Noradrenergic Activation Amplifies Bottom-Up and Top-Down Signal-to-Noise Ratios in Sensory Thalamus

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Thalamocortical cells receive sensory signals via primary sensory afferents and cortical signals via corticothalamic afferents. These signals are influenced by a variety of neuromodulators that are released in the thalamus during specific behavioral states. Hence, different neuromodulators may set different thalamic modes of sensory information processing. We found that noradrenergic activation affects sensory and corticothalamic signals in the whisker thalamus differently than cholinergic activation. Whereas cholinergic activation increases the spontaneous firing (noise) and enlarges the receptive fields of ventroposterior medial thalamus (VPM) cells, noradrenergic activation decreases spontaneous firing and focuses receptive fields. Consequently, for sensory signals, noradrenergic activation sets bottom-up thalamic processing to a focused and noise-free excitatory receptive field, which contrasts with the broad and noisy excitatory receptive field characteristic of cholinergic activation. For corticothalamic signals, noradrenergic activation sets top-down processing to a noise-free high-frequency signal detection mode, whereas cholinergic activation produces a noisy broadband signal detection mode. The effects of noradrenergic activation on signal-to-noise ratios of VPM cells were found to be mediated by nucleus reticularis thalamic (nRt) cells. Hence, a major role of nRt cells is to regulate the noise level of thalamocortical cells during sensory processing. In conclusion, different modulators establish distinct modes of bottom-up and top-down information processing in the sensory thalamus.

Key words: thalamus; vibrissas; sensory processing; vigilance; attention; locus ceruleus; brainstem reticular formation; acetylcholine; norepinephrine; corticothalamic

Introduction

Thalamocortical cells are at the center of a neuronal network that involves sensory, cortical, and modulatory inputs (Sherman and Guillery, 1996). In the ventroposterior medial thalamus (VPM), thalamocortical cells form clusters, called barreloids (Land et al., 1995), that project to clusters of cells in layer IV of somatosensory cortex, called barrels (Woolsey and Van der Loos, 1970). VPM cells receive signals from four main sources. Primary sensory fibers originating from clusters of cells, called barrelelets (Henderson and Jacquin, 1995), in the principal trigeminal nucleus provide sensory (bottom-up) signals. Corticothalamic fibers originating in layer VI of barrel cortex (Bourassa et al., 1995) provide top-down influences. Fibers from nucleus reticularis thalamic (nRt) cells (Ohara and Lieberman, 1985) provide the main inhibitory control in VPM. Finally, a variety of neuromodulator fibers originate mainly in several brainstem nuclei. Among them, cholinergic fibers derive from laterodorsal tegmentum (LDT) or pedunculopontine (PPT) nuclei (Satoh and Fibiger, 1986; Hallanger et al., 1987), and noradrenergic fibers originate in the locus ceruleus (LC) (Lindvall et al., 1974; Simpson et al., 1997).

Early studies in several sensory systems recognized the impact of arousal and behavioral state on the responsiveness and activity of thalamic neurons to sensory inputs (Poggio and Mountcastle, 1963; Livingstone and Hubel, 1981; Swadlow and Weyand, 1985) (for review, see Castro-Alamancos, 2004b). These dynamic changes are primarily caused by neuromodulators released in thalamus during specific behavioral states (Vanderwolf, 1988; Steriade and McCarley, 1990; Jones, 1993). In particular, cholinergic neurons in the LDT/PPT complex discharge vigorously during paradoxical sleep and also during wakefulness (el Mansari et al., 1989; Steriade et al., 1990), and the levels of acetylcholine increase in the thalamus during those states (Williams et al., 1994). Noradrenergic neurons in the LC discharge robustly during high levels of vigilance and attention, reduce their firing during slow-wave sleep, and stop firing during paradoxical sleep (Hobson et al., 1975; Foote et al., 1980; Aston-Jones and Bloom, 1981; Aston-Jones et al., 1991), and their firing produces forebrain activation (Berridge and Foote, 1991). Because both systems, cholinergic and noradrenergic, are active during arousal and induce forebrain activation (Moruzzi and Magoun, 1949), it is particularly relevant to know whether they have similar or, instead, selective effects on thalamocortical information processing.

Many studies have investigated the effects of electrical stimulation of cholinergic (LDT/PPT) and noradrenergic (LC) brainstem nuclei on thalamocortical network responses. However, these effects are difficult to interpret because these nuclei are intricately interconnected between each other and project to ar-
as that also innervate the thalamus. Thus, LC stimulation may influence cholinergic cells in LDT/PPT or other thalamus-projecting cells. A different approach to study the selective effects of these neuromodulators is to infuse them directly into the thalamocortical network.

