Influence of a Subtype of Inhibitory Interneuron on Stimulus-Specific Responses in Visual Cortex

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Inhibition modulates receptive field properties and integrative responses of neurons in cortical circuits. The contribution of specific interneuron classes to cortical circuits and emergent responses is unknown. Here, we examined neuronal responses in primary visual cortex (V1) of adult Dlx1−/− mice, which have a selective reduction in cortical dendrite-targeting interneurons (DTIs) that express calretinin, neuropeptide Y, and somatostatin. The V1 neurons examined in Dlx1−/− mice have reduced orientation selectivity and altered firing rates, with elevated late responses, suggesting that local inhibition at dendrites has a specific role in modulating neuronal computations. We did not detect overt changes in the physiological properties of thalamic relay neurons and features of thalamocortical projections, such as retinotopic maps and eye-specific inputs, in the mutant mice, suggesting that the defects are cortical in origin. These experimental results are well explained by a computational model that integrates broad tuning from dendrite-targeting and narrower tuning from soma-targeting interneuron subclasses. Our findings suggest a key role for DTIs in the fine-tuning of stimulus-specific cortical responses.

Keywords: cortex, dendrite, inhibition, orientation, tuning

Introduction

γ-Aminobutyric acid (GABA)ergic inhibitory interneurons are integral to cortical development and function. While these cells comprise merely 20–30% of neocortical neurons, they exhibit the greatest diversity in morphological, physiological, molecular, and synaptic characteristics (DeFelipe 1993; Markram et al. 2004; Blatow et al. 2005; Ascoli et al. 2008). Among different classes of interneurons, basket cells project to soma and proximal dendrites of excitatory neurons, chandelier cells innervate initial axon segment, whereas double bouquet, bipolar, bitufted, and Martinotti cells target dendrites and dendritic tufts (DeFelipe 1993; Markram et al. 2004). Furthermore, chandelier and typical basket cells express parvalbumin and are fast spiking; other basket cells express cholecystokinin (CCK) and/or vasoactive intestinal peptide (VIP), are adapting/regular spiking or irregular spiking, and fire action potentials with frequency adaptation (Kawaguchi and Kubota 1997, 1998; Kubota and Kawaguchi 1997; Wang et al. 2002; Galarreta et al. 2004); small basket cells express VIP and are regular spiking (Kawaguchi and Kubota 1997; Wang et al. 2002); dendrite-and tuft-targeting interneurons express somatostatin, neuropeptide Y (NPY), and calretinin and are accommodating (regular spiking) or for a small fraction irregular spiking (DeFelipe 1993; Chow et al. 1999; Markram et al. 2004; Toledo-Rodriguez et al. 2004; Xu et al. 2006; Gonchar et al. 2007).

Most GABAergic interneurons in the rodent neocortex derive from subpallial progenitors in the caudal or medial ganglionic eminences (CGE and MGE) (Marin and Rubenstein 2001; Wonders and Anderson 2006); parvalbumin+ and somatostatin+ interneurons originate in the MGE, while many calretinin+ and NPY+ interneurons are generated in the CGE (Xu et al. 2004; Butt et al. 2005). Several genes that encode homeobox transcription factors are critical for development of interneurons, including Nkx2.1 (Sussel et al. 1999; Butt et al. 2008), Lbx6 (Liodis et al. 2007; Zhao et al. 2008), and Dlx1 and Dlx2 (Anderson et al. 1997; Cobos et al. 2007). In particular, Dlx1 is required for survival of somatostatin+, NPY+, and calretinin+ interneurons—mice that lack Dlx1 show a selective reduction in these dendrite-innervating cortical interneurons (Cobos et al. 2005).

Given their diverse properties, different classes of interneurons likely have distinct roles in modulating integrative responses and receptive field properties of excitatory neurons. For example, GABAergic basket cells influence ocular dominance plasticity in visual cortex (Fagiolini et al. 2004; Yazaki-Sugiyama et al. 2009). Furthermore, parvalbumin+ interneurons are critical for generating gamma oscillations that may be important for cortical information processing (Cardin et al. 2009; Sohal et al. 2009). Inhibitory mechanisms involving distinct roles of interneuron subtypes may also underlie receptive field properties in visual cortex. A paradigmatic emergent property created by circuitry in the primary visual cortex (V1) is orientation selectivity (Somers et al. 1995; Ferster and Miller 2000). The role of inhibition in creating orientation selectivity is unresolved, with some studies claiming a critical role for intracortical inhibition in sharpening orientation tuning (Sillito 1975; Eysel et al. 1998), others indicating that thalamocortical excitation provides sufficient selectivity and inhibition has little role (Nelson et al. 1994; Ferster et al. 1996; Chung and Ferster 1998; Finn et al. 2007; Priebé and Ferster 2008), and still others arguing for a combination of thalamocortical plus intracortical excitation together with intracortical inhibition in shaping selectivity (Somers et al. 1995; Vidyasagar et al. 1996; Schummers et al. 2002; Shapley et al. 2003; Marino et al. 2005). Thus, we reasoned that it would be timely and instructive to examine whether the loss of dendrite-innervating interneurons results
in systematic alterations in orientation selectivity and other response properties of V1 neurons.

We have measured the effects of selectively reducing dendrite-innervating interneurons on stimulus-specific V1 responses in Dlx1−/− mice (Cobos et al. 2005)—due to their selective reduction of dendrite-innervating interneurons, these mice provide an opportunity to measure the effects of selectively reducing this subtype of interneuron on stimulus-specific V1 responses. We report that orientation tuned responses are broadened and firing rates are altered in Dlx1−/− mice, while other aspects of V1 physiology and thalamic inhibitory circuits appear normal. A computational model that captures our experimental findings indicates that different interneuron subclasses with specific response tuning can contribute to the generation of feature-selective responses in cortical circuits.

Materials and Methods

Animals

Mutant Dlx1−/− mice (Qiu et al. 1997) were maintained on a C57BL/6 × CD1 mixed genetic background to increase their viability (Cobos et al. 2005). Wild-type litters were used as controls. Mouse colonies were maintained at the Massachusetts Institute of Technology (MIT), the University of California, San Francisco (UCSF), and Stanford University in accordance with guidelines of the National Institutes of Health, MIT, UCSF, and Stanford, and all experiments were performed with approval from Institutional Animal Care and Use Committees. Dlx1−/− mice and their wild-type litters at ages of ≥postnatal days 60 were used in all experiments except in vitro slice recording.

Because Dlx1−/− mice show handle-induced seizures (Cobos et al. 2005), we attempted to avoid such seizure incidences by habituating the animals for 3–5 sessions (−10 min in each session, once a day) prior to the day of experiments. In these habituation sessions, mice were acclimated to the room and handling of the experimenter. The experimenter wore gloves, socialized with the mice, allowing them to walk or run from hand to hand, until the mice showed no apparent signs of stress or fear. We did not observe any seizure activity in Dlx1 mutant animals during the handling. These procedures helped minimize occurrence of seizures in the mutant mice: Very few mice displayed seizure episodes, as assessed by a 5-stage scale (Racine 1972), during the experiments described below. Mice that exhibited seizures were excluded from data analysis.

Histology and Cell Counting

Adult mice (−2 months old) were deeply anesthetized with Avertin (Sigma; 0.2 mL/10 g body weight) and perfused intracardially with 4% paraformaldehyde in phosphate-buffered saline (PBS) solution (pH 7.4). (Sigma; 0.2 mL/10 g body weight) and perfused intracardially with 4%

Experiments were performed in parallel for Dlx1+/+ and Dlx1−/− littermate mice, on at least 3–4 mice per genotype. Each experiment was repeated at least twice.

Nonstereological count methods were used to compare the number of neuronal bodies stained by immunohistochemistry or RNA in situ hybridization in mutant and control mice. The experimenter was blinded to genotype in all histological procedures, including sampling, data collection, and statistic analysis. Counts were performed on digitized images acquired using a Nikon Eclipse 80i microscope and Nikon Elements Software. At a magnification of 10×. Cell counting was performed on 2–3 coronal sections per animal in layers 2–3, 4, 5, and 6 of visual cortex, as well as in the nRT and dLGN. In all cases, samples from similar rostrocaudal levels in mutant and controls were selected. For quantitative in visual cortex, coronal sections were selected using stereotactic coordinates between bregma levels −2.92 and −1.20 mm (Paxinos and Franklin 2001). A 1-mm strip of cortex from the white matter to the pial surface was analyzed in each section (1.2 mm2 each). Values from the 2 trials performed on the same animal were averaged. The values from each mutant were normalized to their wild-type littermate controls. Percentage of change is presented as mean ± standard error of the mean (SEM). Statistical analysis was performed using the Student’s t-test (2-tailed unequal variance, α = 0.05) with Bonferroni corrections to calculate Pvalues.

