory responses were greatly enhanced both for higher and lower frequency stimuli. Second, excitatory responses were only modestly reduced, or sometimes slightly enhanced in
peri-traumatic regions where inhibition was increased. Finally, although spontaneous firing rates were unaffected, previous subthreshold inputs could become suprathreshold, leading to a small but significant shift in best frequency by 1/3 octave and an overall broader spiking receptive field.

These changes in AI synaptic and spiking receptive fields are similar to the effects of deafferentation in the visual (Gilbert and Wiesel, 1992; Eysel et al., 1999) and somatosensory systems (Calford and Tweedale, 1988; He et al., 2004), in that peripheral damage leads almost immediately to network-wide reorganization of cortical receptive fields synaptic circuitry driven largely by disinhibition. It remains to be determined to what degree these changes are pathological or compensatory. Furthermore, it is unclear whether trauma- and deprivation-induced changes in cortical responses reflect modification of intracortical synapses themselves or are predominantly inherited from alterations of subcortical stations. However, the uncoupling between excitatory and inhibitory synaptic strength in vivo and in vitro indicates that, at least in part, some of these changes occur directly within the cortex.

4. Developmental plasticity of AI maps and receptive fields

In rodent AI, adult representations of sound frequency and intensity profoundly depend on the properties of the acoustic environment in early postnatal life. The critical period for AI frequency tuning begins immediately at hearing onset, but the duration and offset seem to be controlled by the patterns of sensory experience. Days of repetitive stimulation with pure tones of a given frequency or within a frequency range leads to an enlarged representation of those presented frequencies within characteristic frequency maps of young rats, but only when exposure occurs during the second postnatal week (de Villers-Sidani et al., 2007; Insanally et al., 2009). Conversely, unmodulated stimuli such as continual white noise (Chang and Merzenich, 2003) or continual pure tones (Zhou et al., 2008) prevent tonotopic organization from emerging in AI, apparently keeping the cortex in an unrefined yet still plastic state.

Developmental changes of cortical frequency tuning at the spiking level are paralleled by modulations of the underlying excitatory and inhibitory synaptic receptive fields. In rodent AI, synaptic maturation occurs between postnatal day (P) P12 to P21 (Oswald and Reyes, 2008; Domn et al., 2010). Excitatory inputs are tuned for sound frequency by approximately P14 (Fig. 3A), likely as a consequence of activity-dependent but experience-independent pre-patternning that occurs before the rodent auditory system becomes functional (Froemke and Jones, 2011). Inhibitory inputs are initially present but untuned at hearing onset, gradually becoming tuned and proportional in strength to excitation across the frequency range of rodent hearing (Domn et al., 2010). In this way, developmental sensory experience throughout the same period (P12–P21) leads to calibration of synaptic circuitry and formation of excitatory–inhibitory balance (Fig. 3B,C). As a consequence, the spiking output of individual AI cells is initially unreliable, imprecise, and at longer latency than in adults (Domn et al.,...
correlation between excitation and inhibition in young (P12 end of the second postnatal week, excitation and inhibition were uncorrelated. By the end of the third week, the correlation improved, and by the end of the first month, the correlation was similar to that measured in adult animals. Adapted from Dorrn et al. (2010).

...as cortical inhibitory circuits become shaped by activity and experience, spike timing precision and receptive field structure substantially increase. It is important to note that the overall amplitude of tone-evoked inhibitory responses in rat AI has been found to be approximately the same in young and older animals (Dorrn et al., 2010; Sun et al., 2010), in contrast to the gradual strengthening of inhibition that seems to occur in the rodent visual cortex (Hensch, 2005).

A fraction of neurons in young AI appear to have highly tuned and balanced excitatory and inhibitory synaptic receptive fields, equivalent to that observed in adult AI (Dorrn et al., 2010; Sun et al., 2010). It is currently unknown why some cells in young rat AI display unusually high correlation and co-tuning between excitatory and inhibitory frequency tuning. However, existing data suggest a few possibilities. For one, Sun et al. (2010) show that at threshold, excitatory and inhibitory tuning are mismatched by approximately one octave. However, while thresholds of AI neurons are higher during development than in adults, these thresholds are still considerably low (~30–45 dB SPL) throughout P12–P14, and at adult levels of ~20 dB SPL thereafter (de Villers-Sidani et al., 2007); these thresholds are far from the sound intensities of 70 dB SPL used in Dorrn et al. (2010) to assess excitatory–inhibitory balance in young AI.