In the vibrissa system of rodents, cholinergic activation enlarges the receptive fields of VPM cells (Aguilar and Castro-Alamancos, 2005) and enhances their responses to high-frequency sensory signals, virtually eliminating rapid sensory adaptation in the sensory thalamus (Castro-Alamancos, 2002a). However, the effects of noradrenergic activation on sensory responses of VPM cells are unknown. Moreover, the effects of either cholinergic or noradrenergic activation on the responses of VPM cells to cortical signals are also unknown. In the visual system, application of norepinephrine to lateral geniculate nucleus cells via iontophoresis enhances their firing and responsiveness to optic nerve electrical stimuli (Rogawski and Aghajanian, 1980a,b) and affects responses to visual stimuli (Funke et al., 1993).

In the present study, we compared the effects of cholinergic and noradrenergic thalamic activation on the responses of VPM cells to sensory and cortical signals in urethane-anesthetized rats. The results show that these modulators have highly selective effects that set different modes of sensory and corticothalamic information processing.

Materials and Methods

Surgery. Sixty adult Sprague Dawley rats (300–350 g) were used in this study and cared for in accordance with National Institutes of Health guidelines for laboratory animal welfare. All experiments were approved by the Drexel University Institutional Animal Care and Use Committee. Rats were anesthetized with urethane (1.5 g/kg, i.p.) and placed in a stereotaxic frame. All skin incisions and frame contacts with the skin were injected with lidocaine (2%). A unilateral craniotomy extended over a large area of the parietal cortex. Small incisions were made in the dura as necessary. Body temperature was automatically maintained constant with a heating pad at 37°C. The level of anesthesia was monitored with a hand probe. Each of the six whisker stimulators were controlled with the hand probe. Each of the six whisker stimulators were controlled with a computer-programmable Master-8 (A.M.P.I., Jerusalem, Israel).

Whisker and corticothalamic stimulation was delivered according to the following protocols. For whisker stimulation, a trial consisted of an initial 2 s without whisker stimulation, followed by stimulation delivered to each whisker at 2 s intervals (the order of whisker stimulation was randomly selected). The first whisker was stimulated 2 s after the trial began, the second whisker was stimulated 4 s after the trial started, and so on, so that the sixth (last) whisker stimulus was delivered 12 s after the start of the trial. Thus, a single trial contained stimuli for all six whiskers and lasted a total of 14 s. Whisker stimulators consisted of several stimuli delivered at 2 Hz or 10 stimuli delivered at 20 or 20 Hz. To ordinary, only four stimuli were used at Hz because little more adaptation is produced in this frequency with additional stimuli. For high-frequency whisker stimulation (e.g., 10 Hz) analyses, the last stimulus in the train was used.

For corticothalamic stimulation, a trial consisted of 3 s of no stimulation, followed by a 10 pulse train delivered at 2.5, 10, 20, and 40 Hz. Each trial lasted 11 s. Each peristimulus time histogram (PSTH) was created with a minimum of 20 trials for corticothalamic stimulation and 30 trials for whisker stimulation using a 1 ms bin size.

Microdialysis. To apply drugs into VPM, a microdialysis cannula (250 μm diameter, 2-mm-long membrane) was placed around the following coordinates (in mm): posterior, 3; lateral, 2–3; depth, 4–6 as described previously (Castro-Alamancos, 2002a). The cannula entered into the brain at an angle (~30°) from the midline, whereas the VPM recording electrode entered lateral to the cannula and parallel to the midline. Artificial CSF (ACSF) was continuously infused through the probe at 2–4 μl/min. ACSF contained the following (in μM): 126 NaCl, 3 KCl, 1.25 NaH2PO4, 26 NaHCO3, 1.3 MgSO4, 7H2O, 10 dextrose, and 1 CaCl2 2H2O. Carbachol was dissolved in the ACSF at 0.5–1 mM. Norepinephrine was dissolved at 0.5–1 μM and protected from light and from oxidation with ascorbic acid. Atropine and phenolamine were used at 1–2 μM. Bicuculline methiodide (BMI) and CGP35348 (p-3-aminopropylp-dieothoxymethyl phosphoric acid) (CGP) were used at 100–300 μM and 5–10 μM, respectively. The diffusion of the drugs using microdialysis was determined by placing the probe and an array of four microelectrodes at distances from the probe. We found that norepinephrine (1 μM) and carbachol (0.5 μM) affected multiunit activity at distances of 0.5–1 μM but not at 1.5 μM or above from the probe. This indicates that the effects of the drugs are confined to a 1 mm area around the probe. This spread includes the VPM and nRt but may also affect parts of adjacent structures, such as the posterior nucleus and the zona incerta, both of which respond to whisker stimulation (Bartho et al., 2002; Trageser and Keller, 2004).

Although potential effects mediated by these structures should be considered, we believe that these higher-order nuclei would be unlikely to explain most effects reported in the present paper.