In vitro Slice Preparation

Brain slices were prepared from Dlx1−/− and Dlx1+/+ mice (2–3 weeks and 1–2 months). Briefly, the mice were anesthetized with pentobarbital (50 mg/kg), decapitated, and the brains rapidly removed and immersed in an ice-cold (4°C) slicing solution containing (in mM): 234 sucrose, 2.5 KCl, 1.25 NaH2PO4, 0.5 MgSO4, 26 NaHCO3, and 11 glucose; equilibrated with 95% O2 and 5% CO2; pH 7.4. In the majority of older mice, the mice were first quickly perfused transcardially with 50 mL of the ice-cold slicing solution prior to decapitation. Horizontal thalamic slices (260 μm) containing the nRT and dLGN were cut and incubated in artificial cerebrospinal fluid containing (in mM): 126 NaCl, 2.5 KCl, 1.25 NaH2PO4, 0.5 MgCl2, 2 CaCl2, 26 NaHCO3, and 11 glucose; equilibrated with 95% O2 and 5% CO2; pH 7.4, initially at 32 °C for 1 h, and subsequently at room temperature, before being transferred to a recording chamber.

In vitro Patch Clamp Recordings

Thalamic neurons from nRT or dLGN were visually identified using infrared video microscopy. In addition, the identity of the cells was confirmed based on electrophysiological criteria related to the action potential firing of nRT cells and T-type current response of dLGN cells. Recordings were performed at room temperature in whole-cell configuration. Electrodes (tip resistance of 2–4 MΩ) were filled with internal solution containing (in mM): 120 K-glucionate, 11 KCl, 1 MgCl2, 10 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 1 ethyleneglycol-bis-2-aminoethyl ether)-N,N,N',N'-tetra acetic acid (EGTA) pH 7.3 adjusted with KOH, 290 mM osm for excitatory postsynaptic current (EPSC) recordings or 135 CsCl, 10 HEPES, 10 EGTA, 2 MgCl2, 5 QX-314 pH 7.3 adjusted with CsOH, 300 mM osm for inhibitory postsynaptic current (IPSC) recordings. For IPSC recordings, EPSC were isolated in 50 μM picrotoxin (Tocris), a GABAa receptor blocker, and for IPSC recordings, IPSCs were isolated in 20 μM 6,7-dinitroquinoxaline-2,3-dione (DQX), an amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) and kainate receptor antagonist and 50 μM 2-aminophosphonotanionic acid (AP5), a N-methyl-d-aspartic acid (NMDA) receptor antagonist (Ascent Scientific), all by bath application. The access resistance was always <18 MΩ and constantly monitored for stability.

In vitro Data Acquisition and Analysis

Signals were amplified with a Multiclamp 700A amplifier (Axon Instruments), low-pass filtered at 20 KHz and sampled at 4 or 10 kHz. A Digidata 1320 digitizer (Axon Instruments) and pClamp9 (Axon Instruments) were used for data acquisition. Locally written software (J. R. Huguenard, Stanford University School of Medicine) was used to detect (Widectecta, a program to detect postsynaptic current), sort, and
Mice were anesthetized using urethane (1.0 g/kg, intraperitoneally [i.p.]) and chlorpromazine (0.2 mg, i.p.), supplemented as necessary throughout imaging. During the experiment, the animal’s physiological condition was continuously monitored: temperature was maintained at 37–38 °C with a homeothermic heating blanket, the eyes were periodically lubricated with silicone oil, and the animal was continuously oxygenated. The skull was exposed and a custom-made plate was glued on the head to minimize movement. The skull was thinned over V1 in the same hemisphere across all animals with a dremel drill and covered with an agarose solution (1.5%) and a glass coverslip. The cortical surface was illuminated with red light (630 nm), and intrinsic signals were imaged with a charge-coupled device camera (Cascade 512B, Roper Scientific).

Intrinsic signal maps from a region of visual space (72° × 72°) were acquired during periodic stimulation (9 s/cycle) with a drifting horizontal or vertical white bar (9° × 72°) over a uniformly gray background, randomly presented to either eye (monocularly) or to both eyes simultaneously. Images of visual cortex were captured at 15 frames/s during each stimulus session of 25 min. A temporal high pass filter (135 frames) was used to remove slow noise components, after which the temporal Fast Fourier Transform (FFT) component at the stimulus frequency was calculated, pixel by pixel, from the whole set of images. The strength of visually driven responses for each eye was measured from the amplitude of the FFT component, and maps of response (ODI) were calculated at each pixel as ODI = (Rcontra − Ripsi)/(Rcontra + Ripsi), and ODI values from within the binocular zone were averaged for comparison across animals. The binocular zone was defined as the region with ipsilateral eye drive as well as contralateral drive. All pixels that had magnitude above 60% of the maximum level were considered responsive and thus included for analysis. To assess retinotopic map quality between wild-type and Dlx1Δ/− mice, a region of 50 × 50 pixels (900 μm²) was selected at a similar retinotopic location within the elevation map for each mouse. Scatter was then calculated as the difference in retinotopy between each pixel and the mean of the surrounding 25 pixels. Differences between wild-type and Dlx1Δ/− mice were compared using Student’s t-tests (2-sided unequal variance, based on normal distributions of the data and heterogeneous variance across groups), with the following exception: the P value for comparison of scatter between V1 and background from nonvisual cortical regions (see Fig. 1 legend) was calculated by a Wilcoxon rank sum test (2-sided, based on absence of normal distributions).

Optical Imaging of Intrinsic Signals

Single units were recorded with tungsten microelectrodes (2-4 MΩ resistance, FHC) as described (Dragoi et al. 2000). Briefly, mice were given atropine (0.54 μg/g) to decrease bronchial and salivary secretions and then anesthetized with Nembutal (pentobarbital, 0.05 mg/g), supplemented as necessary throughout recording. The body temperature of the animal was monitored by a rectal probe and maintained at 37–38 °C by a homeothermic heating pad. The animal’s eyes were covered with ophthalmic ointment until recording began, at which time the ointment was carefully wiped off and a thin layer of silicone oil was applied to the eyes. A metal plate was glued to the head of the animal; this plate was attached to a holder to maintain stability during the craniotomy and recording sessions. A dental drill and sharp forceps were used to perform a craniotomy to expose V1; the brain was covered with 2% agar.

Cluster sorting was performed using DataWave or Plexon software. Peristimulus time histograms (PSTHs) were constructed with 10-ms bins and averaged across all 16 conditions with visual stimuli at different orientations. Units were classified as visually responsive if their instantaneous firing rates (binned over 10-ms intervals) during the first 150 ms after stimulus onset were at least 4 standard deviations above the mean of prestimulus spontaneous firing rates (measured over 350 ms). Cells selected for detailed analysis had >1 Hz average evoked firing rates.
during the late response (defined below) at the preferred direction. Firing rate responses to each orientation, either maximal instantaneous at the initial transient peaks (10 ms average) or averaged over 250 ms of the late "steady-state" responses, were averaged from all trials, and the spontaneous firing rates were subtracted. Tuning curves based on these transient or late responses were generated by aligning the preferred direction of each cell to 90° and averaging these aligned responses. Orientation selectivity index (OSI) was calculated using a vector averaging method, and the preferred orientation was determined from the angle of the mean orientation vector (Swindale 1998; Dragoi et al. 2000). Values for OSIs, firing rates, and ratios (and also those from modeling in the following section) are presented as mean ± SEM. All P values were calculated using Wilcoxon rank sum tests (2-sided, α = 0.05) except for the following: P values for comparisons of individual data points (binned over 10 ms) on the PSTHs and for comparisons of tuning curves were calculated using 2-way analysis of variance (with genotypes and stimulus conditions as the 2 factors; α = 0.05).

Computational Model
We constructed a simplified cortical network model which was organized in a feed-forward sequence of 5 2-dimensional layers of cells. Within each layer, the preferred orientation of the cells varied along the first dimension between 0° and 180° and was constant along the second dimension.

