Effects of Hypocretin/Orexin and Major Transmitters of Arousal on Fast Spiking Neurons in Mouse Cortical Layer 6B

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Abstract

Fast spiking (FS) GABAergic neurons are thought to be involved in the generation of high-frequency cortical rhythms during the waking state. We previously showed that cortical layer 6b (L6b) was a specific target for the wake-promoting transmitter, hypocretin/orexin (hcrt/orx). Here, we have investigated whether L6b FS cells were sensitive to hcrt/orx and other transmitters associated with cortical activation. Recordings were thus made from L6b FS cells in either wild-type mice or in transgenic mice in which GFP-positive GABAergic cells are parvalbumin positive. Whereas in a control condition hcrt/orx induced a strong increase in the frequency, but not amplitude, of spontaneous synaptic currents, in the presence of TTX, it had no effect at all on miniature synaptic currents. Hcrt/orx effect was thus presynaptic although not by an action on glutamatergic terminals but rather on neighboring cells. In contrast, noradrenaline and acetylcholine depolarized and excited these cells through a direct postsynaptic action. Neurotensin, which is colocalized in hcrt/orx neurons, also depolarized and excited these cells but the effect was indirect. Morphologically, these cells exhibited basket-like features. These results suggest that hcrt/orx, noradrenaline, acetylcholine, and neurotensin could contribute to high-frequency cortical activity through an action on L6b GABAergic FS cells.

Key words: acetylcholine, arousal, neurotensin, noradrenaline, sleep

Introduction

Fast spiking (FS) cortical neurons are GABAergic interneurons (McCormick et al. 1985; Connors and Gutnick 1990; Ascoli et al. 2008) that have taken an ever increasing importance in our understanding of cortical network activity. Their denomination is based on their characteristic brief action potentials, their firing with little or no adaptation and their ability to reach a high level of activity when depolarized (McCormick et al. 1985; Connors and Gutnick 1990). Early on, most of them were also shown (Kawaguchi and Kubota 1993) to express the calcium-binding protein parvalbumin (PV). Morphologically, FS parvalbumin positive (FS-PV) cells were found to be of only two types, chandelier or basket cells (Povysheva et al. 2013; Inan and Anderson 2014). In an extension of early studies showing that networks of GABAergic cells can generate fast (40 Hz) synchronized oscillations (Whittington et al. 1995), FS-PV cells have now been shown to participate in the production of a high-frequency rhythm (“gamma band oscillation” or GBO) involved in cognitive processes during the waking state (Freud 2003; Cardin et al. 2009; Sohal et al. 2009; Hu et al. 2014; Kim et al. 2015).
This rhythm was shown to be modulated by the waking state, being reduced during slow wave sleep and increased during waking (Maloney et al. 1997) under the influence of the systems of arousal which notably express noradrenaline, acetylcholine, and the neuropeptide hcrt/orx (Jones 2011). GBO was also shown to be promoted by neurotensin, a peptide colocalized in hcrt/orx neurons (Furutani et al. 2013), through an action on basal forebrain cholinergic neurons (Cape et al. 2000).

Among the transmitter pathways implicated in arousal, the hcrt/orx system holds a special status. It is indeed the only one indispensable for maintaining waking as a deficit in the production of hcrt/orx or an alteration of hypocretin/orexin receptors are the only known conditions which result in narcolepsy in animals and humans (for recent reviews on hcrt/orx, see Siegel and Boehmer 2006; Alexandre et al. 2013; Jones and Hassani 2013; Mahlios et al. 2013; de Lecea and Huerta 2015) of the arousal systems originating from the brainstem and the basal forebrain (for reviews, see Jones 2011; Jones and Hassani 2013).

Beyond its action on the arousal systems, hcrt/orx was also suggested to affect the cerebral cortex more directly. Indeed, of a particular interest was firstly the demonstration that hcrt/orx exerted an action in the thalamus, and that in this structure it targeted specifically, and exclusively, the widespread cortical projection system of the intralaminar and midline nuclei (Bayer et al. 2002). Secondly, it was shown that hcrt/orx directly affected cortical neurons, but again, at that level, it exerted a direct postsynaptic action specifically and exclusively on neurons of a single sublayer, the layer 6b (Bayer et al. 2004), which is thought to exert a widespread action on the overlying cortex through local or distant corticocortical connections (Clancy and Cauller 1999; Clancy et al. 2001; Arimatsu et al. 2003). This layer, a remnant of the subplate (for review, see Kanold and Luhmann 2010), has been shown to persist into adulthood in many species either as a distinct sublayer at the very bottom of the layer 6 or as a collection of scattered neurons (known as “interstitial neurons”) located in the white matter of the corpus callosum (Kostovic and Rakic 1980; Reep and Goodwin 1988; Chun and Shatz 1989; Reep 2000; Clancy et al. 2001; Friedlander and Torres-Reveron 2009).

The goal of the present study was thus to identify FS cells in layer 6b of mouse somatosensory cortical slices and determine whether they are sensitive to hcrt/orx as well as to other transmitters associated with high-frequency cortical rhythms such as noradrenaline, acetylcholine, and neurotensin (Jones 2011). Preliminary results have been published in an abstract form (Wenger et al. 2011).