A more likely explanation is that there are particular spatial regions of AI in which excitation and inhibition become precociously co-tuned. These cells may be localized specifically within cortical layer 4, which is specifically where Sun et al. (2010) concentrated their recordings, suggesting that development of cortical synaptic receptive fields might be heterochronic. In such a model, thalamoreceptive neurons and cell assemblies would mature first, before inhibitory inputs of downstream cells and networks successively become co-tuned to match the statistics of excitatory inputs. Alternatively, or in addition, it is possible that specific subregions of AI are pre-balanced by hearing onset, and that over the auditory cortical critical period, surrounding sectors are progressively integrated into AI. This may be analogous to (or even account for) the experimental reports that the electrophysiologically-defined AI tonotopic map begins to form around a central mid-to-high-frequency sector in P11 rats, and expands in size to include outlying regions during the second and third weeks of postnatal development (Zhang et al., 2001; de Villers-Sidani et al., 2007), despite the observation that thalamic innervation of overall AI is likely complete before hearing onset (Lund and Mustari, 1977). It is plausible that in these earliest well-tuned regions of the AI map, average excitatory–inhibitory balance is at mature levels, while in surrounding, poorly-tuned regions, excitatory–inhibitory balance is much lower.

Regardless of the specific receptive field properties of individual cells in developing AI, over a substantial population of recordings, excitatory and inhibitory frequency tuning is uncorrelated on average between P12 and P16. Excitation and inhibition progressively become balanced until reaching mature levels around P25–P30 (Fig. 3B). However, even in adult AI, excitatory–inhibitory balance is a statistical property of the cortex, with some cells having uncorrelated or anti-correlated synaptic frequency tuning (Fig. 3B). This is similar to results from the visual cortex, where in vivo intracellular recording studies have revealed untuned or cross-
tuned inhibitory inputs (Ferster, 1986; Douglas and Martin, 1991; Pei et al., 1991; Schummers et al., 2002; Monier et al., 2003; Sohya et al., 2007). Finally, we predict that in older animals, excitatory–inhibitory balance breaks down again, decreasing the temporal precision and spectral selectivity of AI neurons in the aged brain (Turner et al., 2005). However, rather than resulting from strong but untuned inhibitory circuitry (as in developing AI), this may instead be due to weakening of cortical inhibition and loss of GABAergic cells, including parvalbumin-positive interneurons (Caspary et al., 2008; de Villers-Sidani et al., 2010).

The maturation of inhibitory frequency tuning and excitatory–inhibitory balance in AI can be accelerated by certain types of developmental auditory experience (Dorrn et al., 2010). A few minutes of patterned stimulation with repetitive, pulsed pure tones of a given frequency modifies excitatory and inhibitory synapses, such that synaptic tuning curves rapidly shift toward the presented tone frequency (Fig. 4A). This form of receptive field plasticity is spectrally and temporally complex, seemingly involving coordinated changes orchestrated across multiple inputs over minutes to hours. First, excitatory and inhibitory responses evoked by tones of the presented frequency are potentiated (Fig. 4B, top). Second, enhancement at these inputs seems to spread to spectrally-proximal inputs within one octave of the presented frequency (Fig. 4B, top). Finally, responses to the original best frequencies of excitation and inhibition are suppressed (Fig. 4B, bottom). Collectively, these long-term synaptic modifications substantially increase the correlation between excitatory and inhibitory frequency tuning curves. Although induced by a few minutes of patterned tonal stimulation, synaptic modifications and enhanced excitatory–inhibitory balance are maintained for over an hour (Fig. 4C). Moreover, these changes can persist for much longer if additional patterned stimulation is provided, and prevent future episodes of patterned stimulation from inducing other changes to synaptic receptive fields (i.e., effectively bring the critical period for AI frequency tuning to a close). Importantly, the changes to inputs other than that presented during patterned stimulation were predominantly responsible for the increase to excitatory–inhibitory balance (Dorrn et al., 2010).