Data analysis. Spontaneous cell firing was computed by counting the number of spikes during the 2–3 s period at the beginning of each trial and for a minimum of 20–30 trials. Population data are presented as mean ± SD. Statistics for comparing population data between quiescent (control) and cholinergic or noradrenergic states consisted of paired t tests if the populations were normal according to the Shapiro-Wilk normality test. Nonparametric tests consisted of Wilcoxon’s signed-rank test for within-group comparisons and the Mann–Whitney test for between groups comparisons. Note that each cell is studied in at least two different states (n refers to number of cells).

Histology. At the end of the experiments, the animals were given an overdose of sodium pentobarbital and each perfused through the heart.
with saline followed by paraformaldehyde (4%) or the brain was directly extracted and placed in the fixative. The brains were then sectioned in the coronal plane using a vibratome (80–100 μm) and processed for Nissl staining. For the cells included in the study, subsequent analysis confirmed the location of electrode tracts within VPM and nRt.

**Results**

**Dataset**

VPM cells were identified by their short latency to a whisker deflection (<7 ms) and by tracing the track of the recording electrode in histological sections confirming that the electrode was inside VPM. The data shown in the present study are part of a growing dataset consisting of >200 well isolated single-unit thalamic recordings. Because the amplitude of these spikes is generally in the range of 5–10 mV, we presume that the electrode tip is very close to the membrane of the cell. Generally, these recordings are fairly stable so that the cells can be studied for several hours. All of the cells presented here are naïve cells, unless otherwise indicated. That is, drug applications were done to one cell per animal.

**Effect of noradrenergic activation on whisker-evoked responses in VPM cells**

To produce noradrenergic activation in the thalamus, we infused norepinephrine (0.5–1 mM) using microdialysis, as described previously (Castro-Alamancos, 2002a). To produce cholinergic activation, we infused carbachol (0.5–1 mM). In these experiments, up to six whisker stimulators were used simultaneously to map the responses of the PW and several AWs. An example of the effects of carbachol and norepinephrine on the same VPM cell is shown in Figure 1. During the quiescent (control) state, the spontaneous firing rate of this cell was 0.5 Hz, whereas during carbachol, the firing rate increased to 6 Hz, and during norepinephrine the cell had no spontaneous firing. PSTHs of responses to low-frequency stimulation (0.1 Hz) of each of five individual whiskers revealed that, during the quiescent state, the cell responded primarily to two of the stimulated whiskers, the PW and one AW. However, during carbachol, the cell responded to stimulation of all five whiskers, albeit with different latencies and strengths. In contrast, during norepinephrine, the cell responded mostly to two of the five whiskers, with some responses present also for two other whiskers. Although cells responded exceedingly well to whisker stimuli during either carbachol or norepinephrine, during norepinephrine, the only activity present was that evoked by the whisker stimulus.

Figure 2A shows population data of spontaneous firing and whisker-evoked responses from a group of cells that were subjected to either cholinergic (n = 8 cells) or noradrenergic (n = 9) activation. The spontaneous firing was measured as the spikes per stimulus during a 20 ms time window after the whisker deflection. The values during the control (quiescent) state for these cells were not significantly different for cells subjected to carbachol or norepinephrine, and thus they were averaged together in Figure 2A for illustration purposes (control; n = 17). However, all statistical comparisons are made within subjects by using the corresponding control values for each cell. As with the cell shown in Figure 1, all VPM cells increased their spontaneous firing during carbachol (p < 0.01) and all decreased their spontaneous firing during norepinephrine (p < 0.01). In fact, most cells had no spontaneous firing during norepinephrine. Despite the reduction of spontaneous firing during norepinephrine, the responses to whisker stimulation were robust. Figure 2A illustrates whisker responses evoked by stimulating the PW and four AWs at low frequencies (0.1 and 2 Hz) and at high frequencies (10 and 20 Hz) during control states and during either cholinergic or noradrenergic activation. For low-frequency whisker stimulation, PW responses were robust during any of the three states, but AW responses were significantly stronger during carbachol than during the control state for all four AWs, indicating that the size of the receptive field had enlarged (p < 0.01). In contrast, during norepinephrine, only the first AW (i.e., the AW with the strongest response) produced a response that was significantly stronger than during the
control state ($p < 0.01$). The other three AWs during norepinephrine did not change their responses significantly compared with control. Thus, receptive fields are larger during cholinergic than during noradrenergic thalamic activation.

For high-frequency whisker stimulation, during the control state, responses showed strong adaptation (i.e., responses are suppressed by high-frequency whisker stimulation). During carbachol, responses to high-frequency stimulation were significantly enhanced for the PW and all AWs ($p < 0.05$), resulting in a virtual elimination of adaptation (Castro-Alamancos, 2002a) for the PW. During norepinephrine, the response to high-frequency stimulation was significantly enhanced when compared with the control state only for the PW ($p < 0.01$ for 10 and 20 Hz PW responses). However, the effect of norepinephrine at enhancing high-frequency PW responses was not as strong as that of carbachol; PW high-frequency responses during norepinephrine were significantly smaller than during carbachol ($p < 0.05$ carbachol vs norepinephrine for 10 and 20 Hz PW responses, independent $t$ test). Thus, cholinergic activation is more effective than noradrenergic activation at reducing sensory adaptation in the thalamus.