Materials and Methods

Slice Preparation

Brain slices were obtained from juvenile C57Bl/6 mice (Charles River Laboratories, France) and GAD67-GFP (line G42) transgenic mice (courtesy of Dr Alan Carleton, Geneva University, Geneva, Switzerland) from 17 to 21 day old. These transgenic mice express GFP under the control of a GAD67 bacterial artificial chromosome (BAC) clone. In this line, GFP-positive cells selectively express parvalbumin (although only about 50% of the PV neurons express GFP) in the cortex and most of them have been described as FS basket cells (Chattopadhyaya et al. 2004; Xu and Callaway 2009). Mice were treated according to the rules of the Swiss Federal Veterinary Office (approval ID: 31.1.1007/3248/0). They were anesthetized by isoflurane (Forene, Abbott AG, Baar, Switzerland), decapitated and the brain was extracted. Coronal sections (300 µm-thick), at the level of the primary somatosensory (SSp) cortex, were then cut on a vibrating-blade microtome (Leica VT1200s; Leica Biosystems, Nussloch, Germany) into a cold slicing medium (Nashmi et al. 2002) containing (in mM): 135 N-Methyl-D-glucamine (NMDG), 0.5 CaCl₂, 20 choline bicarbonate, 10 glucose, 1 KCl, 1.2 KH₂PO₄, 1.5 MgCl₂ (NMDG was first dissolved and then titrated to pH 7.4 with HCl). Following the slicing, tissue was left to recover at 37°C for 1 h and then kept at room temperature (RT), until use, in artificial cerebrospinal fluid (ACSF) containing (in mM): 2.4 CaCl₂, 10 glucose, 5 KCl, 1.2 KH₂PO₄, 130 NaCl, 20 NaHCO₃, and 1.3 MgSO₄ aerated with carbogen (95% O₂ and 5% CO₂). During the recovery period and the time left at RT, 1 mM of kynurenic acid was added to the ACSF. Individual slices were then transferred to a thermoregulated (32°C) recording chamber placed on the stage of an upright microscope (Axioskop, Zeiss, Oberkothen, Germany) equipped with an infrared camera (TILL-Photonics) and a mercury lamp (HBO 100, Zeiss) as well as selective filter sets (filter sets 20 and 38, Zeiss) for fluorescent cell identification. Slices were maintained immersed and were continuously superfused (4–5 mL/min.) with ACSF bubbled with 95% O₂ and 5% CO₂.

Electrophysiological Recordings

Recordings were performed on cells located in cortical L6b (Fig. 1A). This layer is defined as a thin and compact band of neurons lying immediately above the white matter (WM) of the corpus callosum, in the deepest part of the gray matter, below the layer 6a (Valverde et al. 1989; Clancy and Cauller 1999; Reep 2000). In our experiments, such a compact cell layer could be identified under infrared microscopy and neurons therein were preferably recorded the closest to the WM/L6b boundary. Regarding the electrophysiological experiments on GAD67-GFP (line G42) transgenic mice, fluorescent cells in the L6b were first

![Figure 1. Recordings in L6b. (A) Schematic representation (adapted from the Allen Mouse Brain Atlas; mouse brain-map.org) of the different layers of the primary somatosensory cortex (SSp) and the underlying white matter (WM), together with an infrared microscope image of a neuron recorded in this study. (B) Localization of an Alexa Fluor dye-injected neuron in layer 6b (arrow) with fluorescent Nissl counterstaining. (C1–3) Confocal snapshot images of a L6b cell recorded in a GAD67-GFP (line G42) transgenic mouse, showing Alexa Fluor dye-injected fluorescence (C1) and GFP fluorescence (C2) colocalized (C3) in the same cell. Scale bar in C1–3: 10 µm.]
visualized with a selective filter set for GFP (filter set 38, Zeiss). Once a GFP-positive cell was selected, it was visualized with infrared illumination and recorded as described below.

The recordings were performed in the whole-cell configuration in either current-clamp or voltage-clamp mode (Ushakov et al. 2011) using an Axopatch 200A amplifier (Molecular Devices, Sunnyvale, CA, USA) and pClamp 10.0 software (Molecular Devices). Patch pipettes (7–12 MΩ for current-clamp and 4–5 MΩ for voltage-clamp) were pulled on a DMZ universal puller (Zeit-Insititut, Munich, Germany) from borosilicate glass capillaries (GC150F-10, Harvard Apparatus, Holliston, MA, USA) and filled with an intrapipette solution containing (in mM): 3 ATP, 0.1 BAPTA, 0.1 GTP, 10 HEPES, 4 KCl, 126 KMeSO₄, 5M MgCl₂, 8 phosphocreatine disodium salt, pH 7.3 (285 mOsm). In order to validate the colocalization of the recorded cell with the GFP-positive one and allow for a subsequent morphological analysis, Alexa Fluor 555 hydrazide (200 μM; Molecular Probes by Life Technologies, Carlsbad, CA, USA) was generally added to the intrapipette solution. The database for the present study consisted of 65 L6b FS cells. Membrane potential, measured immediately after obtaining whole-cell configuration, was (mean ± SEM) −58.7 ± 0.8 mV (n = 61). Membrane potential values were not compensated for junction potential (estimated at −9.6 mV; Eggermann et al. 2003).