These synaptic changes induced by patterned stimulation seem somewhat similar to the set of modifications described by Scholl and Wehr (2008) after acoustic trauma, although the effects on spike generation are qualitatively different. Acoustic trauma was found to delay membrane potential responses, while patterned stimulation accelerates responses and improves spike timing precision at the presented frequency (Dorrn et al., 2010). Other types of disruptive or noxious stimuli also impair the development of cortical excitatory–inhibitory balance, including white noise stimulation (Chang et al., 2005; Dorrn et al., 2010) and perinatal exposure to environmental toxins (Kenet et al., 2007). In contrast to patterned tonal stimulation, continual or pulsed white noise stimulation for an equivalent exposure period did not improve the correlation between tone-evoked excitation and inhibition (Dorrn et al., 2010). Other forms of neonatal sensory exposure with different statistics have not yet been investigated at the synaptic level.

It is unclear what mechanisms contribute to the formation of cortical synaptic receptive fields and excitatory–inhibitory balance. In the young rat visual cortex, pairing visual stimulation with...
postsynaptic spiking strengthened sensory-evoked excitatory responses, in a manner that depended both on the pairing interval and postsynaptic Ca\(^{2+}\) influx (Meliza and Dan, 2006). This suggests that spike-timing-dependent synaptic modifications, similar to those characterized in vitro throughout the cortex and elsewhere in the central nervous system (Markram et al., 1997; Froemke and Dan, 2002; Tzounopoulos et al., 2004; Feldman, 2009), are induced by repetitive patterned stimulation in developing AI. However, given that spike timing is imprecise in young AI, synaptic plasticity induced by patterned stimulation may instead depend on local ‘hotspots’ of excitability where the excitation-inhibition ratio is particularly high. In each case, though, we predict that whenever sensory stimuli are paired with strong postsynaptic depolarization, NMDA receptors are activated, leading to increases in intracellular Ca\(^{2+}\) and subsequent long-term changes in synaptic strength.

Thus early in life, environmental factors including the patterns of acoustic experience control the strengths of excitatory and inhibitory synapses, which in turn govern the organization of receptive fields, the output of cortical circuitry, and the perception of auditory stimuli. The perceptual consequences of changes to sensory representations are not always obvious, and require more investigation by studies combining auditory exposure, behavioral sensitivity, and thalamocortical electrophysiology. In an important study, it was shown that young rats exposed to several days of patterned stimulation had enlarged cortical representations around the presented frequency, but these animals were impaired in perceptual discrimination of this over-represented frequency. By contrast, discrimination of nearby under-represented frequencies was substantially improved (Han et al., 2007). It remains to be determined how other forms of developmental experience affect sensory perception, and how the shapes and relative structures of tuning profiles are related to detection and discriminative abilities.

5. Neuromodulation and synaptic receptive field plasticity in adult AI

After the end of the critical period, passive sensory stimulation is generally insufficient for long-term synaptic modifications and persistent changes in the organization of AI receptive fields. Instead, adult cortical plasticity seems to depend more strongly on stimulus history and internal state variables such as arousal level and motivation. This behavioral context is often conveyed by activation of subcortical modulatory systems that directly project to AI, e.g., the cholinergic nucleus basalis (Rasmusson, 2000; Weinberger, 2007) or the noradrenergic locus coerules (Edeline et al., 2010). Whereas neuromodulation can influence development plasticity as well as adult plasticity (Bear and Singer, 1986), and neuromodulator antagonists also prevent induction of some forms of long-term synaptic plasticity in cortical slices (Choi et al., 2005), Acetylcholine plays a central role in arousal, selective attention, and modulation of cortical responses (Mesulam, 1998; Weinberger, 1998; Yan and Zhang, 2005; Disney et al., 2007; Froemke et al., 2007; Parikh et al., 2007; Herrero et al., 2008; Silver et al., 2008; Groard and Dan, 2009). Cholinergic modulation has a wide range of effects on cortical neurons, but a consistent observation is increased excitability (Woody and Gruen, 1987) and suppression of intracortical synaptic transmission (Xiang et al., 1998; Metherate et al., 2005; Sarter and Parikh, 2005), including release of GABA from cortical interneurons (Kruglikov and Rudy, 2008). Extracellular recording studies in vivo have shown that pairing pure tones of a specific frequency with electrical stimulation of nucleus basalis induces large, long-lasting enhancements of spontaneous and tone-evoked spiking (Bakin and Weinberger, 1996; Rasmusson and Dykes, 1988; Kilgard and Merzenich, 1998). Although electrical stimulation of nucleus basalis should activate a heterogeneous population of projection neurons, including those that release acetylcholine, glutamate, GABA, and various peptides (Henny and Jones, 2008; Lin and Nicolelis, 2008), pharmacological evidence indicates that cortical acetylcholine receptors are specifically required for the long-term effects on AI receptive fields of this pairing procedure (Bakin and Weinberger, 1996; Froemke et al., 2007).