A measure of the response evoked by a sensory stimulus that considers the saliency of that stimulus with respect to the background noise is the signal-to-noise ratio. The signal-to-noise ratio was calculated by dividing the whisker-evoked response value by the mean spontaneous firing rate for the equivalent 20 ms period for each cell. Because cells could have no spontaneous activity (e.g., during norepinephrine) or may not respond to a whisker, we substituted zeros with a minimum value (0.0016) for these calculations only. This allowed calculating signal-to-noise ratios for all of the cells, including those that had no spontaneous firing. Figure 2B compares the signal-to-noise ratios during cholinergic and noradrenergic activation. Norepinephrine produced signal-to-noise ratios that were significantly stronger than those of carbachol for all stimulation frequencies of the PW and the first AW and also for low-frequency whisker stimulation of the second AW ($p < 0.05$). Thus, noradrenergic activation in the thalamus leads to significantly higher signal-to-noise ratios than cholinergic activation (Fig. 2B).

In a previous study, we determined that the enlargement of the receptive fields of VPM cells and increased spontaneous firing caused by carbachol is mediated by muscarinic receptors because the effect is blocked by applying atropine (1–2 mM) in the thalamus (Aguilar and Castro-Alamancos, 2005). Here, we found that the effects of norepinephrine were reversed by the $\alpha$-adrenergic antagonist phentolamine (1 mM; $n = 3$) applied directly into the thalamus via the microdialysis cannula (supplemental Fig. S1, available at www.jneurosci.org as supplemental material).

### Multiwhisker versus single-whisker responses during noradrenergic activation

During thalamic cholinergic activation, VPM cells respond to a PW and several AWs. However, when the PW and AWs are stimulated simultaneously (multiwhisker stimulation), the response resembles the PW response, as if the AWs had not been stimulated (Aguilar and Castro-Alamancos, 2005). In the present study, we addressed whether noradrenergic activation also leads to differences between single-whisker and multiwhisker stimulation. Figure 3A shows PSTHs comparing the responses of VPM cells ($n = 5$) to simultaneous stimulation of six whisks with the sum of the responses to each of the six individual whiskers stimulated at low (0.1 Hz) frequencies. As with cholinergic activation, noradrenergic activation resulted in a sublinear summation of multiwhisker responses; the response to all six whiskers stimulated together was significantly smaller than the sum of the responses to each of the six whiskers ($p < 0.001$). Moreover, when the PSTHs to all six whiskers stimulated together are overlaid with those of the PW whisker alone, they are very similar during both the control state and norepinephrine (Fig. 3B). When these responses are compared (Fig. 3C), they are not significantly different for low- or high-frequency whisker stimulation. Note that there is a tendency for the response to all six whiskers to be smaller than the PW response; this is attributable to the near absence of long-latency spikes during multiwhisker stimulation (Aguilar and Castro-Alamancos, 2005). Thus, during either cho-
linergic or noradrenergic thalamic activation, the responses to stimulation of all six whiskers together resemble the responses to stimulation of the PW alone, as if the AWs had not been stimulated.

**Effect of cholinergic and noradrenergic activation on corticothalamic responses**

Up to this point, we described the effects of noradrenergic activation on sensory-evoked responses in VPM cells and compared them with the effects of cholinergic activation. From this point forward, we describe the effects of cholinergic and noradrenergic activation on corticothalamic responses evoked in VPM cells by stimulating the thalamic radiation.

Single-unit recordings were obtained from VPM cells, and a stimulating electrode was placed in the thalamic radiation to evoke corticothalamic responses. A hallmark of the corticothalamic pathway is that it produces strong frequency-dependent facilitation (for review, see Castro-Alamancos, 2004b). Accordingly, stimulating and recording electrodes were carefully aligned so that low-intensity stimulation (< 150 μA) elicited virtually no response to low-frequency stimulation (0.1 Hz) but a near 100% firing to stimulation at 10 Hz. Moreover, the responses produced at 10 Hz were phase-locked to the stimulus and of short latency (< 7 ms).

During the alignment, we purposefully ensured that none of the recorded cells were driven antidromically by the thalamic radiation stimulation at the intensities used. Obviously, other thalamocortical cells, whose axons are coursing closer to the stimulating electrode, are being antidromically discharged. The main consequence of this antidromic discharge in the thalamus will be the recruitment of recurrent (feedback) inhibition from the nRt via thalamocortical fibers. However, because the nRt is already being recruited by feedforward corticothalamic fibers, this should simply add more of the same, meaning that nRt-mediated inhibition could be enhanced (for a detailed discussion, see Castro-Alamancos, 2004b). Also, the stimulation of thalamocortical axons in the radiation means that action potentials travel to neocortex and stimulate corticothalamic cells via thalamocortical synapses in layer VI. However, this thalamocortical EPSP may arrive in layer VI very close in time with the backpropagating action potential of the stimulated corticothalamic cells, which could impede the generation of additional action potentials because of refractory periods. In conclusion, although these issues must be kept in foresight, we believe that they do not significantly affect the conclusions derived from the present study.