In several experiments, synaptic transmission was blocked by either adding TTX (10⁻⁶ M) to the ACSF or increasing its magnesium and lowering its calcium concentration (respectively, 10 mM Mg²⁺ and 0.1 mM Ca²⁺).

Since no qualitative differences in either intrinsic properties, synaptic activity or the effects of transmitters were observed between the cells recorded from wild type or from the GAD67-GFP (line G42) transgenic mice, data from these 2 mouse strain lines were pooled.

Synaptic Activity Recordings and Analysis

Postsynaptic currents (PSCs) were recorded using voltage-clamp mode. During these experiments, access resistance (10–20 MΩ) was monitored regularly and recordings were discarded if it varied more than 20% during an experiment. In order to investigate the nature of the PSCs, either IVs in ACSF (holding membrane potential Vₐ₀ = −60 mV; 1 s-long steps between −100 and −20 mV) or IVs in the absence or the presence of DNQX (holding membrane potential Vₐ₀ = −60 mV; 9.5 s-long steps at −80, −60, and −40 mV) were performed.

To characterize the effect of hcrt/orx on FS L6b neurons, spontaneous (sPSCs) or miniature (mPSCs) postsynaptic currents were recorded (Vₐ₀ = −60 mV; 10 s-long sweeps) in ACSF or in presence of TTX, respectively. The amplitude and the frequency of the PSCs were then analyzed with a custom-written software (Detector, Dr J.R. Huguenard, Stanford University, Stanford, CA, USA) in which individual events were detected with a threshold-triggered process from a sweep of 10 s both before bath-application and during the maximal effect of the transmitter (due to the important decrease in the number of PSCs in the presence of TTX, 3 successive sweeps of 10 s each were analyzed in TTX experiments). Two detection criteria, namely the threshold and the duration of the trigger, were adjusted to ignore electrical noise and allow maximal discrimination of PSCs. In both conditions (ACSF and TTX), each event detected before application of the transmitter was checked visually in order to avoid artifactual detection (i.e., an event which did not show a typical PSC waveform) and to optimize and validate the adjustment of the program parameters.

For each cell, frequency and mean amplitude of the same number of detected events (n = 250) before application and during the maximal effect of the transmitter were calculated and plotted as scatter plots for each condition (ACSF and TTX). The average of frequencies and mean amplitudes of PSCs obtained were plotted as histograms. Changes in cumulative PSC interevent interval and amplitude distributions were also analyzed. Data are presented as mean ± SEM. Statistical analyses were carried out using either unpaired two-tail t-test for analyses involving comparison of means or Kolmogorov–Smirnov (KS) test (XLStat, Addinsoft, France) to compare cumulative distributions. Statistical significance was set to P < 0.01.

Chemicals

The peptide hypocretin1/orxinA (hcrt1/orxA; Bachem, Bubendorf, Switzerland) was used exclusively in the present study since it is known that hcrt1/orxA peptide is as potent as hypocretin2/orxinB (hcrt2/orxB) on OX2R but more active than hcrt2/orxB on OX1R (Sakurni et al. 1998). For brevity, hcrt1/orxA will be referred to as hcrt/orx. Noradrenaline, carbachol, and neurotransin were all obtained from Sigma-Aldrich (Buchs, Switzerland), while AP5, DNQX, and bicuculline were obtained from Tocris Bioscience (Tocris Bioscience, Bristol, UK), and tetrodotoxin (TTX) from Latoxan (Latoxan, Valence, France). All the drugs were bath-applied.

Histology

Following electrophysiological recordings, a subset of slices with only 1 or 2 dye-injected cells were fixed in a solution of 4% paraformaldehyde for 1–2 h. Slices were then counterstained with NeuroTrace 435/455 blue-fluorescent Nissl stain (Molecular Probes by Life Technologies) to confirm the location of Alexa Fluor-filled neurons in L6b (Fig. 1B). For that purpose, after washing in PBS, the slices were incubated twice for 2 h at RT, first in PBS–Trition 0.3% and then in NeuroTrace 1/100 in PBS–Trition 0.3%. After an overnight washing step in PBS, the slices were mounted in FluorSave (Calbiochem) and coverslipped. Images were acquired on an inverted epifluorescence microscope (Nikon Eclipse Ti, Nikon Instruments, Melville, NY, USA) using a 10× magnification objective.

In experiments with transgenic mice, the colocalization of GFP and the Alexa Fluor dye in the recorded cell was always verified at the end of the recording under microscope while the slice was still in the recording chamber. To confirm this colocalization, a subset of dye-injected GFP-positive neurons were fixed and counterstained as described above and snapshot images were acquired (Fig. 1C1–C3) using a confocal laser scanning microscope (Zeiss LSM700, Zeiss).