There were 2 input layers and 3 processing layers representing 3 different types of cortical neurons. The first input layer provided stimulus-independent "background" (B) activity while the second layer represented stimulus-dependent inputs (I). Each input layer consisted of 512 cells. The 3 processing layers consisted of excitatory pyramidal cells, soma-targeting interneurons (STIs), and dendrite-targeting interneurons (DTIs), respectively. STIs were modeled as parvalbumin+ fast-spiking basket cells, while DTIs properties were consistent with somatostatin+, NPY+, and/or calretinin+, regular-spiking cells. The time constants for DTIs and STIs were tuned to match the physiological properties of fast-spiking versus regular-spiking GABAergic neurons. There were a total of 684 cells including 512 pyramidal cells (75% of total cells), 112 STI cells (15%), and 60 DTI cells (10%), matching the cortical proportions of these different cell types (Markram et al. 2004). Both B and I input layers provided excitatory inputs to all 3 processing layers. Both STI and DTI layers provided inhibition to pyramidal cells. Synaptic connections between layers were made such that each postsynaptic cell received input from presynaptic cells with similar orientation preferences.

Cell Parameters
Input (B and I) cells were modeled as independent variable rate Poisson process spike generators with a time step of 100 μs. Pyramidal (P) cells had a single somatic compartment with 6 dendritic compartments attached. The somatic compartment was based on the single compartment Hodgkin-Huxley type model (Golomb and Amitai 1997), and adapted to include additional dendritic compartments. The membrane potential was determined by the membrane equation summing currents from different intrinsic currents through channels present in the membrane with synaptic and injected currents.

For the soma:

\[ \frac{dV_m}{dt} = I_L + I_K + I_Na + I_Na \text{rev} + I_K + I_K + I_GABA \text{rev} + I_GABA \text{syn} + I_{inj} \]

With:

\[ C_m = 1.5 \mu F/cm^2 \]

Leak current:

\[ I_L = g_L (E_L - V_m) \]

\[ g_L = 0.20 \text{ ms}^2/cm^2, \ E_L = -65 \text{ mV} \]

Fast sodium channels:

\[ I_{Na} = g_Na (E_Na - V_m) m^b \]

\[ m = \frac{1}{1 + e^{-\frac{V_m - E_Na}{h}} \ b} \]

\[ \tau_m = 0.37 + \frac{2.78}{1 + e^{-\frac{V_m - 65}{h}}} \]

\[ \frac{dN}{dt} = \frac{m_n - n}{\tau_n} \]

\[ g_{Na} = 48.0 \text{ ms}^2/cm^2, \ E_{Na} = 55 \text{ mV}, \ \theta_n = -50.0 \text{ mV}, \ \sigma_n = 9.5 \text{ mV}, \ \theta_p = -53.0 \text{ mV}, \ \sigma_p = 7.0 \text{ mV}, \ \theta_s = -40.5 \text{ mV}, \ \sigma_{s} = 6.0 \text{ mV} \]

Persistent sodium channels:

\[ I_{NaP} = g_{NaP} (E_{NaP} - V_m) \]

\[ g_{NaP} = 0.14 \text{ ms}^2/cm^2, \ E_{NaP} = 55 \text{ mV}, \ \theta_p = -40.0 \text{ mV}, \ \sigma_p = 5.0 \text{ mV} \]

Delayed rectifier potassium channels:

\[ I_{Kd} = g_{Kd} (E_K - V_m) n^4 \]

\[ g_{Kd} = 0.14 \text{ ms}^2/cm^2, \ E_K = -90 \text{ mV}, \ \theta_n = -30.0 \text{ mV}, \ \sigma_n = 10.0 \text{ mV}, \ \theta_s = -27.0 \text{ mV}, \ \sigma_s = -15.0 \text{ mV} \]

A-type potassium channels:

\[ I_{A} = g_a (E_A - V_m) \]

\[ g_a = 28.0 \text{ ms}^2/cm^2, \ E_A = -90 \text{ mV}, \ \theta_a = -50.0 \text{ mV}, \ \sigma_a = 20.0 \text{ mV}, \ \theta_h = -80.0 \text{ mV}, \ \sigma_h = 6.0 \text{ mV}, \ \tau_p = 15.0 \text{ ms} \]

Slow potassium channels:

\[ I_{K} = g_k (E_K - V_m) \]

\[ g_k = 0.80 \text{ ms}^2/cm^2, \ E_K = -90 \text{ mV}, \ \theta_s = -39.0 \text{ mV}, \ \sigma_s = 5.0 \text{ mV}, \ \theta_h = 75.0 \text{ ms} \]

Synaptic currents:

\[ I_{syn} = g_{syn} (E_{syn} - V_m) \]

\[ E_{syn} = 0 \text{ mV}, \ E_{GABA} = -80 \text{ mV}, \ E_{AMPA} = -90 \text{ mV} \]

The dendrites were modeled analogously but only contained passive (leak) channels.

DTIs (D) were modeled as single compartments. These neurons usually have regular-spiking characteristics similar to pyramidal neurons (Beierlein et al. 2003; Ma et al. 2006; Caputi et al. 2008) and were thus modeled using the same set of equations, adjusted for the lack of dendritic compartments. \( g_l = 0.10 \text{ ms}^2/cm^2, \ g_{Na} = 48.0 \text{ ms}^2/cm^2, \ g_{NaP} = 0.14 \text{ ms}^2/cm^2, \ g_{Kd} = 6.0 \text{ ms}^2/cm^2, \ g_{A} = 28.0 \text{ ms}^2/cm^2, \ g_{K} = 0.20 \text{ ms}^2/cm^2 \).

STIs (S) show very different properties including fast spikes, higher firing rates, lack of spike rate adaptation (Beierlein et al. 2003; Ma et al. 2006).
were modeled as single compartments after the Hodgkin-Huxley type equations in a previous model (Wang and Buzsáki 1996):

\[
\frac{dV_m}{dt} = I_e + I_h + I_{K_D} + I_{syn} + I_{ij}
\]
\[
C_m = 1.5 \ \mu F/cm^2
\]

Leak current:

\[
l_h = g_h (E_h - V_m)
\]
\[
g_h = 0.40 \ \text{mS/cm}^2, E_h = -70 \ \text{mV}.
\]

Fast sodium channels:

\[
l_{Na} = g_{Na} (E_Na - V_m) m^3 h
\]
\[
g_{Na} = 0.60 \cdot (V_m + 35)/cosh(0.1 \cdot (V_m + 35)) - 1
\]
\[
\beta_h = 4c \ \frac{1}{cosh(0.1 \cdot (V_m + 35))}
\]
\[
m = m_s = \frac{\alpha_m}{\alpha_m + \beta_m}
\]
\[
\alpha_m = 0.07c \ \frac{(V_m + 40)}{cosh(0.1 \cdot (V_m + 40))}
\]
\[
\beta_m = 1 \ \cosh^{-1}(0.1 \cdot (V_m + 20)) + 1
\]
\[
g_{Na} = 70.0 \ \text{mS/cm}^2, E_{Na} = 55 \ \text{mV}.
\]

Delayed rectifier potassium channels:

\[
l_{K_D} = g_{K_D} (E_k - V_m) n^4
\]
\[
g_{K_D} = 0.1 \cdot (V_m + 34)/cosh(0.1 \cdot (V_m + 34)) - 1
\]
\[
\beta_n = 0.02c \ \frac{1}{cosh(0.1 \cdot (V_m + 34))}
\]
\[
m = m_s = \frac{\alpha_n}{\alpha_n + \beta_n}
\]
\[
\alpha_n = 0.125c \ \frac{1}{cosh(0.1 \cdot (V_m + 34))}
\]
\[
\beta_m = 1 \ \cosh^{-1}(0.1 \cdot (V_m + 20)) + 1
\]
\[
g_{K_D} = 180.0 \ \text{mS/cm}^2, E_k = -90 \ \text{mV}.
\]

Synaptic Dynamics

The model included 3 major synapse types, excitatory AMPA synapses and inhibitory GABA synapses including both GABA_A and GABA_B subtypes. All synapses were modeled by kinetic models with 2 states representing the open and closed states of ion channels (Destexhe et al. 1994). The fraction of open channels for a synapse \( r \) increases with the rate constant \( \alpha \) when neurotransmitter is present \((T = 1 \ \text{mM})\) and decreases with the dissociation rate \( \beta \):

\[
\frac{dr}{dt} = \alpha r (1 - r) - \beta r
\]