Once the recording and stimulating electrodes had been aligned, the thalamic radiation was stimulated with 10 pulses at different frequencies (0.1, 2, 5, 10, 20, and 40 Hz) for a minimum of 20 trials for each frequency. This protocol was repeated at least twice for every frequency during control conditions and then again twice after application of the drugs via microdialysis. The spontaneous firing of each cell was measured by counting the number of spikes during the 3 s period at the beginning of each for the 20 stimulation trials before the corticothalamic stimulation was delivered (i.e., 60 s period). This value is then converted to either spikes per second (in Hertz) or spikes per 20 ms for comparison with the whisker-evoked responses. As reported above, carbachol significantly enhanced the spontaneous firing of the cells (p < 0.01; n = 8) (Fig. 4A).

**Figure 3.** Population PSTHs of multiwhisker responses during noradrenergic activation. A, Population PSTH (n = 5) of responses evoked by simultaneously stimulating six whiskers at 0.1 Hz, including the PW and five AWs (top row), and PSTHs consisting of the algebraic sum of the responses to each of these six whiskers stimulated alone (middle row) during control conditions and during application of norepinephrine in thalamus. The right column and bottom row overlay the PSTHs of responses shown in the vertically or horizontally aligned panels for comparison. B, Population PSTH (n = 5) of responses evoked by stimulating the PW alone (thin traces) or all six whiskers simultaneously, including the PW (thick traces). Different panels correspond to responses during control conditions and noradrenergic activation, during low-frequency (0.1 Hz) and high-frequency (10 Hz) whisker stimulation. C, Comparison of responses evoked by the PW and all six whiskers during the different conditions. Responses were measured by summing the spikes evoked during a 20 ms time window after the whisker stimulus (n.s., not significant).
were mostly absent for all cells. As the stimulation frequency increased, corticothalamic responses became apparent, showing strong frequency-dependent facilitation. During application of carbachol into the thalamus, responses to low-frequency stimulation increased significantly ($p < 0.05$ carbachol vs control for 0.1 and 2 Hz), whereas high-frequency responses did not change significantly.

In another population of cells, we tested the effects of noradrenergic activation on corticothalamic responses by applying norepinephrine into the thalamus via microdialysis. As reported above, norepinephrine significantly reduced the spontaneous firing of the cells ($p < 0.01; n = 9$) (Fig. 5A). Figure 5A displays population data ($n = 9$) of corticothalamic responses corresponding to the number of spikes evoked during a 20 ms window after each stimulus in a train delivered at different frequencies during control conditions and during application of norepinephrine in the thalamus. The PSTHs correspond to the 10th stimulus in the train.
effects should lead to changes in the amount of facilitation of corticothalamic responses. Indeed, Figure 6A plots the amount of facilitation during control and either cholinergic or noradrenergic activation. Facilitation was calculated for each cell as the percentage change of the steady-state response (response to the 10th stimulus) as a function of the low-frequency response at 0.1 Hz. As above, to make these calculations, zeros were substituted with a minimum value (0.0016). Also, for illustration purposes only, the control values for cells subjected to carbachol or norepinephrine were averaged together and are displayed in Figure 6A, but all statistics were computed within subjects. Carbachol resulted in a significant reduction of facilitation for high frequencies ($p < 0.01$; 5, 10, 20, and 40 Hz). In contrast, norepinephrine resulted in a significant reduction of facilitation for low frequencies ($p < 0.05$; 2 Hz) and in the enhancement of facilitation for the highest frequency ($p < 0.01$; 40 Hz).

In addition, we computed signal-to-noise ratios for each cell during cholinergic and noradrenergic activation by dividing the steady-state response for each frequency by the spontaneous firing and compared these values during cholinergic and noradrenergic activation. As shown in Figure 6B, during noradrenergic activation in the thalamus, VPM cells have significantly larger signal-to-noise ratios for stimulation frequencies above 5 Hz than during cholinergic activation (between subjects, $p < 0.05$; 5–40 Hz). Thus, noradrenergic activation sharply increases signal-to-noise ratios for high-frequency corticothalamic responses. We also tested whether the effects produced by cholinergic and noradrenergic activation on corticothalamic responses were blocked by specific receptor antagonists (supplemental Fig. S2, available at www.jneurosci.org as supplemental material). The enhancement of VPM cell firing caused by cholinergic activation was blocked by atropine ($n = 3$), whereas the suppression of spontaneous VPM cell firing caused by noradrenergic activation was blocked by phentolamine ($n = 3$). In addition, the results revealed that atropine reversed the effects of carbachol on both facilitation and signal-to-noise ratios, whereas phentolamine reversed the effects of norepinephrine on the signal-to-noise ratios (supplemental Figs. S1, S2, available at www.jneurosci.org as supplemental material).