Neurolucida Reconstruction and Morphological Analysis

Alexa Fluor-filled neurons were imaged using a confocal laser scanning microscope (Zeiss LSM700, Zeiss). Confocal image stacks were acquired at 3 different magnifications (20×, 40×, and 63×) to cover the entire dendritic arborization. Dendritic and axonal trees were then reconstructed in 3D from the confocal stacks with Neurolucida 11 software (MBF Bioscience, Williston, VT, USA). Once the tracing was completed, several somatodendritic and axonal features were analyzed using Neurolucida Explorer 11 software (MBF Bioscience). The reconstructions of neurons were aligned to have the WM/L6b boundary parallel to the x-axis. Only neurons with minimal extracellular dye leakage and with a total axonal length above 1.0 mm were considered. Five somatic parameters were quantified: the cell body perimeter and its area, the maximum and minimum ferets as well as the aspect ratio.
Max/min ferets were defined, respectively, as the longest/smallest diameter of the soma, measured between two parallel tangential lines drawn along the soma boundaries (as if measured with a caliper). The aspect ratio, defined by the ratio of the maximal feret to the minimal feret, gives an indication about the elongation of the cell body. In order to investigate the dendritic arborization, the following parameters were quantified: the number of primary dendrites, the total dendritic length, the number of nodes, the dendritic branching frequency (i.e., number of nodes/100 µm) and the ratio of the total dendritic length to the total dendritic surface. In order to further analyze the spatial distribution of the dendrites, a Sholl analysis and a polar histogram were also performed. The Sholl analysis, used to represent the dendritic density around the cell body, was expressed as a fraction of the total dendritic length contained in a series of spheres centered at the centroid of the soma with incremental radii (100, 200, and 300 µm). The polar histogram, on the other hand, describes the overall orientation of the dendritic processes in which their 3D reconstruction was reduced to a 2D polar histogram with dendritic lengths plotted, in bins, as a function of direction. The dendritic arborization around the soma was indeed divided in 120 bins of 3° with cubic spline smoothing for each dendritic reconstruction. The average of the dendritic reconstructions of the whole set of cells was calculated and a mean polar plot was obtained with Statgraphics Centurion XVII (Statpoint Technologies, Warrenton, VA, USA) in which the 0–180° axis was parallel to the WM/L6b boundary. On the polar plot, processes pointing to 90° were oriented toward the pial surface whereas those directed to 270° were oriented toward the white matter. In order to characterize the axonal arborization and its spatial distribution, a procedure similar to the one outlined above for the dendritic trees, was applied. Each parameter is presented as mean ± SEM.

Results

Whole-cell recordings were performed on FS neurons in L6b (n = 65) of the mouse SSp cortex (Fig. 1A,B). About half of the recordings (n = 30) was obtained from transgenic GAD67-GFP (line G42) mice (Fig. 1C1–C3; see Materials and Methods), while the rest was obtained from wild-type C57Bl/6 mice.

Intrinsic Properties and Spontaneous Synaptic Activity of L6b FS Cells

Fast spiking neurons in L6b of the SSp cortex did not differ from FS cells in other cortical regions (McCormick et al. 1985; Connors and Gutnick 1990; Ascoli et al. 2008) as illustrated in Figure 2. Challenged by a small depolarizing step, they fired in an irregular way (Fig. 2A1), with action potentials typically followed by a deep single after-hyperpolarization (Fig. 2A2). With increasing amplitude of stimulation, they typically showed a clustering of the action potentials (Fig. 2B1, enlarged in 2B2), in a pattern previously described as “stuttering” (Ascoli et al. 2008) as well as subthreshold membrane potential oscillations (Fig. 2B2, dots). Finally, at an even higher amplitude of stimulation, the cells discharged tonically with little adaptation (Fig. 2C1, enlarged in 2C2) at a high frequency (107.8 ± 6.0 Hz; range 50.3–195.1 Hz; n = 35). These cells also presented a small voltage- and time-dependent inward rectification (Fig. 2D1, dot) compatible with the presence of an I_h current.

As illustrated in Figure 2D1–2, the L6b FS cells, while usually displaying no spontaneous action potentials at rest, always exhibited a very high level of spontaneous synaptic activity. This activity consisted of EPSPs as demonstrated with the following arguments. First, in voltage-clamp mode, only inward synaptic currents could be recorded (n = 5/5; holding potentials ranging from −100 to −20 mV, n = 4, and −100 to −30 mV, n = 1) and outward currents were never unmasked (not shown). Second, as illustrated in Figure 3, these inward synaptic currents depended on AMPA/kainate glutamatergic transmission as they were completely blocked by the AMPA/kainate receptor antagonist DNQX (n = 5/5, at 2 × 10^{-5} M) at 3 different holding potentials (−80, −60, and −40 mV). In contrast, neither AP5 (n = 3/3), an NMDA antagonist (applied at 5 × 10^{-5} M), nor bicuculline (n = 2/2), a GABA_A receptor antagonist (applied at 10^{-5} M), had any effect (not shown). These data therefore suggest an absence of spontaneous IPSPs in these L6b FS cells in our conditions.