The total conductance \( g \) of a synapse was the product of its maximal conductance \( \hat{g} \) the fraction of open channels \( r \) and factor that determined the depression state of the synapse \( D \). Synaptic depression was modeled similar to the open channel fraction, adapted from a previous approach (Varela et al. 1997). In the nondepressed state, the full conductance of the synapse was available, represented by \( D = 1 \). Each time a synapse was active, its availability was decreased, governed by the rate constants \( \xi_D \) and \( \beta_D \):

\[
\frac{dD}{dt} = \xi_D T(1 - D) - \beta_D D
\]

The rate constants for the different synapse types (Destexhe et al. 1994) were:

- AMPA synapses: \( \alpha = 1.1 \times 10^4 \ \text{M}^{-1} \ \text{s}^{-1}, \beta = 190 \ \text{s}^{-1} \); GABA_A: \( \alpha = 0.53 \times 10^8 \ \text{M}^{-1} \ \text{s}^{-1}, \beta = 180 \ \text{s}^{-1} \);
- GABA_B: \( \alpha = 1.6 \times 10^8 \ \text{M}^{-1} \ \text{s}^{-1}, \beta = 4.7 \ \text{s}^{-1} \).

For depressing synapses: \( \alpha_D = 0.1 \times 10^5 \ \text{M}^{-1} \ \text{s}^{-1}, \beta_D = 4.0 \ \text{s}^{-1} \); for nondepressing synapses: \( \alpha_D = 0, \beta_D = 0 \).

Connectivity

Synaptic connections between layers were made such that each postsynaptic cell receives input from presynaptic cells with similar orientation preferences. The maximal distance between the orientation preferences of a pair of connected cells was determined by the input tuning. The standard input tuning for each connection in the model was 30°, that is, the maximal difference between the preferred orientation of the potential presynaptic cell and its postsynaptic target was 30°. Within this pool of potential presynaptic cells, the probability of an actual synaptic connection fell off with distance and was determined by a normal distribution with \( \sigma = 15° \).

All excitatory synapses were modeled as AMPA/kainate synapses. STIs made GABA_ergic synapses onto the soma of pyramidal cells. DTRs made both GABA_A- and GABA_B-ergic synapses onto the dendrites of pyramidal cells. For each pair of connected layers, the total synaptic conductance \( g \) was fixed as a parameter and the conductance of each individual synapse was computed by dividing the total conductance by the number of synapses for each postsynaptic neuron. The total conductances were: \( g_{AGA} = 3.0 \ \text{nS}, g_{GABA} = 12.0 \ \text{nS}, g_{SP} = 5.0 \ \text{nS}, g_{GABA} = 30.0 \ \text{nS}, g_{GABA} = 7.0 \ \text{nS}, g_{SP} = 40.0 \ \text{nS}, g_{GABAA} = 10.0 \ \text{nS}, g_{GABAD} = 0.8 \ \text{nS} \).

Inputs

For each B cell, the mean firing rate was fixed at a constant 10 Hz throughout the simulation. The firing rate of 1 cell varied with the stimulus. For cells with preferred orientations \( \theta \) and a stimulus with orientation \( \theta_s \), the firing rate of the cell was determined by a Gaussian curve centered on \( \theta_s \) with a width of 18°, baseline firing rate of 20 Hz and peak firing rate of 30 Hz:

\[
f_B = c \ \frac{e^{-\frac{(\theta_s - \theta)^2}{2 \ \sigma^2}}}{\sqrt{2 \pi \ \sigma}} \ \cdot f_0 + f_0
\]

with \( f_0 = 10 \ \text{Hz}, f_0 = 20 \ \text{Hz}, \) and \( \sigma = 18° \).

Simulation Environment and Global Parameters

All simulations were carried out using Cortical Network Simulator (Mutch et al. 2010) written in Matlab (The Mathworks) and running on a custom PC with 2 GeForce 8800 Ultra GPUs (EVGA). The time step for intracellular and synaptic computations was 10 \( \mu \)s, intercellular spike-based communication was performed with a time step of 100 \( \mu \)s.

Results

Subtype-Specific Reduction of Interneurons in Dlx1/Mice

Visual Cortex

We first examined whether there is subtype-specific loss of interneurons in the adult visual cortex of Dlx1/Mice, as reported in the somatosensory cortex of these mice (Cobos et al. 2005). Immunohistochemistry on the visual cortex of adult Dlx1/Mice mice showed a reduction in calretinin+ (27.7 ± 2.7%, \( P < 0.05 \)), NPY+ (30.2 ± 1.3%, \( P < 0.01 \)), somatostatin+ (16.1 ± 7.0%, \( P < 0.05 \)) neurons compared with wild types (Fig. 1A-G). The total number of parvalbumin+ neurons was unchanged in the mutant mice (Fig. 1G). These data are similar to those previously described in other cortex of Dlx1/Mice (Cobos et al. 2005). Furthermore, analysis by cortical laminar in the visual cortices of mutant mice and their control littersmates revealed that the reduction in calretinin+, NPY+, and somatostatin+ neurons occurred predominantly in layers 2–4 (Fig. 1G). Interestingly, while the total number of parvalbumin+ neurons appeared unchanged in the Dlx1/Mice visual cortex, there was an increase of parvalbumin+ neurons in layers 2/3 (27.6 ± 5.5%, \( P < 0.05 \)). Thus, we identified a subtype-specific reduction of interneurons, particularly in the superficial layers, in the visual cortex of Dlx1 mutant mice.

Normal Thalamic Function in Dlx1/Mice

Input to the visual cortex from the retina is first processed by the dLGN of the thalamus (Sherman and Spear 1982; Sherman...
Influence of an Interrneuron Subtype on V1 Responses

Dlx1 is expressed in the inhibitory neurons of the nRT (and its anlage, the ventral thalamus) and is not expressed in the dLGN (Bullone et al. 1993; Jones and Rubenstein 2004), we assessed whether there is loss of GABergic neurons in both the nRT and the dLGN of adult Dlx1−/− mice. The number of Gad67+ cells in these regions appeared normal in Dlx1−/− mice (Fig. 2A,B). We also examined the membrane and synaptic properties of nRT and dLGN neurons in Dlx1 mutants. We found no significant differences in these properties between Dlx1+/− mice and their wild-type littermates at adult or juvenile ages (Table 1). In particular, no difference was detected in the frequency or amplitude of sEPSCs recorded from nRT cells (Fig. 2C-F and Table 1). Similarly, we found no difference in the frequency or amplitude of sIPSCs recorded from dLGN relay neurons in Dlx1−/− mice compared with wild-type littermates (Fig. 2G,H and Table 2). Based on these observations, we did not find any evidence for altered properties of the thalamic relay to visual cortex in the Dlx1 mutant mice.

Intact Map Organization and Ocular Dominance in Dlx1−/− Mice

Given the reduction in dendrite-targeting cortical interneurons in Dlx1−/− mice, we next asked whether the overall organization of thalamocortical inputs to visual cortex is altered in these mice. We carried out intrinsic signal optical imaging in V1 of Dlx1−/− mice and their wild-type littermates and

Table 1

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<th>nRT</th>
<th>Passive membrane properties</th>
<th>Spontaneous excitatory postsynaptic currents</th>
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<tr>
<td></td>
<td>Capacitance (pF)</td>
<td>Input resistance (MΩ)</td>
</tr>
<tr>
<td>+/+ (adult; 10)</td>
<td>40.3 ± 5.8</td>
<td>330.2 ± 27.7</td>
</tr>
<tr>
<td>−/− (adult; 16)</td>
<td>40.9 ± 4.8</td>
<td>230.5 ± 34.4</td>
</tr>
<tr>
<td>+/+ (juvenile; 12)</td>
<td>56.3 ± 4.5</td>
<td>341.5 ± 36.8</td>
</tr>
<tr>
<td>−/− (juvenile; 19)</td>
<td>62.0 ± 3.8</td>
<td>342.2 ± 33.2</td>
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Note: No differences were observed in cell size (capacitance), input resistance or time constant, or in measure of synaptic excitation (as assessed via analysis of sEPSCs). Numbers in the first column indicate number of cells.
examined the retinotopic map organization, scatter of these maps, and distribution of eye-specific inputs (ocular dominance). None of these 3 properties differed significantly between the 2 mouse groups (Fig. 3).