Intracellular recordings in vivo have revealed the mechanisms by which stimulation of the nucleus basalis neuromodulatory system activates cortical networks (Metherate et al., 1992; Metherate and Ashe, 1993) and enables receptive field plasticity (Froemke et al., 2007). In these latter experiments, in vivo whole-cell voltage-clamp recordings were obtained from neurons in anesthetized adult rat AI (Fig. 5A), and excitatory and inhibitory synaptic frequency tuning profiles were initially measured (Fig. 5B,C). Afterward, tones of a specific non-preferred frequency were paired with electrical stimulation of nucleus basalis. Several seconds after the start of pairing, there was a large suppression of inhibitory events evoked by the paired tone, followed by a more gradual enhancement of tone-evoked excitation (Fig. 5B,D). These changes were long-lasting, persisting at least 20 min or more after the end of the pairing procedure (Fig. 5D–F). While nucleus basalis stimulation has immediate effects on both thalamocortical and intracortical transmission, longer-term synaptic modifications appear to be specific to intracortical connections and not to the primary thalamic input to AI (Metherate and Ashe, 1993; Froemke et al., 2007).

Due to the cooperative effects of suppression of inhibition and enhancement of excitation, nucleus basalis pairing disrupted excitatory–inhibitory balance in adult AI (Fig. 5B–E). Over a longer time period of several hours, however, synaptic modifications continually evolved, with inhibition progressively increasing to a higher level than before, eventually re-balancing the persistent increase of excitation at the paired frequency (Fig. 5F). These results indicate that the dynamics of inhibitory transmission could serve as a cortical memory trace of the relatively brief pairing episode (Froemke et al., 2007). The duration of input-selective disinhibition may permit self-re-organization of AI receptive fields, emphasizing the new preference for paired stimuli in a manner independent of further evoked neuromodulator release. Under natural conditions, this memory trace could represent sensory objects or events that have acquired new behavioral meaning, or might be similar to the sorts of cortical changes that occur during perceptual learning, especially for those tasks requiring focal attention and sensory discrimination. In this way, neuromodulatory systems allow cortical networks to selectively respond to important or novel stimuli, and appropriately update internal models of the external world.
Transient, focal suppression of inhibition may be a general mechanism for induction of receptive field modification in the adult cortex. During developmental critical periods, the high level of plasticity may be due to a less-refined inhibitory tone or imbalance between excitation and inhibition (Hensch, 2004; Chang et al., 2005; Dorrn et al., 2010), permissive for alterations of cortical networks by passive stimuli. In adult cortex, however, receptive field plasticity also requires activation of neuromodulator systems, reflecting the importance of behavioral context in associative learning and memory provided by subcortical systems (Weinberger, 2007). This is further demonstrated by a series of studies from Fritz and colleagues (Fritz et al., 2003, 2005), using single-unit recordings in AI of head-restrained behaving ferrets. Receptive fields of AI neurons were powerfully modified after behavioral conditioning. Excitatory and suppressive subregions of spectrotemporal receptive fields evoked by specific stimuli were altered when those stimuli were followed by tail-shock. The predominant changes to spectrotemporal receptive fields were increases of excitatory regions and reductions of suppressive regions around the conditioned tone (Fritz et al., 2003), strikingly similar to the synaptic effects of nucleus basalis pairing in anesthetized adult rats (Froemke et al., 2007).

Future work will be required to determine if developmental forms of synaptic receptive field plasticity also depend in some way on neuromodulatory systems.

6. A mechanistic model for long-term cortical synaptic receptive field plasticity

These recent data from in vivo intracellular recordings help connect previous extracellular experiments of receptive field plasticity to a large in vitro literature on the mechanisms of long-term synaptic plasticity. Taken together, these studies suggest a general model of plasticity by which changes to sensory experience affect and remodel cortical circuits. In particular, we hypothesize that there are three core phases to the microdynamics of cortical synaptic receptive field plasticity: an initial disinhibition...
and stimulus-selective enhancement of excitation, followed by network-wide reorganization of excitatory inputs, concluded by a protracted period of inhibitory plasticity in order to balance (or rebalance) excitation and inhibition (Fig. 6). We find evidence of this progression from recordings made in rat AI following noise-induced hearing loss (Scholl and Wehr, 2008), after pairing a tone with electrical stimulation of the cholinergic basal forebrain (Froemke et al., 2007), and in response to passive patterned stimulation early in life (Dorn et al., 2010). We emphasize that many of the details of this model remain to be determined, but it may prove useful as a framework or working hypothesis for the design of future experiments.