Hcrt/orx Increases Spontaneous Excitatory Activity in L6b FS Cells

We next tested the effect of hcrt/orx on L6b FS cells (n = 53) in either current-clamp (CC) or voltage-clamp (VC) mode. In CC mode, when neurons were challenged with hcrt/orx (at 10^{-6} M), there was nearly always (n = 36/37) a large increase of spontaneous EPSPs as illustrated in Figure 4A (enlarged in 4B1 and 4B2). In some cells showing this increase of EPSPs, there was an associated membrane depolarization (3.2 ± 0.5 mV; n = 15/36), whereas in others no membrane potential deflection could be observed (n = 21/36). Hcrt/orx, in contrast, had no effect when applied in...
the presence of either a high Mg/Low Ca solution (Fig. 4C; \( n = 3/3 \)) or a TTX-containing ACSF (Fig. 4D; \( n = 2/2 \)). In order to evaluate the effect of h_nr transmitted more precisely on L6b FS cells, it was then tested at \( 10^{-6} \) M in the VC mode. As shown in Figure 5A, h_nr strongly increased the occurrence of spontaneous synaptic currents (\( n = 15/16 \)) in a reversible manner. This effect was entirely explained by a significant increase in the frequency of the synaptic currents with no effect on their amplitude (Fig. 5A). Indeed, a significant change in inter-event interval cumulative distributions (\( P_{\text{KS}} = 10^{-3} \), \( n = 9 \); Fig. 5A) together with a significant increase in mean frequencies (from 107.2 ± 10.7 Hz, \( P_{\text{t-test}} = 2.6 \times 10^{-5} \), \( n = 9 \); histogram in Fig. 5A) were observed, whereas no significant change was seen in either amplitude cumulative distributions (\( P_{\text{KS}} = 10^{-1} \), \( n = 9 \); Fig. 5A) or mean amplitudes (from 8.8 ± 0.5 to 8.7 ± 0.7 pA, \( P_{\text{t-test}} = 0.88 \), \( n = 9 \); histogram in Fig. 5A).

We next considered the effect of h_nr in the presence of TTX at \( 10^{-6} \) M. In this condition, as illustrated in Figure 5B1 (left panel), some residual synaptic activity could still be recorded, indicating the presence of miniature synaptic currents, but this activity did not appear to be sensitive to h_nr (\( n = 5/5 \); Fig. 5B1, middle panel), thereby suggesting that the effect was presynaptic. This was confirmed by analyzing further the results in presence of TTX (Fig. 5B2–B5) which indeed showed an absence of effect of h_nr on inter-event interval cumulative distributions (\( P_{\text{KS}} = 1.0 \), \( n = 5 \); Fig. 5B4), mean frequencies (from 20.5 ± 6.7 to 20.3 ± 6.1 Hz, \( P_{\text{t-test}} = 0.99 \), \( n = 5 \); histogram in Fig. 5B4), amplitude cumulative distributions (\( P_{\text{KS}} = 1.0 \), \( n = 5 \); Fig. 5B5) as well as on mean amplitudes (from 5.6 ± 0.7 to 5.5 ± 0.9 pA, \( P_{\text{t-test}} = 0.94 \), \( n = 5 \); histogram in Fig. 5B5).

Altogether the data suggest a presynaptic action of h_nr but not on axon terminals impinging on L6b FS cells, but rather on neighboring neurons possibly activated directly by h_nr (Bayer et al. 2004).

### Postsynaptic Actions of Noradrenaline and Acetylcholine of L6b FS Cells

As a next step, we then tested the effect of 2 other major wake-promoting transmitters on L6b FS cells that had previously responded to h_nr.

We first tested the effect of noradrenaline (from \( 10^{-5} \) to \( 10^{-4} \) M) and, as illustrated in Figure 6A, the effect was a significant membrane depolarization in 5/6 cells (range 2.1–6.7 mV; \( n = 5 \)) associated in the majority of cases (\( n = 3/5 \)) with the firing of action potentials. In the presence of either a high Mg/Low Ca solution (Fig. 6B) or a TTX-containing ACSF (Fig. 6C), noradrenaline (respectively, from \( 10^{-5} \) to \( 2 \times 10^{-5} \) M and from \( 10^{-5} \) to \( 10^{-4} \) M), in contrast to h_nr, had still an effect as it induced, in both conditions, a significant membrane depolarization (range 2.7–3.9 mV in 3/3 cells in high Mg/Low Ca and range 2.5–5.8 mV in 3/4 cells in TTX). These results indicate a main postsynaptic effect for this transmitter on L6b FS cells.

We then tested the effect of the cholinergic agonist, carbaclo (from \( 10^{-5} \) to \( 10^{-4} \) M). As illustrated in Figure 7A, as for noradrenaline, the effect was a significant membrane depolarization (range 3.0–11.6 mV; \( n = 5/5 \)) associated with the firing of action potentials in the majority of cases (\( n = 3/5 \)). In the presence of either a high Mg/Low Ca solution (Fig. 7B) or a TTX-containing ACSF (Fig. 7C), carbaclo, applied at \( 10^{-5} \) or \( 10^{-4} \) M had still an effect as it induced a significant membrane depolarization in both cases (range 2.3–2.4 mV in 2/3 cells in high Mg/Low Ca and range 1.7–4.2 mV in 3/4 cells in TTX). These results indicate a main postsynaptic effect for acetylcholine on L6b FS cells as was the case for noradrenaline.