Broadened Orientation Tuning in V1 of Dlx1<sup>−/−</sup> Mice

Next, we focused our investigation on orientation selectivity in V1 of Dlx1<sup>−/−</sup> mice and their wild-type littermates using single-unit recordings. We recorded 95 single units from 16 Dlx1<sup>+/+</sup> mice and 122 single units from 10 Dlx1<sup>−/−</sup> mice. In order to ensure reliable data and to minimize contamination by artifacts, we chose a conservative strategy for data selection: We eliminated cells with extremely low visually evoked spike counts, which can introduce statistical bias or error. Moreover, our results from laminar analysis of interneuron subtypes showed that reduction in dendrite-innervating interneurons are mostly restricted to the superficial layers in the visual cortices of mutant mice. Thus, we focused our analysis on units in layers 2 through 4; these included 62 of 95 units in Dlx1<sup>+/+</sup> mice and 91 of 122 units in Dlx1<sup>−/−</sup> mice. Of these units, 69.4% (43 of 62 units) in Dlx1<sup>+/+</sup> mice and 76.9% (70 of 91 units) in Dlx1<sup>−/−</sup> mice were responsive to visual stimuli. Among these visually responsive units, 61.4% (27 of 43 units) in Dlx1<sup>+/+</sup> mice and 62.9% (44 of 70 units) in Dlx1<sup>−/−</sup> mice had highly reliable evoked firing rates (> 1 Hz evoked firing rates, see Materials and Methods). All of the following analyses were performed on these cells.

We first examined whether temporal dynamics of visually evoked responses are affected in the Dlx1<sup>−/−</sup> visual cortex. PSTHs of Dlx1<sup>−/−</sup> units were distinct from those of Dlx1<sup>+/+</sup> units (Fig. 4A). The significantly different components of the Dlx1<sup>−/−</sup>-PSTHs included lower baseline firing rates prior to visual stimulus onset (Fig. 4B), lower transient response (single peaks with large amplitudes) to the stimulus onset (Fig. 4C) but higher firing rates in the late response to the stimulus (Fig. 4D). Thus, even a partial removal of dendrite-innervating interneurons alters the amplitude and dynamics of baseline and stimulus-driven responses.

<table>
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<th>Table 2</th>
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<tr>
<td>Properties of sIPSCs in Dlx1&lt;sup&gt;+/+&lt;/sup&gt; and Dlx1&lt;sup&gt;−/−&lt;/sup&gt; mice.</td>
</tr>
<tr>
<td>dLGN</td>
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<tr>
<td>Frequency (Hz)</td>
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<tr>
<td>+/+ (adult; 7)</td>
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<td>+/+ (adult; 6)</td>
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<tr>
<td>–/+ (adult; 6)</td>
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<tr>
<td>–/+ (adult; 7)</td>
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Note: No differences were observed in frequency, amplitude, or kinetics of sIPSCs in dLGN relay neurons. Numbers in the first column indicate number of cells.

Figure 3. Adult Dlx1<sup>−/−</sup> mice showed normal retinotopic maps and binocular responses in V1. (A) Intrinsic signal optical imaging in V1 (from the same hemisphere across all animals, as indicated by the cyan area in the illustration at right) showed that Dlx1<sup>−/−</sup> mice have normal retinotopy in response to a drifting horizontal or vertical bar displayed to the contralateral eye. As examples, elevation and azimuth maps from a wild-type and Dlx1<sup>−/−</sup> littermate are displayed as a function of amplitude, normalized to the maximum pixel response in the field of view. Scale bar: 500 µm. (B) Average pixel scatter of retinotopic maps did not significantly differ between wild-type (n = 3) and Dlx1<sup>−/−</sup> (n = 2) mice. For comparison, scatter from nonvisual cortical regions was significantly higher than scatter in V1 of wild-type and Dlx1<sup>−/−</sup> mice (P < 0.05). (C) ODI pixel histograms from representative mice showed similar distributions in wild-type and Dlx1<sup>−/−</sup> binocular cortex. Average ODI values from V1 of Dlx1<sup>−/−</sup> versus wild-type mice (0.36 ± 0.08 vs. 0.30 ± 0.02) were not significantly different. For all panels, *P < 0.05.
We next analyzed orientation tuning of V1 units in Dlx1+/− and Dlx1−/− mice. Based on our findings of altered response dynamics (described above), we separately analyzed orientation tuning for the early transient and late components of the visual response (for definition of early and late responses, see Materials and Methods). Tuning curves based on early transient response and late response were significantly different between Dlx1+/− and Dlx1−/− cells (Fig. 4A, early response: $P < 0.05$; Fig. 4F, late response: $P < 0.01$), likely due to differences in firing rates between the 2 curves. OSI values from the early response were significantly lower in the Dlx1−/− cells (Fig. 4G; Dlx1+/−, 0.13 ± 0.02; Dlx1−/−, 0.10 ± 0.01; $P < 0.05$). Similarly, OSIs from the late response in Dlx1+/− cells showed a trend toward lower values (Fig. 4H, Dlx1+/−, 0.20 ± 0.02; Dlx1−/−, 0.16 ± 0.02; $P = 0.06$). (Orientation tuning was less selective during the transient responses regardless of genotype. This likely reflects the contribution of an orientation-independent “onset transient” to these early responses. Tuning curves of late responses for individual cells from the top 20th percentile of OSI distributions in each of the Dlx1+/− and Dlx1−/− groups are shown in Supplementary Figs 1a and 1b, respectively). In sum, both early and late responses of V1 cells in Dlx1−/− mice are less orientation selective than cells in wild-type controls.
While the averaged evoked firing rates of the overall populations did not differ (Fig. 4E,F), there was a trend toward lower transient responses in Dlx1−/− mice (Dlx1+/−, 12.38 ± 1.44 Hz; Dlx1−/−, 11.21 ± 1.15 Hz; P > 0.05) and higher late steady-state responses (Dlx1+/−, 1.44 ± 0.23 Hz; Dlx1−/−, 2.31 ± 0.37 Hz; P > 0.05). The ratios of late responses to early responses were significantly higher in Dlx1−/− mice compared with Dlx1+/− mice (Fig. 4K; Dlx1+/−, 0.12 ± 0.01; Dlx1−/−, 0.20 ± 0.02; P < 0.05). Comparisons of peak evoked firing rates revealed similar results to those of averaged evoked firing rates (Supplementary Fig. 2A,B). Spontaneous firing rates of Dlx1−/− cells were significantly lower than those of Dlx1+/− cells (Fig. 4L; Dlx1+/−, 0.77 ± 0.12 Hz; Dlx1−/−, 0.48 ± 0.14 Hz; P < 0.05).

Furthermore, while examining the evoked firing rates, we noticed a clear divergence in the distribution of evoked firing rate differences (based on late responses) between wild-type and Dlx1−/− cells (Supplementary Fig. 3)—this divergence point was evident at the 30th percentile level of the distribution. Thus, approximately 30% of the cells (those with the highest evoked firing rates) seemed to be affected more severely than the rest of the population in the mutant mice. This breakpoint between the top 30% and bottom 70% of cells is consistent with the divergence point in the cumulative density function (CDF) in Figure 4J (indicated by an arrow).

We then examined in further detail the response properties of the 2 subgroups of cells in Dlx1−/− mice and compared each with the equivalent fraction of cells in Dlx1+/− mice. Interestingly, the top 30% and the bottom 70% of mutant cells showed drastically different changes in response properties (Fig. 5). For the top 30% of cells with the highest firing rates (Fig. 5A-C), it was the tuning curve based on late response that was more severely affected (early: P < 0.05; late: P < 0.01), whereas in the bottom 70% of cells (Fig. 5D-F), it was the tuning curve based on early response that was more affected (early: P < 0.01; late: P > 0.05). Spontaneous firing rates of the top 30% Dlx1−/− cells were significantly lower than those of Dlx1+/− cells (Fig. 5G; Dlx1+/−, 1.55 ± 0.20 Hz; Dlx1−/−, 0.91 ± 0.44 Hz; P < 0.05). The top 30% Dlx1−/− cells showed a trend toward higher evoked early response (measured by average firing rates; for similar results from analysis of peak evoked firing rates, see Supplementary Fig. 2) than those of Dlx1+/− cells (Fig. 5H; Dlx1+/−, 14.00 ± 3.29 Hz; Dlx1−/−, 16.19 ± 1.99 Hz; P > 0.05). As expected, the evoked late response of the top 30% Dlx1−/− cells was significantly higher than that of Dlx1+/− cells (Fig. 5I; Dlx1+/−, 2.18 ± 0.56 Hz; Dlx1−/−, 5.14 ± 0.84 Hz; P < 0.05). It is possible that some of our recorded cells are inhibitory neurons, but none of the putative fast-spiking cells identified based on their waveforms were in the top 30% population.