In this scheme, the initial factor that controls the induction of cortical plasticity is the mechanism of disinhibition. Inhibition has long been known to limit the induction of excitatory synaptic plasticity by preventing activation of NMDA receptors, subsequent postsynaptic Ca\textsuperscript{2+} influx, and the biochemical signal transduction pathways that lead to expression and consolidation of long-term synaptic modifications (Feldman, 2009). Artola et al. (1990) used different concentrations of the GABAA receptor antagonist bicuculline to demonstrate that, in slices of rat visual cortex, 50 Hz tetanic stimulation did not induce long-term changes in synaptic strength when inhibitory circuitry was intact, but induced long-term depression (LTD) for a low concentration of bicuculline and long-term potentiation (LTP) for a higher concentration. Thus the relative level of inhibitory control in cortical circuits can control the sign, magnitude, and presence or absence of synaptic modifications. However, synaptic receptive field plasticity is a complex process with several aspects beyond LTP of various excitatory synapses. What phenomena have disinhibitory effects in local cortical circuits to initiate synaptic modifications, and what other changes occur within cortical networks as a consequence?

In the adult auditory cortex, delayed and balanced inhibition allows tone-evoked excitation, dominated by AMPA receptor activation, to produce a transient spiking response (Wehr and Zador, 2003), but brief enough to prevent effective depolarization of NMDA receptors. This organization—perhaps the normative processing mode of adult cortex—allows sensory information to be rapidly computed without necessarily modifying the existing circuitry. Nevertheless, there may be many ways to effectively disinhibit adult cortical networks and/or activate NMDA receptors to induce LTP. Several different neuromodulators, including acetylcholine, seem to reduce GABAergic transmission in cortex (Froemke et al., 2007; Kruglikov and Rudy, 2008). We postulate that this neuromodulator-based disinhibition allows for remodeling of synaptic receptive fields, when paired with specific sensory stimuli (Fig. 6A, ‘Phase 1’). In some cases though, cholinergic activity or neuromodulation in general may not be required for adult cortical receptive field plasticity (Ramanathan et al., 2009). For example, repetitive pairing of a non-preferred stimulus with a preferred stimulus effectively shifts AI frequency tuning curves in anesthetized and awake ferrets, in a manner that depends on the precise timing between the two tones (Dahmen et al., 2008), analogously to spike-timing-dependent plasticity (Froemke et al., 2006; Meliza and Dan, 2006; Feldman, 2009). Here, the second of the two stimuli during the pairing procedure might overcome an inhibitory

![Fig. 6. Spatiotemporal dynamics of synaptic receptive field plasticity in rat AI neurons. A, Progression of synaptic modifications in adult rat AI after pairing electrical basal forebrain stimulation with tonal presentation. Initially, the strength of excitatory (‘Exc’) and inhibitory (‘Inh’) events are correlated across frequencies (‘Baseline’). During pairing, inhibition at the paired frequency is reduced in a manner that depends on cortical acetylcholine receptors, while NMDA receptor-dependent LTP is induced at excitatory inputs evoked by the paired frequency (‘Phase 1’). Green arrows represent changes in excitatory strength, while red arrows represent changes in inhibitory strength; size and dendritic locations of synaptic inputs are simply illustrative. Approximately 30 min after pairing, excitation and inhibition at the original best frequency are depressed, while inhibition at the paired input begins to recover (‘Phase 2’). Finally, inhibition at the paired frequency increases to a new, higher level to rebalance enhanced excitation (‘Phase 3’). B, Synaptic modifications in P12–P21 rats after patterned tonal stimulation. Before patterned stimulation, excitation and inhibition are imbalanced (‘Baseline’). At the presented input, both excitation and inhibition are enhanced for minutes to hours (‘Phase 1’). Potentiation seems to spread to inputs activated within one octave of the presented tone frequency (‘Phase 2a’), while the strengths of excitation and inhibition at their respective best frequencies is reduced (‘Phase 2b’). For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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threshold to provide sufficient depolarization for NMDA receptor activation. Alternatively, certain forms of repetitive stimulation might selectively reduce inhibitory transmission relative to excitation if the dynamics of short-term excitatory and inhibitory plasticity are distinct.