### Neurotensin Effect on L6b FS Cells

We finally tested the effect of neurotensin, a transmitter which is colocalized in h_nr neurons (Furutani et al. 2013) and which was shown to promote high-frequency cortical rhythms by an action on basal forebrain cholinergic neurons (Cape et al. 2000). When it was applied (from \( 5 \times 10^{-7} \) to \( 2 \times 10^{-6} \) M) on cells that had previously responded to h_nr, it induced a membrane depolarization in 5/7 tested cells (range 2.7–8.5 mV; \( n = 5 \); Fig. 8A) accompanied in most cells (\( n = 4/5 \)) by the firing of action potentials. This depolarizing effect was however absent when neurotensin was applied in either a high Mg/Low Ca solution (Fig. 8B; \( n = 3/3 \)) or in the presence of TTX (Fig. 8C; \( n = 3/3 \)). These results suggest that the effect of neurotensin must be indirect in this case, in contrast to what was observed for noradrenaline and acetylcholine.

### NeuroLucida Reconstructions of L6b FS Neurons

In order to analyze the morphology of the L6b FS neurons, the recorded cells were injected with a fluorescent red dye, Alexa Fluor 555 hydrazide, and then reconstructed using confocal microscopy and NeuroLucida software, as illustrated in Figure 9A1 and A2. As FS cells are generally classified as basket or chandelier...
cells mainly on the basis of their axonal configuration, only neurons whose axon could be followed for more than 1 mm were included in these analyses. As a consequence of this condition, artifacts due to the truncation of processes, known to occur during the slicing procedure, are minimized, thus leading to neurons with a concomitant better integrity of their dendritic processes. Four cells satisfied this criterion and their morphological characteristics are summarized in Table 1.

These L6b FS cells appeared multipolar with 4–7 primary dendrites radiating in all directions (Fig. 9A1 and A2). Their soma appeared fusiform (elongation 2.0 ± 0.4 μm) with mean feret diameters of, respectively, 18.9 ± 2.6 and 9.9 ± 0.9 μm. They had extensive dendritic and axonal arborizations (respective mean length 2323.3 ± 214.2 and 3239.2 ± 952.9 μm) both running beyond 300 μm away from the soma, according to the Sholl analyses. Dendritic trees were smooth (see inset in Fig. 9A1) with a branching frequency of 1.1 ± 0.2 nodes/100 μm and the main fraction of dendritic arborization (94.4%) was restricted to a 200 μm radius around their soma with no preferential orientation (see the

**Figure 5.** Hcrt/orx increases spontaneous synaptic currents in L6b FS cells. (A1) Effect of hcrt/orx on spontaneous synaptic activity, in normal ACSF, with panels illustrating control (left), effect (middle), and wash (right). (A2–5) Scatter plots of results of control (C) versus hcrt/orx (H/O) with respect to synaptic current frequencies (A2) and amplitudes (A3). (A4–5) Cumulative distributions of interevent interval (A4) and amplitude (A5) together with histograms for mean frequencies and amplitudes in control (C) versus hcrt/orx (H/O). (B1–5) Same as A1–5 but in the presence of TTX. Bars in histograms illustrate the mean ± SEM and significant differences are marked by 3 asterisks (P < 0.01).

**Figure 6.** Noradrenaline depolarizes and excites L6b FS cells. (A) Depolarizing effect of a brief bath-application of noradrenaline (NA). (B) Depolarizing effect of NA in the presence of a high Mg/low Ca ACSF. (C) Depolarizing effect of NA in the presence of TTX.

**Figure 7.** Carbachol depolarizes and excites L6b FS cells. (A) Depolarizing effect of a brief bath-application of the cholinergic agonist, carbachol (carb). (B) Depolarizing effect of carbachol in the presence of a high Mg/low Ca ACSF. (C) Depolarizing effect of carbachol in the presence of TTX.
The neurons in the present study were identified as being in L6b by their location within a thin layer of packed cells lying immediately above the white matter of the corpus callosum, in the deepest part of the gray matter (Valverde et al. 1989; Clancy and Cauller 1999; Reep 2000). Their identification was first done visually under infrared microscopy during the electrophysiological recordings (see Materials and Methods section) and then by confirming the location of a subset of dye-filled cells using a fluorescent Nissl stain procedure. Their location corresponded to that of the neurons of the subplate that survive into adulthood (Valverde et al. 1989; Clancy and Cauller 1999; Reep 2000; Watakabe et al. 2007; Hoerder-Suabedissen et al. 2009).

The L6b neurons studied here were characterized as fast spiking (FS) GABAergic neurons with basket-like characteristics, based on their electrophysiological properties, neurotransmitter content, and morphological reconstruction. With respect to electrophysiology, their identification as FS cells relied on several lines of evidence that include a rather brief action potential followed by a single deep after hyperpolarization, the clustering of action potentials or “stuttering” (as seen in some other FS cortical cells; Ascoli et al. 2008), the high-frequency subthreshold oscillations between clusters and, ultimately, their high-frequency tonic firing with little adaptation when challenged with stronger depolarizing current steps (for review, see Ascoli et al. 2008).

