

Parallel Processing of Sensory Input by Bursts and Isolated Spikes

Anne-Marie M. Oswald,¹ Maurice J. Chacron,^{1,2} Brent Doiron,^{1,2} Joseph Bastian,³ and Leonard Maler¹

¹Department of Cellular and Molecular Medicine, University of Ottawa, Ottawa, Ontario K1G 2K4, Canada, ²Physics Department, University of Ottawa, Ottawa, Ontario K1N 6N5, Canada, and ³Department of Zoology, University of Oklahoma, Norman, Oklahoma 73019

Burst firing is commonly observed in many sensory systems and is proposed to transmit information reliably. Although a number of biophysical burst mechanisms have been identified, the relationship between burst dynamics and information transfer is uncertain. Electrosensory pyramidal cells have a well defined backpropagation-dependent burst mechanism. We used *in vivo*, *in vitro*, and modeling approaches to investigate pyramidal cell responses to mimics of behaviorally relevant sensory input. We found that within a given spike train, bursts are biased toward low-frequency events while isolated spikes simultaneously code for the entire frequency range. We also demonstrated that burst dynamics are essential for optimal feature detection but are not required for stimulus estimation. We conclude that burst and spike dynamics can segregate a single spike train into two parallel and complementary streams of information transfer.

Key words: burst; dendritic backpropagation; neural code; stimulus estimation; feature detection; sensory systems

Introduction

During sensory processing, peripheral input is processed so that relevant aspects of the sensory signal are encoded in the sequences of action potentials that constitute the neural code. In theory, if the stimulus input and the neural output are known, then the transformation between the two defines the coding mechanism. However, because of the complexity of most sensory input and the variability of neural responses, only a few of the mechanisms by which stimuli are represented in neural codes have been identified (Rieke et al., 1997; Eggermont, 1998; Lestienne, 2001).

Action potential sequences can consist of single spikes, high-frequency (HF) spike clusters known as bursts, or both. When coding for comparable signals, bursts, in comparison with single spikes, have been shown to improve the signal-to-noise (SNR) ratio (Sherman, 2001) and transmit specific or additional information about the stimulus (Reinagel et al., 1999; Keat et al., 2001; Kepecs et al., 2002; Martinez-Conde et al., 2002). Bursts also have been suggested to enhance the reliability of information transfer between neurons (Malinow et al., 1994; Lisman, 1997). However, because the behavioral significance of specific stimulus patterns is often unknown; it is difficult to determine whether bursts code for particular categories of relevant stimuli that are distinct from those coded for by single spikes (Krahe and Gabbiani, 2004).

Behaviorally relevant electrosensory inputs have been characterized and easily mimicked (Zupanc and Maler, 1993; MacIver et

al., 2001). Specifically, prey stimuli are low-frequency (LF) (<10 Hz) inputs, whereas communication stimuli are often high frequency (>40 Hz). The ability of electrosensory pyramidal cells to encode these stimuli depends on both the frequency content and the spatial extent of the stimulus (Bastian et al., 2002; Chacron et al., 2003). *In vivo* studies of electrosensory pyramidal cell bursting have revealed that spontaneous burst production varies with dendritic structure and spontaneous firing rate (Bastian and Nguyenkim, 2001), and stimulus-evoked bursts extract stimulus features better than single spikes (Gabbiani et al., 1996; Metzner et al., 1998). In addition, pyramidal cells can switch from a non-oscillatory to oscillatory and burst firing pattern contingent on spatial stimulus characteristics (Doiron et al., 2003a). However, the mechanism underlying burst production was not addressed in these studies.

In vitro studies have established that the biophysical burst mechanism of electrosensory pyramidal cells is dependent on the active dendritic backpropagation of somatic spikes (Turner et al., 1994; Lemon and Turner, 2000), and theoretical analyses also have characterized these burst dynamics (Doiron et al., 2001, 2002; Laing and Longtin, 2003; Laing et al., 2003). These previous studies suggest that the electrosensory system is an ideal preparation in which to elucidate the role of a biophysically and dynamically characterized burst mechanism in sensory coding.

This study addresses bursting and stimulus encoding in the weakly electric fish, *Apteronotus leptorhynchus*. Using both *in vivo* and *in vitro* experiments and an accurate burst model (ghost-burster), we demonstrated that bursts exclusively code for low-frequency stimuli, whereas isolated spikes code for high-frequency stimuli. We also demonstrated that optimal burst coding is dependent on burst dynamics. Thus, the electrosensory system creates two “streams” of frequency-specific information that could be differentially decoded by higher brain centers.

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Correspondence should be addressed to Anne-Marie Oswald, Department of Cellular and Molecular Medicine, University of Ottawa, 451 Smyth Road, Ottawa, Ontario K1G 2K4, Canada. E-mail: ammoswald@yahoo.com.

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Materials and Methods

In vivo. The preparation and surgical techniques for recording from *Apterionotus leptorhynchus* have been described previously (Bastian, 1996a,b) and were in accordance with the University of Oklahoma animal care and use guidelines. Stimulation consisted of random amplitude modulations (RAMs) of the animal's own electric organ discharge (EOD) produced by adding an amplitude-modulated train of sinusoidal signals (EOD mimic) to the animal's normal discharge. The RAMs were produced by multiplying the sinusoids with band-limited Gaussian noise (0–60 Hz bandwidth; eighth order Butterworth filter) and applied with local geometry (Bastian et al., 2002) using a small dipole (2 mm tip spacing) at a distance of 2–3 mm lateral to the skin surface. Extracellular and intracellular recordings of 33 pyramidal cells were obtained primarily from the centrolateral (CLS) and lateral (LS) electrosensory lateral line lobe (ELL) segments. Extracellular single-unit recordings were made with metal-filled electrodes, and intracellular recordings were made with 40–100 M Ω , KCl-filled micropipettes. Recordings were preamplified and acquired with Cambridge Electronics Design (Cambridge, UK) 1401plus hardware and Spike II software.