**Effect of cholinergic and noradrenergic activation on nRt cell firing**

The present results indicate that noradrenergic activation and cholinergic activation produce significantly different effects on sensory and corticothalamic signals to the thalamus. The major difference is that norepinephrine has a strong inhibitory effect on spontaneous firing, AW responses, and low-frequency corticothalamic responses of VPM cells, whereas carbachol has the opposite effects.

The effects of norepinephrine could be caused by an increase in the firing of nRt cells, which would then tonically inhibit VPM cells. To test this possibility, we recorded from nRt cells, which were identifiable by their typical long-lasting responses to sensory stimulation and long burst firing (Hartings et al., 2000, 2003; Castro-Alamancos, 2004b). Interspike interval histograms and autocorrelations for an nRt cell are shown in supplemental Figure S3 (available at www.jneurosci.org as supplemental material). nRt cells strongly enhanced spontaneous firing during noradrenergic activation and suppressed firing during cholinergic activation. An example of the effect of norepinephrine and carbachol application on an nRt cell is shown in Figure 7A (supplemental Fig. S3, available at www.jneurosci.org as supplemental material). During the control condition, the cell fired at $\pm 2.1$ Hz, whereas during norepinephrine, the cell firing was 72 Hz. Subsequent application of carbachol to the same cell reduced the spontaneous firing to nil. The suppression of spontaneous firing caused by carbachol was completely reversed by atropine (1 mM; data not shown). Figure 7B shows an example of the responses of the nRt cell to whisker stimulation delivered at 0.1 and 5 Hz before and after norepinephrine. Note that, although the spontaneous firing increased, the response to the whisker stimulus was also salient, suggesting that recurrent inhibition was still present on top of the enhanced tonic inhibition. Figure 7C shows population data from nRt cells subjected to either cholinergic or noradrenergic activation. Data from a group of VPM cells are also included for comparison. Norepinephrine significantly enhanced the spontaneous firing of nRt cells but decreased the firing of VPM cells, and carbachol significantly reduced the spontaneous firing of nRt cells while significantly enhancing the firing of VPM cells. Thus, cholinergic and noradrenergic activation has opposite effects on nRt and VPM cells.

These data suggest that, during noradrenergic activation, VPM cells are being inhibited by the robust firing of nRt cells. To test this possibility directly, we recorded from VPM cells and infused GABA$_A$ and GABA$_B$ receptor antagonists (BMI and CGP35348, 100–300 $\mu$M and 5–10 mM, respectively) directly into the thalamus via microdialysis and then applied norepinephrine. Norepinephrine no longer suppressed the spontaneous firing of VPM cells in the absence of thalamic inhibition. In fact, during disinhibition, norepinephrine significantly enhanced the firing of VPM cells from $3.6 \pm 1.9$ Hz (mean $\pm$ SD) to $6.5 \pm 2.9$ Hz (BMI–CGP vs BMI–CGP–norepinephrine; $n = 8$ cells; $p < 0.05$). These results demonstrate that the suppression of spontaneous VPM cell firing produced during norepinephrine is caused by tonic inhibition of VPM cells attributable to increased firing of nRt cells. Consequently, these results indicate that the enhancement in signal-to-noise ratios characteristic of noradrenergic activation is caused by tonic inhibitory input on VPM cells produced by nRt firing.

**Discussion**

Neuromodulators are released in the thalamus during different behavioral states. For example, acetylcholine is released during...
paradoxical sleep and during waking, whereas norepinephrine is released during waking in relation to high levels of vigilance and attention. Here we show that noradrenergic activation affects sensory and corticothermal signals in the whisker thalamus differently than cholinergic activation. For sensory signals, noradrenergic activation sets bottom-up thalamic processing to a focused and noise-free excitatory receptive field, which contrasts with the broad and noisy excitatory receptive field characteristic of cholinergic activation. For corticothermal signals, noradrenergic activation sets top-down processing to a noise-free high-frequency signal detection mode, whereas cholinergic activation produces a noisy broadband signal detection mode. Apparently, noradrenergic activation sets thalamocortical information processing to a mode that seems more appropriate for sensory discrimination and selective attention.