Early on, FS cells were suggested to be GABAergic (McCormick et al. 1985) and later were actually identified as a subtype of cortical GABAergic neurons (for reviews, see Ascoli et al. 2008; Rudy et al. 2011; DeFelipe et al. 2013). The L6b FS cells recorded here were identified as GABAergic as evident from their recording in GAD67-GFP mice. They were also, most probably, parvalbumin-positive cells. Indeed, parvalbumin has been identified as an additional marker in most GABAergic FS cells (Kawaguchi and

### Table 1 Morphological parameters of L6b FS cells

| Morphological parameters | Mean   | SEM   |
|--------------------------|--------|-------|
| Somatic perimeter (µm)   | 46.5   | 4.5   |
| Somatic area (µm²)       | 127.2  | 11.3  |
| Max. feret of the soma (µm) | 18.9  | 2.6   |
| Min. feret of the soma (µm) | 9.9   | 0.9   |
| Somatic elongation        | 2.0    | 0.4   |
| Number of primary dendrites | 5.5   | 0.7   |
| Total dendritic length (µm) | 2323.3 | 214.2 |
| Number of dendritic nodes | 24.5   | 1.9   |
| Dendritic branching frequency (nodes/100 µm) | 1.1   | 0.2   |
| Dendritic length/dendritic surface | 0.7   | 1.3 $\times$ 10^{-5} |
| Dendritic Sholl (0–100 µm) (%) | 58.1  | 2.7   |
| Dendritic Sholl (100–200 µm) (%) | 36.3  | 2.8   |
| Dendritic Sholl (200–300 µm) (%) | 5.4   | 2.1   |
| Dendritic Sholl (>300 µm) (%) | 0.2   | 0.2   |
| Total axonal length (µm) | 3239.2 | 952.9 |
| Number of axonal nodes | 40.3   | 12.9  |
| Axonal branching frequency (nodes/100 µm) | 1.3   | 0.1   |
| Axonal length/axonal surface | 1.6   | 5.1 $\times$ 10^{-3} |
| Axonal Sholl (0–100 µm) (%) | 58.2  | 7.9   |
| Axonal Sholl (100–200 µm) (%) | 37.0  | 5.1   |
| Axonal Sholl (200–300 µm) (%) | 4.5   | 2.9   |
| Axonal Sholl (>300 µm) (%) | 0.3   | 0.3   |

Each parameter is given as a mean ± SEM of the cells reconstructed according to the criterion described in Materials and Methods section (n = 4).

The neurons in the present study were identified as being in L6b by their location within a thin layer of packed cells lying immediately above the white matter of the corpus callosum, in the deepest part of the gray matter (Valverde et al. 1989; Clancy and Cauller 1999; Reep 2000). Their identification was first done visually under infrared microscopy during the electrophysiological recordings (see Materials and Methods section) and then by confirming the location of a subset of dye-filled cells using a fluorescent Nissl stain procedure. Their location corresponded to that of the neurons of the subplate that survive into adulthood (Valverde et al. 1989; Clancy and Cauller 1999; Reep 2000; Watakabe et al. 2007; Hoerder-Suabedissen et al. 2009).

The L6b neurons studied here were characterized as fast spiking (FS) GABAergic neurons with basket-like characteristics, based on their electrophysiological properties, neurotransmitter content, and morphological reconstruction.

With respect to electrophysiology, their identification as FS cells relied on several lines of evidence that include a rather brief action potential followed by a single deep after hyperpolarization, the clustering of action potentials or “stuttering” (as seen in some other FS cortical cells; Ascoli et al. 2008), the high-frequency subthreshold oscillations between clusters and, ultimately, their high-frequency tonic firing with little adaptation when challenged with stronger depolarizing current steps (for review, see Ascoli et al. 2008).

Early on, FS cells were suggested to be GABAergic (McCormick et al. 1985) and later were actually identified as a subtype of cortical GABAergic neurons (for reviews, see Ascoli et al. 2008; Rudy et al. 2011; DeFelipe et al. 2013). The L6b FS cells recorded here were identified as GABAergic as evident from their recording in GAD67-GFP mice. They were also, most probably, parvalbumin-positive cells. Indeed, parvalbumin has been identified as an additional marker in most GABAergic FS cells (Kawaguchi and

### Discussion

In a previous study in rat brain slices, we have shown that in the neocortex only L6b neurons were directly sensitive to the wake-promoting transmitter hcrt/orx (Bayer et al. 2004), a result later confirmed in mice (Wenger et al. 2010). Here, in mouse cortical brain slices, we now describe a set of L6b GABAergic FS cells that are indirectly excited by hcrt/orx and neurotensin and directly depolarized and excited by noradrenaline and acetylcholine, thereby suggesting a possible role for these neurons in promoting cortical high-frequency activity.

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Cortical FS-PV cells have been recently determined to fall into only one of two different morphological classes, chandelier and basket cells (Rudy et al. 2011). In the present study, the Neuronal reconstructions that were carried out on injected neurons indicated that they were not of the chandelier type but rather of the basket cell type. Indeed, none of the reconstructed cells displayed visible vertical axon cartridges, a hallmark feature of chandelier axon (Kawaguchi 1995; Xu and Callaway 2009). In addition, it is noteworthy that chandelier cells are quite rare and that most of the GFP-positive cells of the transgenic mouse line used here had been described previously as basket cells (Chattopadhyaya et al. 2004; Xu and Callaway 2009).