In comparison, spontaneous firing rates of the bottom 70% Dlx1−/− cells were significantly lower than those of Dlx1+/− cells (Fig. 5J; Dlx1+/−, 0.51 ± 0.11 Hz; Dlx1−/−, 0.29 ± 0.05 Hz; P < 0.05). These cells showed a trend toward lower evoked early response (Fig. 5K; Dlx1+/−, 11.66 ± 1.53 Hz; Dlx1−/−, 9.13 ± 1.23 Hz; P = 0.07) and lower late response (Fig. 5L; Dlx1+/−, 1.11 ± 0.17 Hz; Dlx1−/−, 1.12 ± 0.10 Hz; P > 0.05). We did not detect a significant difference between the OSIs of the top 30% Dlx1+/− and Dlx1−/− cells but did notice a trend toward lower OSIs in the bottom 70% Dlx1−/− cells (Supplementary Fig. 4). Thus, lack of Dlx1 results in changes in firing rates (especially during the late response) in the Dlx1−/− mice, and these changes particularly affect a subpopulation of cells.

Furthermore, we performed similar analysis on responses of cells in layers 5–6; these included 20 of 33 units (60.6%, from 16 Dlx1−/− mice) and 15 of 31 units (48.4%, from 10 Dlx1−/− mice) that were responsive to visual stimuli. Among these visually responsive units, 10 of 20 units (50%) in Dlx1−/− mice and 8 of 15 units (53.3%) in Dlx1−/− mice had reliably evoked firing rates (>1 Hz evoked firing rates, see Materials and Methods). Interestingly, compared with units in layers 2–4 (Fig. 4E,F and Supplementary Fig. 5A,B), only the tuning curves based on early transient response of cells in layers 5–6 were significantly different between Dlx1+/− and Dlx1−/− mice (Supplementary Fig. 5D,E, early response: P < 0.01), where Dlx1−/− units exhibited higher firing rates than Dlx1+/− units at all orientations. This is consistent with our observation that in layers 5–6 of Dlx1−/− mice, the numbers of calretinin+, NPY+, and somatostatin+ neurons are relatively unchanged (Fig. 1G and Supplementary Fig. 5F). In contrast, Dlx1−/− cells in layers 2–4 showed higher late response levels than Dlx1+/− units (Supplementary Fig. 5A,B), consistent with a reduction in calretinin+, NPY+, and somatostatin+ neurons and an increase in parvalbumin+ neurons in these layers (Fig. 1G and Supplementary Fig. 5C; see also Discussion). We did not detect significant differences in the other response properties, including OSI or evoked firing rates during early or late responses as well as spontaneous firing rates, between Dlx1+/− and Dlx1−/− units in layers 5–6 (data not shown).

Modeling the Contribution of Inhibitory Neuron Subtypes to Orientation Tuning

To further understand the changes in orientation selectivity and response level in Dlx1 mutant mice, we constructed a simple computational model that reproduced our experimental results. Specifically, we focused on building a model that captured the differential effects of the Dlx1−/− interneuron loss on early and late responses, as characterized by their firing rates and OSI values. The model predicts the corresponding tuning properties of different interneuron subtypes and provides quantitative insight into potential mechanisms underlying reduced orientation selectivity in the Dlx1 mutant mice.

We based our model on a feed-forward architecture (Fig. 6A) that consisted of 3 cell types: excitatory pyramidal cells, STIs, and DTIs. Because the majority of the interneurons lost in the Dlx1−/− cortex, namely somatostatin+, NPY+, and a fraction of calretinin+ cells, target dendrites of pyramidal cells, we did not explicitly include a fraction of calretinin+ cells that have been reported to target inhibitory neurons (Gonchar and Burkhalter 1999). All of these cells were modeled with Hodgkin–Huxley style dynamics, multiple compartments for pyramidal cells, and single compartments for inhibitory neurons, with some of the putative fast-spiking cells identified based on their waveforms in the top 30% population.

In comparison, spontaneous firing rates of the bottom 70% Dlx1−/− cells were significantly lower than those of Dlx1+/− cells (Fig. 5J; Dlx1+/−, 0.51 ± 0.11 Hz; Dlx1−/−, 0.29 ± 0.05 Hz; P < 0.05). These cells showed a trend toward lower evoked early response (Fig. 5K; Dlx1+/−, 11.66 ± 1.53 Hz; Dlx1−/−, 9.13 ± 1.23 Hz; P = 0.07) and lower late response (Fig. 5L; Dlx1+/−, 1.11 ± 0.17 Hz; Dlx1−/−, 1.12 ± 0.10 Hz; P > 0.05). We did not detect a significant difference between the OSI of the top 30% Dlx1+/− and Dlx1−/− cells but did notice a trend toward lower OSIs in the bottom 70% Dlx1−/− cells (Supplementary Fig. 4). Thus, lack of Dlx1 results in changes in firing rates (especially during the late response) in the Dlx1−/− mice, and these changes particularly affect a subpopulation of cells.
increased the impact of STIs onto pyramidal cells in the mutant model, based on the following rationale: 1) the number of parvalbumin+ cells was increased by 27.6 ± 5.48% in the superficial layers of Dlx1−/− visual cortex (Fig. 1G); 2) reduction in calretinin+ cells, some of which target GABAergic neurons in superficial layers (Gonchar and Burkhalter 1999), may result in decreased inhibition of inhibitory neurons, including STIs, in Dlx1−/− visual cortex; and 3) consistent with the above, spontaneous activity was lower in the Dlx1 mutant cells (Fig. 4L). To account for an increase in the number of parvalbumin+ cells and possible homeostatic compensation in their synaptic inputs (see Discussion), we chose a 50% increase in STI impact. Among the different combinations, the only one that showed reduced OSI values for both early and late responses in mutant mice, as indicated by our experimental data (Fig. 4G,H), contained STIs with narrower input tuning (with a maximum difference between presynaptic and postsynaptic preferred orientations of 30°, resulting in a mean OSI of 0.13) and DTIs with broader input tuning (with input tuning range of 90°, resulting in a mean OSI of 0.02) (Fig. 6A,C). Models with more sharply tuned dendrite-targeting inhibition led to changes in tuning that are less consistent with our experimental observations.

In addition, we also tested models with lower amounts of STI upregulation (e.g., 30% increased impact). These models replicated the increase in late response of mutant cells (this
The effect is robust and does not depend on the exact amount of STI upregulation, as long as DTIs are broadly tuned) but did not perform as well in replicating the changes in the spontaneous and early responses as the models with a 50% increase in STI impact.

In the best-fit model, tuning curves of the mutant cells showed changes that were highly consistent with our experimental results (compare Fig. 6D–G with Fig. 4E–H). In particular, the tuning curves based on late responses showed a major broadening and DC shift in the mutant model (Fig. 6D). OSI values calculated from early transient responses ($Dlx1^{+/+}$, 0.15 ± 0.003; $Dlx1^{-/-}$, 0.13 ± 0.003) and from late response ($Dlx1^{+/+}$, 0.34 ± 0.004; $Dlx1^{-/-}$, 0.29 ± 0.006) were both lower in the mutant cells (Fig. 6F–G). Similar to the experimental data, the mutant cells showed lower early responses ($Dlx1^{+/+}$, 11.81 ± 0.07 Hz; $Dlx1^{-/-}$, 10.77 ± 0.10 Hz) and higher late responses ($Dlx1^{+/+}$, 1.52 ± 0.01 Hz; $Dlx1^{-/-}$, 2.29 ± 0.05 Hz).