During development, however, the average total strengths of excitation and inhibition are approximately balanced overall (Sun et al., 2010), but are locally imbalanced. This suggests that there may be particular ‘hotspots’ in neonatal AI naturally sensitive to repetitive, patterned stimulation with pure tones (Dorrn et al., 2010), and these locations within the cortical network will be the first sites of modification during patterned tonal stimulation (Fig. 6B, ‘Phase 1’). In particular, if inhibition is globally balanced with excitation, but locally imbalanced, this would lead to particular spatial locations within AI in which the excitatory–inhibitory ratio is unusually high. A slight change in the amount of excitatory input, as during patterned stimulation, might then be sufficient to strongly depolarize these cells, gate NMDA receptors, and set in motion the mechanisms of long-term synaptic modification. Neurromodulation may be important for developmental plasticity, as lesions of the cholinergic and noradrenergic system together prevent ocular dominance shifts after monocular deprivation (Bear and Singer, 1986), and neurotransmitter agonists can gate short-term dependency in slices of rodent visual cortex (Seol et al., 2007). Changes to the inputs or firing patterns of modulatory centers might be an important step in closing developmental critical periods, along with expression of molecules such as the cholinergic prototoxin Lynx1 that act as regulatory elements over adult cortical plasticity (Morishita et al., 2010).

During both adult and developmental forms of synaptic receptive field plasticity in rat AI, excitatory synapses tuned to the repetitively-presented stimulus are enhanced within approximately 1 min (Froemke et al., 2007; Dorrn et al., 2010). This form of LTP enabled by cholinergic modulation is highly stimulus specific in adult AI, but is less selective in young AI; nearby stimuli within one octave are also enhanced after patterned stimulation (Dorrn et al., 2010). There is precedent for spreading of LTP to neighboring synapses within tens of microns (Fig. 6B, ‘Phase 2a’), possibly by release of extracellular or intracellular messengers (Engert and Bonhoeffer, 1997; Harvey and Svoboda, 2007). Another possibility is that in the developing nervous system, there is intrinsically less stimulus specificity in the set of afferents activated by patterned stimulation, such that inputs within an octave of the presented stimulus are also reliably engaged by repetitive tonic exposure.

Changes to the presented input alone may not be sufficient to allow cortical networks to differentially respond to new or updated sensory information. Therefore, modifications to other inputs may also be important to take advantage of changes to cortical representations. One consistent observation in studies of receptive field plasticity is a reduction in the responses to the original best stimuli (Bakin and Weinberger, 1996; Froemke et al., 2007; Dahmen et al., 2008; Scholl and Wehr, 2008; Dorrn et al., 2010). We have found that, regardless of position within the tuning curve, these initially-preferred inputs are selectively depressed over tens of minutes, both in adult (Fig. 6A, ‘Phase 2’) and developing rat AI (Fig. 6B, ‘Phase 2b’), and this process is considerably slower than the expression of LTP at the paired or repetitively-presented stimulus (Froemke et al., 2007; Dorrn et al., 2010). As a number of theoretical studies have emphasized the importance of tuning curve shape for information processing (Pouget et al., 1999; Zhang and Sejnowski, 1999), these coordinated positive and negative changes in excitatory inputs might not only help conserve net excitation, but also preserve the general structure of synaptic receptive fields, shifting the peak rather than distorting or flattening cortical tuning curves. While this LTD of the original best stimulus could be related to the types of homeostatic synaptic scaling documented in the visual cortex (Desai et al., 2002), we think that instead, given the high degree of stimulus specificity and relatively fast dynamics, original best stimulus depression may be a form of heterosynaptic LTD (Scanziani et al., 1996; Royer and Paré, 2003). The mechanisms of best stimulus depression remain to be determined, especially the mechanisms by which AI neurons and cell assemblies are able to identify and selectively downregulate their particular local maxima. We note that, although the relative excitatory–inhibitory ratio may play a predominant role in AI plasticity (Froemke et al., 2007) and ocular dominance plasticity in developing visual cortex (Hensch, 2005; Southwell et al., 2010), these canonical examples of cortical reorganization are fundamentally different. In particular, shifts of AI frequency tuning begin with enhancement of a weaker input followed by a delayed reduction of the original best input; conversely, ocular dominance shifts are initiated by suppression of the originally preferred, deprived input followed by strengthening of the weaker, spared input (Frenkel and Bear, 2004). Rather than reflecting a basic difference between the visual and auditory systems, these opposite synaptic dynamics are probably related to the precise manipulation of sensory input in each case: monocular deprivation leads to a shift away from the deprived input, while repetitive tonal exposure leads to a shift toward the over-represented stimulus.