The L6b FS cells of the present study were strongly activated by hcr/orx, an effect which was always indirect. This is suggested first by the evidence that hcr/orx increased the frequency of spontaneous excitatory PSPs, which are conspicuous in these cells (while inhibitory PSPs are absent as evident in DNXQ), without increasing their amplitude. It is confirmed by the absence of hcr/orx effect on miniature synaptic current amplitude in the presence of TTX. The absence, in that latter condition, of hcr/orx effect on miniature synaptic current frequency, suggests however that it is not due to action on glutamatergic terminals, but rather on neighboring cells activated by the peptide. These results stand in contrast to other studies of hcr/orx presynaptic excitatory effects in the cortex that have shown an increase in miniatures and thus, a modulatory presynaptic effect on glutamatergic terminals (Lambe and Aghajanian 2003; Aracri et al. 2015). Finally, one must stress that the reported prevalence of electrical coupling among FS neurons elsewhere in the cortex (Galarreta and Hestrin 1999; Gibson et al. 1999), one cannot exclude that some of the spontaneous activity reported here was actually due to such coupling.

Based on our previous studies that have demonstrated an exclusive postsynaptic action of hcr/orx in L6b neurons in rat brain slices (Bayer et al. 2004) as well as in mice (Wenger et al. 2010), a possible interpretation is that the increased EPSPs recorded here originate from contiguous glutamatergic L6b neurons that are directly (postsynaptically) activated by hcr/orx. The strength of this effect suggests the presence of a tight control of the firing of presumed glutamatergic and secondarily GABAergic neurons in L6b by hcr/orx. Neurotensin, a neuropeptide which is colocalized in hcr/orx neurons (Furutani et al. 2013) and which was shown to be able to promote high-frequency cortical rhythms (Cape et al. 2000), could possibly exert a similar control, given that its strong depolarizing and excitatory action on L6b FS neurons is also entirely indirect.

L6b FS cells were also excited by noradrenaline and acetylcholine but, in contrast to hcr/orx and neurotensin, it was the result of a direct, postsynaptic action. This contention is based on the persistence of depolarizing and/or excitatory responses to these transmitters in conditions of either TTX or synaptic uncoupling. These results are in agreement with numerous studies since the early 80s (for review, see McCormick 1989) which have identified postsynaptic excitatory effects of these transmitters on cortical neurons, including, more recently, effects of noradrenaline on FS neurons (Kawaguchi and Shindou 1998). With respect to acetylcholine, however, the data of the present study seem to be the first to show a postsynaptic depolarizing and excitatory effect on cortical FS neurons. A recent thorough investigation of acetylcholine effects has actually concluded, to the contrary, that FS neurons in the cortex are not directly sensitive to acetylcholine (Gulledge et al. 2007). Nevertheless, it is noteworthy that in the hippocampus, a structure also prone to FS-dependent GBO, FS cells were found to be directly sensitive to acetylcholine (Chiang et al. 2010).

Functionally, one could speculate that the effects of the neurotransmitters on L6b FS neurons described here could contribute to the cortical activation associated with arousal. Fast cortical rhythms and arousal have indeed been known since the early studies of Moruzzi and Magoun (1949) to depend on the influence of the “ascending reticular activating system” even though the exact nature of the system was not understood at that time. Later on, the transmitters implicated in this action, and the pathways containing them, were sorted out and among them, noradrenaline from the locus coeruleus and acetylcholine from the basal forebrain, have been held as prominent players since then (for review, see Jones 2003). More recently, an hypothalamic peptide, hcr/orx (de Lecea et al. 1998; Sakurai et al. 1998), was found to be a major wake-promoting transmitter and as such to be also involved in enabling the cortical high-frequency rhythms associated with the waking state (for recent reviews, see Siegel and Boehmer 2006; Alexandre et al. 2013; Jones and Hassani 2013; Mahlios et al. 2013; Sakurai 2013; de Lecea and Huerta 2014), including possibly by an action on adrenergic (Horvath et al. 1999) and cholinergic (Eggermann et al. 2001; Burlet et al. 2002; Ishibashi et al. 2015) neurons.

In recent years, many studies have proposed that a network of cortical FS-PV cells plays a major role in the genesis of high-frequency rhythms (Freund 2003; Cardin et al. 2009; Sohal et al. 2009; Hu et al. 2014; Kim et al. 2015) but, with the exception of a recent study concerning acetylcholine (Pafundo et al. 2013), the link between this network and the arousal systems referred to above, has thus far remained elusive. It is thus tempting to speculate that what was observed in the present study bears particular significance. It is indeed noteworthy that hcr/orx, which in the somatosensory cortex is thought to exert a direct action only in layer 6b (Bayer et al. 2004; Wenger et al. 2010), appears to directly transfer this excitatory action within L6b to that very class of GABAergic neurons, the FS-PV cells, which today is thought to play a central role in the generation of high-frequency rhythms. It is also noteworthy that these L6b FS cells that are sensitive to hcr/orx, are, at the same time, directly and strongly excited by noradrenaline and acetylcholine. Finally, it is intriguing that the direct effect of acetylcholine on FS cells, as reported here in L6b, appears to be rather unique to this layer, as it was reported to be absent elsewhere in the cerebral cortex (Gulledge et al. 2007).

Altogether these data speak in favor of a potential role of layer 6b on cortical activation but in the absence of knowledge concerning the correlative projections of the L6b FS cells, the mechanisms involved remain unknown.

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