Furthermore, both the best-fit model and the experimental data showed a divergence point between the wild-type and mutant distributions of their late response firing rates. In the model results, this divergence point was located at ~50th percentile of firing rates (the location of the divergence depends heavily on the exact connectivity between DTIs and pyramidal neurons). Since the absolute value is not important for our argument, we did not attempt to fit the model to reproduce the exact 30%/70% split as in our experimental data. Instead, we used the same method of splitting the population at the divergence point to compare the model results to the experimental results.

**Figure 6.** A network model, in which we varied the orientation tuning of DTIs and STIs, explains the key results in $Dlx1^{-/-}$ mice and demonstrates the role of these interneuron subtypes in generating orientation selectivity and response levels of pyramidal neurons. (A) The basic architecture of our feed-forward model (see Materials and Methods). (B,C) After removing 33% of DTIs, changes in the OSI of pyramidal neurons during the early (B) and late responses (C) differed among 9 combinations of DTI and STI tuning properties. Numbers indicate ranges of differences (presented as ratios) between presynaptic and postsynaptic preferred orientations; the greater the number, the broader the tuning of the inhibitory neuron. Colors indicate directions (red: increase; blue: decrease; green: no change) and magnitudes (the darker the color, the greater the change) of changes in OSI. The only combination that produced lower OSI values during both early and late responses was the model with DTI = 90 and STI = 30 (both colors were blue). (D–G) In this best-fit model, tuning curves and CDFs for OSIs based on early (D,E) and late responses (F,G) showed similar changes to those in our experimental data (Fig. 4; see text for details). Tuning curves spanning 180° are presented because we did not include direction selectivity in our model. (H–O) The best-fit model displayed a divergence point of evoked firing rates at the 50th percentile. Changes in tuning curves (H–K) and CDFs for the firing rates (L–O) based on early (H,J,L,M) and late responses (H,K,M,O) of the top 50% of neurons (HJ and LM) and the bottom 50% of neurons (I,J and N,O) were similar to those in our experimental results (Fig. 5; see text for details). For all panels in D–O, blue lines depict wild-type cells; red lines depict mutant cells (after silencing one-third of DTIs).
experimen tal results). For the experimental data, we hypothesized that the 30% of Dlx1<sup>+/−</sup> cells with the highest late firing rates are cells that have lost most of their inputs from DTIs. We tested this hypothesis in the analysis of our model. We found that out of 512 cells, 264 (51.6%) did not lose any of their DTI inputs, whereas 197 (38.5%) lost at least 30% of their inputs. Over 90% (183/197) of these latter cells were in the top 50th percentile with the highest firing rates, thus supporting our hypothesis.

The model and experimental data were also in agreement with regard to changes in orientation selectivity and firing rates of these subpopulations of neurons (compare Fig. 6H–K with Fig. 5B, C, E, F and Fig. 6L–O with Fig. 5H, I, K, L). In the top 50% of cells with the highest firing rates, the tuning curves based on late responses were more severely affected (Fig. 6H, I), whereas in the bottom 50% of cells, the tuning curves based on early response were more affected (Fig. 6J, K). The top 50% mutant cells showed similar early responses to that of the control cells (Fig. 6L; Dlx1<sup>+/−</sup>, 11.87 ± 0.10 Hz; Dlx1<sup>+/+</sup>, 11.85 ± 0.13 Hz) but higher late responses (Fig. 6M; Dlx1<sup>+/−</sup>, 1.67 ± 0.01 Hz; Dlx1<sup>+/+</sup>, 3.36 ± 0.07 Hz). The bottom 50% Dlx1<sup>+/−</sup> cells exhibited lower early (Fig. 6N; Dlx1<sup>+/−</sup>, 12.25 ± 0.11 Hz; Dlx1<sup>+/+</sup>, 10.19 ± 0.11 Hz) and late responses (Fig. 6O; Dlx1<sup>+/−</sup>, 1.87 ± 0.01 Hz; Dlx1<sup>+/+</sup>, 1.72 ± 0.02 Hz).

Although the Dlx1<sup>+/−</sup> mouse represents a model with long-term loss of interneuron subtypes, we extended our modeling predictions to those from systems with acute silencing of one-third of the DTIs or STIs, in which the impact of STIs or DTIs, respectively, onto pyramidal cells remained unchanged (and therefore no homeostatic compensation was included). In these simulations, we used the combination of DTI and STI tuning properties in the best-fit model described above. Both manipulations resulted in increases in firing rates during early and late responses (Supplementary Fig. 6A, B and D, E). However, as shown by the ratios of late response firing rates to early response firing rates, while silencing STIs affected early and late responses to similar extents, silencing DTIs had a much stronger effect on late responses (Supplementary Fig. 6C, F). Furthermore, we observed this greater effect of DTI loss on late responses regardless of the tuning properties of DTIs and STIs: compared with an 8.8 ± 0.9% increase in the firing rate ratio after silencing STIs (averaged across all 9 combinations of DTI and STI tuning properties, as described above), silencing DTIs resulted in a 49.8 ± 1.6% average increase in this ratio (Supplementary Fig. 6G). Thus, DTIs may preferentially impact the sustained firing of target excitatory neurons, as suggested by our experimental data (Fig. 4A).

Discussion

We show that a subtype-specific reduction of dendrite-innervating interneurons in V1 leads to decreased orientation selectivity, lower spontaneous activity, and higher firing rates in late evoked responses. The effects on spontaneous and late evoked firing rates are more prominent in a subpopulation of neurons (~30%) with higher evoked firing rates than the remaining cells. Our computational model reproduces the experimental data and allows us to make predictions about possible tuning properties of DTI and STI interneuron subtypes. We suggest that the defects in V1 are due to intrinsic disruption of the visual cortical network, as no altered function of other components of the visual pathway was detected.

Role of Inhibition in Visual Receptive Field Properties

Inhibition has important functions in shaping ongoing and evoked cortical activity, and hence inhibitory interneurons have a key role in emergent properties of neuronal circuits (Mitchell et al. 2007; Chen et al. 2008). In the visual (and other sensory) cortex, inhibition appears to have a critical role in influencing the receptive field properties of neurons. The role of inhibition in shaping response features such as orientation selectivity (Ferster and Miller 2000) has been difficult to resolve unequivocally, due in part to multiple nonlinear properties (importantly, the spike threshold) available to cortical neurons for integrating inputs and generating selective responses (Priebe and Ferster 2008). Removal of inhibition leads to broadening of orientation and direction tuning (Sillito 1975; Sillito 1977; Jirmann et al. 2009), likely through mechanisms of network action (Somers et al. 1995).

The tuning of inhibitory inputs to an individual neuron is variable (Vidyasagar et al. 1996; Anderson et al. 2000; Monier et al. 2003; Shapley et al. 2003) and depends at least partly on the location of neurons within relevant feature maps (Nelson et al. 1994; Schummers et al. 2002, 2007; Marino et al. 2005). Even less is known about the role of inhibition in the mouse visual cortex, which lacks orientation columns (Drager 1974; Antonini et al. 1999; Hubener 2003) and may therefore require additional inhibitory mechanisms such as higher synaptic specificity for sharpening orientation selectivity. Thus, inhibition likely plays diverse roles via specific synaptic connectivity from different cell types, in order to shape receptive field tuning properties of excitatory neurons. In this report, our data on Dlx1<sup>+/−</sup> mice provide evidence for a specific role of inhibition from subtypes of interneuron in producing sharp orientation tuning. Importantly, we found an increase in the firing rate of late responses in the mutant mice. This suggests that dendritic inhibition has temporally specific effects on neuronal responses to visual stimuli. Still, in the Dlx1<sup>+</sup> mutant mice, the impact of a 30% reduction of a subset of interneurons on cortical responses is modest, perhaps due to compensatory mechanisms (see below), the possibility that orientation selectivity may be generated prior to V1, and/or that a 30% reduction is not enough to cause a stronger effect.