Noradrenergic activation suppresses spontaneous firing of VPM cells

Previous work, mostly in slices, revealed the effects of acetylcholine and norepinephrine on thalamic cells (for review, see McCormick, 1992). Thalamocortical cells are depolarized by either acetylcholine or norepinephrine, whereas nRT cells are hyperpolarized by acetylcholine (Ben Ari et al., 1976; McCormick and Prince, 1986; Pinault and Deschenes, 1992) and depolarized by norepinephrine (Kayama et al., 1982; McCormick and Wang, 1991). The depolarizing effect of acetylcholine and norepinephrine on thalamocortical cells is mediated by muscarinic and α-adrenergic receptors, respectively, which block a resting K⁺ conductance (McCormick and Prince, 1987, 1988). In addition, activation of β-adrenergic receptors enhances the hyperpolarization-activated cation current, which also slightly depolarizes thalamocortical cells (McCormick and Pape, 1990). The hyperpolarizing effect of acetylcholine on nRT cells is produced by activation of a K⁺ conductance (McCormick and Prince, 1986), whereas the depolarizing effect of norepinephrine on nRT cells is mediated by α-adrenergic receptors and attributable to a decrease of a resting K⁺ conductance (McCormick and Wang, 1991).

We found that cholinergic activation leads to an increase of VPM cell firing, whereas noradrenergic activation leads to a reduction of VPM cell firing. The effect of cholinergic activation is easily explained by both a direct depolarization of VPM cells and the suppression of nRT cell firing. However, the effect of noradrenergic activation is more complex. It would be explained if the nRT was suppressing VPM cell firing during noradrenergic activation. Indeed, we found that the suppressive effect of norepinephrine on VPM cell firing was completely mediated by the nRT because, during thalamic disinhibition (block of GABA receptors), norepinephrine no longer suppressed VPM cells. In fact, during disinhibition, VPM cells were excited by norepinephrine. Thus, noradrenergic activation strongly excites nRT cells, which inhibit VPM cells. The differential effects of these neuromodulators on spontaneous firing results in different signal-to-noise ratios in VPM cells (see below).

The considerable differences in spontaneous firing of thalamocortical cells during noradrenergic and cholinergic activation suggest that these neuromodulators may set two different modes of cortical rapid sensory adaptation, because a major determinant of adaptation in the thalamocortical pathway is the firing of thalamocortical cells (Castro-Alamancos and Oldford, 2002; Chung et al., 2002; Castro-Alamancos, 2004a,b). Perhaps cortical sensory adaptation would be reduced during cholinergic activation but present during noradrenergic activation. Future work will need to address this intriguing possibility.
Compared with cholinergic activation noradrenergic activation focuses receptive fields

VPM cells respond to deflections of multiple whiskers during light anesthesia (Simons and Carvell, 1989; Armstrong-James and Callahan, 1991; Diamond et al., 1992; Nicolesis and Chapin, 1994; Friedberg et al., 1999; Brecht and Sakmann, 2002; Mennery et al., 2003). Thus, excitatory receptive fields of VPM cells consist of an excitatory center, the PW, and an excitatory surround, the AWs. Cholinergic activation enlarges the excitatory surround of VPM cells (Aguilar and Castro-Alamancos, 2005). In contrast, we found here that noradrenergic activation caused by application of norepinephrine enhanced AW responses but only for one whisker and for low-frequency responses. Consequently, VPM receptive fields are more focused during noradrenergic activation than during cholinergic activation. The selectivity of noradrenergic activation is also present at the temporal level; norepinephrine was less effective than carbachol at facilitating high-frequency responses to whisker stimulation. Whereas carbachol enhanced high-frequency responses for several AWs, norepinephrine only enhanced high-frequency responses for the PW. This indicates that high-frequency sensory inputs are highly focused to the center of the receptive field during noradrenergic activation. The present results demonstrate that different neuromodulators provide the means to dynamically modify receptive field sizes during arousal, from a more focused receptive field characteristic of noradrenergic activation to a broader receptive field typical of cholinergic activation.

When the PW and several AWs are stimulated simultaneously, the response to the PW prevails and AW responses are absent, as if only the center (PW) of the excitatory receptive field is represented during multiwhisker stimulation. This nonlinear summation of responses occurs during both quiescent states and thalamic cholinergic activation (Aguilar and Castro-Alamancos, 2005). In the present study, we found that noradrenergic activation also leads to a nonlinear summation of responses during multiwhisker stimulation. In fact, all response components other than the short-latency PW response are abolished during multiwhisker stimulation in all conditions investigated so far. Therefore, VPM cells represent only the center of the receptive field when center and surround are simultaneously stimulated during either cholinergic or noradrenergic activation.