Finally, after excitatory tuning curves have shifted to prefer the paired or repetitively-presented stimuli, inhibitory inputs are adjusted in proportion to excitation (Fig. 6A, ‘Phase 3’). This balancing of synaptic circuitry seems to unfold over minutes to hours, and in adult AI, does not occur in absence of sensory experience (Froemke et al., 2007), indicating that specific patterns of activity are required to guide reorganization of cortical microcircuitry. After nucleus basalis pairing, the direct cholinergic suppression of tone-evoked inhibition at the paired input is converted into an intermediate-term depression that lasts roughly 10–30 min, before progressively increasing to match the rapid increase of excitation evoked by the same stimulus. Similar orchestration of excitatory LTP and inhibitory LTD has been previously described in vitro (Lu et al., 2000; Ivenshitz and Segal, 2006), but little is known about this process in vivo, or how this inhibitory depression is then transformed into an enduring potentiation. Upregulation of BDNF release (Huang et al., 1999) or activity-dependent transcription factors such as Npas4 (Lin et al., 2008), which increases GABA receptor expression after postsynaptic increases in intracellular Ca2+ concentration, are two likely candidates important for balancing excitatory and inhibitory inputs. An important avenue for future research will be to determine the set points for excitatory–inhibitory balance, and how inhibitory circuitry is locally calibrated with high precision across subregions of synaptic receptive fields.

7. Conclusion

Intracellular recordings in vivo have been essential for describing the dynamics of modifications to cortical microcircuitry at the synaptic level. During development, perturbations in the sensory environment drive changes in synaptic strength, organizing cortical receptive fields around the statistics of sensory inputs. In the adult brain, receptive field plasticity is controlled by behavioral context and motivational state, acting through neuromodulators such as acetylcholine and noradrenaline to gate long-term changes in excitatory and inhibitory synaptic receptive fields, perhaps through a common disinhibitory and/or NMDA receptor signaling pathway. In each case, an extensive set of positive and negative adjustments are coordinated across multiple synaptic
inputs, to update cortical representations of the external world and ensure that excitatory inputs are balanced by a proportional amount of inhibition.

It remains unclear how distinct elements of cortical networks and subcortical neuromodulatory systems are recruited by various forms of sensation, experience, and internal drive for the control of synaptic modifications, circuit dynamics, perception, and cognition. It will be important to determine the contributions to AI synaptic receptive fields not only from local intracortical connections and the MGB, but also from higher cortical areas including prefrontal cortex. In addition, recordings from subcortical nuclei in awake animals will be necessary to understand which feedback and modulatory inputs are activated under different behavioral contexts. A number of different neuromodulators have disinhibitory effects (Kruglikov and Rudy, 2008), but others — particularly noradrenalin (Edeline et al., 2010) and dopamine (Bao et al., 2001) — may act in other ways or on other elements of cortical networks, including glial cells and vascular processes. Modifications to non-neuronal aspects of the nervous system might be crucial for consolidating changes to cortical circuitry or ensuring that such changes are organized over considerable distances.

Careful analysis of the dynamics of synaptic receptive field modifications will be critical for understanding the key mechanisms and putative behavioral consequences of cortical plasticity. Chronic disruption of excitatory–inhibitory balance is also postulated to play a role in neuropathological conditions such as epilepsy and autism spectrum disorders, as well as hearing loss, tinnitus, and language impairments (Rubenstein and Merzenich, 2003). The ability to selectively increase or decrease specific inputs, through some combination of behavioral training, pharmacological approaches, and more invasive techniques (e.g., electrical stimulation or cell transplantation), provides a powerful means to potentially remediate a large number of nervous system disorders and improve cognitive functions, given an appreciation of the diverse mechanisms engaged during cortical remodeling.

References


Iversen, S.D., Segal, P.P., 2008. Simultaneous NMDA-dependent long-term potentiation of EPSCs and long-term depression of IPSCs in cultured rat hippocampal neurons. Journal of Neuroscience 26, 1199–1210.