Role of Interneuron Subtypes in Modulating Visual Responses

Orientation tuning of inhibitory interneurons in mouse visual cortex has been described as generally broader than that of noninhibitory neurons (Sohya et al. 2007; Niell and Stryker 2008; Liu et al. 2009; Kerlin et al. 2010). However, careful analysis of parvalbumin-expressing interneurons using targeted recordings from single marked cells demonstrates that they have a range of tuning properties, including very sharp orientation selectivity (Runyan et al. 2010). Intracellular recordings from cat V1 neurons also indicate diversity in electrophysiologically defined interneuron classes (Azouz et al. 1997). Thus, a subgroup of fast-spiking neurons, which are soma targeting, has been described as having broad orientation tuning (Nowak et al. 2008), although synaptic inputs to fast-spiking neurons in layer 4 are as narrowly tuned as those to regular-spiking neurons (Cardin et al. 2007). Clarifying properties and functional roles of interneuron subclasses will require more specific molecular-genetic markers, and refinement of technologies for selective removal of specific
subclasses (Luo et al. 2008). Our model predicts broader tuning for DTIs than for STIs, which is at least partly consistent with findings from soma-targeting parvalbumin-expressing interneurons (Runyan et al. 2010). We also show that DTIs may have a precise role in sharpening orientation selectivity of neurons in V1.

The different subclasses of interneurons likely modify pyramidal cell responses in functionally precise and distinct ways. Previous studies in the somatosensory cortex have characterized the synaptic dynamics of DTIs in vitro (Kapfer et al. 2007; Silberberg and Markram 2007) and the involvement of this interneuron subclass in gain modulation of a sensory response (Murayama et al. 2009), but the contribution of dendritic inhibition to feature-specific sensory information processing had remained unknown.

In visual cortical processing, DTIs may contribute specific inhibitory subfields to excitatory neuron receptive fields, whereas STIs may contribute broad complex-cell-like inhibition (Hirsch et al. 2003; Lauritzen and Miller 2003; Palmer and Miller 2007). Our results suggest that the 2 major interneuron classes (DTIs and STIs) both sharpen orientation tuning, but the nature of their roles may be different, depending on their tuning properties and the location of their synapses. Based on predictions from our computational model, we postulate a possible mechanism that may account for our experimental data: STIs provide inhibitory input to the early response of the pyramidal neurons, are more narrowly tuned, and achieve sharpening of the early response by suppressing predominately the peak response to preferred orientations, DTIs provide inhibitory input to the late response, are more broadly tuned, and achieve sharpening of the late response by suppressing response to the preferred as well as nonpreferred orientations. In the mutant, a decrease in DTI inhibitory input leads to an overall (DC) increase in firing rates of the late responses, and hence a lower OSI calculated from the late response. Because input from STIs is increased, the peak response to preferred orientations will be selectively decreased relative to the responses to nonpreferred orientations, resulting in a lower OSI calculated from the early response.

**Time Course of Responses**

Our experimental observations and computational predictions provide convergent evidence that supports a model in which DTIs are critical for shaping the late responses of neurons to a stimulus: in layers 2–4 of Dlx1 mutant mice, the most prominent phenotype is an increase in evoked late response of all cells, while the initial transient response is only moderately reduced. This can arise from 2 possible but not mutually exclusive mechanisms. First, the observation that the main effect is on late sustained responses could indicate that the early transient response is the result of the input and not affected by local processing, whereas sustained activity is the result of local (intralayer) computations and hence more sensitive to loss of interneurons. Second, this is consistent with the observation that fast-spiking parvalbumin+ STIs, which likely exert the initial inhibition of the evoked response, show a small increase in number in superficial layers of the mutant visual cortex, whereas the slower-spiking DTIs, which may be critical for suppressing the late response, are partially lost. Furthermore, in layers 5–6 where DTIs are unchanged (but STIs may be slightly increased), only the evoked early response is increased. The distinct temporal effects of the 2 major interneuron subtypes are consistent with their intrinsic properties and synaptic connections. Fast-spiking parvalbumin+ STIs have membrane time constants that are comparable to or faster than pyramidal cells, while several subtypes of DTIs such as calretinin+ and somatostatin+ interneurons have longer membrane time constants (Beierlein et al. 2003; Ma et al. 2006; Cardin et al. 2007; Caputi et al. 2008). In contrast to the synapses onto or originating from parvalbumin+ cells, which are strongly depressing, synapses involving dendrites are often facilitating (Beierlein et al. 2003; Caputi et al. 2008). Furthermore, the location of synapses on the postsynaptic neuron supports a rapid action of STIs compared with a slower, delayed impact from DTIs.

Another line of support for these distinct temporal roles of interneuron subtypes arises from different postsynaptic GABAergic receptor types. GABA_A-mediated inhibition predominantly suppresses the early transient response, while GABA_B receptors are involved in inhibition of the later sustained response (Pfleger and Bonds 1995; Allison et al. 1996). Several studies have suggested a differential distribution of GABA_A and GABA_B receptors in pyramidal neurons: GABA_A receptors are localized both at the soma and in dendrites, while GABA_B receptors may be restricted to dendrites (Newberry and Nicoll 1985; Connors et al. 1988; Kang et al. 1994). Moreover, GABA_A and GABA_B synapses may originate from distinct subpopulations of interneurons (Segal 1990; Otis and Mody 1992; Sugita et al. 1992, Benardo 1994), although a class of GABAergic interneurons, the neurogliaform cells, in contrast to other GABA-releasing cells, elicits combined GABA_A and GABA_B receptor–mediated responses (Tamas et al. 2005). Nonetheless, neurogliaform neurons express NPY (Karagiannis et al. 2009), which are among the reduced interneurons in the Dlx1+/− cortex (Cobos et al. 2005). Together, these findings present possible mechanisms underlying our model, that slow dendritic GABA_B-mediated inhibition from DTIs (or a subclass thereof) is critical for the specific suppression of late evoked responses.

Furthermore, Dlx1−/− neurons exhibit lower baseline firing rates (CA1 neurons in Dlx1 mutants also exhibit reduced sEPSC amplitudes; D Jones, S Baraban, personal communication). We speculate that this decreased baseline activity may be attributable to a small increase in parvalbumin+ neurons in the superficial layers where most of the reduction in calretinin+, NPY+, and somatostatin+ neurons occurs and/or decreased disinhibition resulted from loss of the fraction of calretinin+ interneurons that innervate other interneurons in layer 2/3 of the cortex (Meskenaite 1997; Gonchar and Burkhalter 1999; Caputi et al. 2008). We found a distinct stratification in our data such that 30% of cells have higher firing rates and show much higher late responses, whereas the other 70% of cells have lower firing rates and are affected only in their early transient response. This result suggests that individual excitatory cells may have lost proportionally more, or less, DTI input, and as a result, may have experienced more, or less, homeostatic compensation from their remaining STI inputs. It is likely that one interneuron innervates multiple neurons, and one neuron receives inputs from multiple interneurons. Nevertheless, the similarity in the proportion of most affected cells (30%), and the proportion of DTI neurons lost (30%) in the mutant mice, suggests the possibility that the subset of lost DTI neurons had a specific target population of excitatory neurons, perhaps due to sparse distribution and small dendritic trees of these DTIs.
This hypothesis can be tested with future technical innovations, such as postnatal conditional mutagenesis of Dlx1, or strategies to selectively inactivate specific interneuron subtypes with optical-genetic tools.

Supplementary Material
Supplementary material can be found at: http://www.cercor.oxfordjournals.org/

Funding
National Institute of Health (EY007023 and EY018648) to M.S., National Eye Institute Kirschstein-NRSA Fellowship 1 F32 EY017243 to R.M., National Eye Institute Kirschstein-NRSA Fellowship 1 F32 EY018065 to A.V.W., NS34774, NS39579, and NS06477 to J.R.H., and MH049428 to J.L.R.R.); Nina Ireland Fund (to J.L.R.R.).

Notes
We thank Jaimie Chia Min Lee and Georgia Panagiotakos for their assistance with histological analyses, Jitendra Sharma, Caroline Runyan, and Scott Gorlin for technical assistance with single-unit recording, Daniela Tropea for advice on antibodies and immunohistochemistry, Orsolya Kuti, Bibek Karki, Travis Emery, Tim Lew, and Astra Bryant for helping in maintaining mouse colonies, Jim Mutch for providing the modeling framework, and Carla Shatz, Cristopher Niell, Scott Gorlin for technical assistance with single-unit recording, and Scott Gorlin for technical assistance with histological analyses, Jitendra Sharma, Caroline Runyan, and Scott Gorlin for technical assistance with single-unit recording.

We thank Jason Ritt, Beata Jarosiewicz, Hiroki Sugihara, Robert Haslinger, and Cortina McCurry for insightful discussions on data analysis and interpretation. We are grateful to Sonal Jhaveri for valuable comments on the manuscript. Conflict of Interest: None declared.

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