Noradrenergic activation high-pass filters corticothalamic signals

In addition to studying how cholinergic and noradrenergic activation affects sensory inputs, we studied how they affect corticothalamic inputs (for a recent review on corticothalamic pathways, see Castro-Alamancos, 2004b). Morphologically and physiologically, VPM cells and upper layer VI corticothalamic cells form closed loops for the flow of information between a thalamic barreloid and a cortical barrel column (Bourassa et al., 1995; Temereanca and Simons, 2004). Corticothalamic synapses display strong frequency-dependent facilitation (Turner and Salt, 1998; Castro-Alamancos and Calcagnotto, 1999, 2001; Turner and Salt, 1999; Golshani et al., 2001), which contrasts with the frequency-dependent depression of primary sensoryafferents. Corticothalamic EPSPs are suppressed by both acetylcholine and norepinephrine, an effect that is independent of the postsynaptic actions of these modulators (Castro-Alamancos and Calcagnotto, 2001).

In a previous study, we recorded corticothalamic field potential responses in somatosensory thalamus and found that thalamic activation produced by brainstem reticular formation stimulation led to the high-pass filtering of corticothalamic responses (Castro-Alamancos and Calcagnotto, 2001); low-frequency responses were suppressed during activated states. This clearly distinguishes corticothalamic from sensory inputs, because sensory inputs drive thalamocortical cells very effectively at low and high frequencies during activated states. Thus, the high-pass filtering is very useful because it ensures that thalamocortical cells are not driven by cortical signals unless those signals arrive at high frequencies.

The present study demonstrates that the high-pass filtering of corticothalamic responses is a selective effect of noradrenergic activation states, which are related to high levels of vigilance and selective attention that should favor sensory relay. In contrast, cholinergic activation augments low-frequency corticothalamic responses, which reduces the amount of facilitation in corticothalamic responses, making VPM cells responsive to a wide frequency band of cortical signals. Hence, during cholinergic activation, the selectivity of VPM cells for high-frequency corticothalamic signals is lost. Certainly, this causes a major problem for thalamocortical sensory processing, because it allows low-frequency cortical signals to become as effective as sensory inputs in driving thalamocortical cells. Such an effect seems undesirable during sensory processing, because thalamocortical cells would not be able to distinguish sensory and cortical inputs.

One possibility is that the enhanced responsiveness to low-frequency cortical signals during cholinergic activation is related to sensory experiences that are driven by internal, top-down, representations during paradoxical sleep (when cholinergic activation is strong and noradrenergic activation is absent). During paradoxical sleep, cortical cells would be strong drivers of thalamocortical neurons, which may serve to feed top-down representations to upper layers of primary sensory cortex via the thalamus, perhaps related to sensory experiences during this phase of sleep. An important, yet unresolved, issue relates to the firing of corticothalamic cells during behavior. Corticothalamic cells are notorious for being difficult to drive with sensory stimulation (Swadlow and Weyand, 1987; Swadlow, 1990) and have low firing rates even during certain behaviors (Sirota et al., 2005). Hence, it is important to determine when corticothalamic cells fire at high frequencies.

nRt regulates signal-to-noise levels in thalamocortical cells

nRt cells that are responsive to whisker stimulation fire spontaneously in two different modes, a continuous tonic mode and a burst mode (Hartings et al., 2003). Noradrenergic activation produces the continuous tonic firing mode, whereas cholinergic activation suppresses nRt cell firing. In turn, cholinergic activation enhances the spontaneous firing of VPM cells, causing signal-to-noise ratios to drop, whereas noradrenergic activation reduces the spontaneous firing of VPM cells, causing signal-to-noise ratios to increase sharply. These effects were found to be caused by the firing of nRt cells because, during block of GABA receptors in the thalamus, norepinephrine no longer suppressed the spontaneous firing of VPM cells or enhanced signal-to-noise ratios for sensory or cortical inputs. In fact, during disinhibition, norepinephrine excited VPM cells. Our results indicate that a primary role of the nRt during sensory processing is to regulate the noise in thalamocortical networks.

An important question related to the effects of norepinephrine is why whisker responses remain in VPM cells if they are strongly inhibited by GABAergic inputs from the nRt. The reason for this seems to be the all-or-none, fast-rising and large-amplitude EPSPs that are triggered by whisker stimulation at
lemniscal synapses (Castro-Alamancos, 2002a,b), which would overcome the inhibition by nRT. Additional intracellular work is needed to address this question.

**Functional implications**

LC cells, the source of norepinephrine to the thalamus, discharge robustly during high vigilance and attentive states, less intensely during waking, and do not fire during sleep, particularly during paradoxical sleep (see Introduction). Thus, noradrenergic activation may provide a dynamic mechanism to (1) focus thalamocortical receptive fields, (2) high-pass filter corticothalamic signals, and (3) enhance signal-to-noise ratios, as cognitive processing demands. We speculate that the more focused receptive fields and signal-to-noise ratios observed during noradrenergic activation compared with cholinergic activation reflect a more appropriate, or at least different, information processing mode for spatial discrimination of sensory inputs, whereas the larger receptive fields, lower signal-to-noise ratios, and broad-frequency spectrum responses to cortical inputs characteristic of cholinergic activation may be related to activated states during paradoxical sleep and nonattentive wakefulness.

**References**