In vitro. Transverse brain slices from the ELL of *A. leptorhynchus* were prepared in accordance with animal care protocols of the University of Ottawa as described previously (Berman et al., 1997). The slices were maintained at room temperature (20–22°C) in oxygenated artificial CSF (ACSF) for 1–2 hr before recording. The ACSF contained (in mM): 124 NaCl, 3 KCl, 0.75 KH₂PO₄, 2 CaCl₂, 2 MgSO₄, 24 NaHCO₃, 10 D-Glucose. All chemicals were provided by Sigma (St. Louis, MO).

The pyramidal cell layer is located just below the stratum fibrosum (StF), an easily identifiable landmark in the ELL slice (Berman et al., 1997). Intracellular recordings from 37 ELL pyramidal cells from the entire pyramidal cell layer were made using 80–120 M Ω borosilicate glass electrodes, pulled by a Brown-Flaming P-87 puller (Sutter Instruments, Novato, CA), and filled with 2 M KAc. Of these cells, 24 were identified as bursting neurons by direct current injection and analyzed.

Recordings were obtained primarily from centromedial segment (CMS). Differences in oscillatory and burst activity across the ELL segments have been reported both *in vitro* (Turner et al., 1996) and *in vivo* (Metzner et al., 1998). Consequently, a comparison of 16 CMS, 6 CLS, and 2 LS *in vitro* pyramidal cell responses was conducted. The cells showed no significant differences in any parameters (such as firing rate, membrane potential, and coherence) studied, including bursting, and consequently the data were pooled.

After impalement, the neurons were allowed to stabilize for ~5–10 min. If necessary, a constant 0.1–0.5 nA hyperpolarizing current was injected through the recording electrode to stabilize the cells. A series of depolarizing current steps was injected to determine the current at which the firing rate was ~10–20 Hz (0.1–0.5 nA), which is within the range of spontaneous activity *in vivo*.

Previous studies have used Gaussian noise stimuli to stimulate the receptive fields of pyramidal cells *in vivo* (Bastian et al., 2002; Chacron et al., 2003; Doiron et al., 2003a). The membrane potential of pyramidal cells can follow these fluctuations (Chacron et al., 2003) presumably because electroreceptor EPSPs are brief (Berman and Maler, 1998). We therefore injected similar noise signals via the recording electrode into pyramidal cells *in vitro*. The noise stimuli were Gaussian noise with cutoff frequencies of 0–60, 0–20, or 40–60 Hz and SD of 0.5 nA, unless otherwise indicated. This was added to a 100 sec depolarizing current (0.1–0.5 nA, as determined above). Stimulation was a computer-controlled (IgorPro and Pulse Control; WaveMetrics, Lake Oswego, OR) (Herrington et al., 1995), and pyramidal cell responses were amplified (Axoclamp 2A; Axon Instruments, Burlingame, CA), sampled at 10 KHz, and stored for analysis (IgorPro and Pulse Control).

Drug application. Tetrodotoxin (TTX; 15 μ M in ACSF; Sigma) was focally pressure ejected in the ventral molecular layer above the Stf in a region corresponding to the apical dendrites of the pyramidal cells (Turner et al., 1994).

The model neuron. The previously described two-compartment model of an ELL pyramidal cell known as the ghostbuster model (Doiron et al., 2002) contains all the essential elements to reproduce bursting as detailed

by Lemon and Turner (2000) and Doiron et al. (2001). The model neuron is comprised of an isopotential soma (s) and a single dendritic compartment that are joined through an axial resistance of 1/gm_{s,d}, allowing for the electrotonic diffusion of currents from the soma to dendrite (d) and vice versa. Both compartments contain the essential spiking currents; fast inward Na⁺ ($I_{Na,s}$, $I_{Na,d}$) and outward delayed rectifying (Dr) K⁺ ($I_{Dr,s}$, $I_{Dr,d}$), and passive leak currents (I_{leak}). The presence of spiking currents in the dendrite enables the active backpropagation of somatic action potentials required for bursting. The membrane potentials at the soma, V_s , and the dendrite, V_d , are determined using a Hodgkin–Huxley-like formalism (Koch, 1999). The original ghostbuster model (Doiron et al., 2002) comprised six nonlinear differential equations; this study expands on this model to incorporate persistent sodium currents ($I_{NaP,s}$, $I_{NaP,d}$) and a slower activation of spiking sodium in the dendrite. The input current $I_s(t)$ delivered to the soma was comparable with the zero-mean Gaussian noise (0–60 Hz) that was added to constant depolarizing current I_0 and injected *in vitro*. $I_s(t)$ was produced by a fourth-order Butterworth filter, and the SD ($\sigma = 3.7 \mu$ A/cm²) of $I_s(t)$ and the intensity of the offset current I_0 (0.21 μ A/cm²) were adapted so that the firing rate (32 Hz) and burst fraction (0.24) were similar to the *in vitro* recording seen in Figure 1. V_s and V_d are described by the following equations:

$$C_m \frac{dV_s}{dt} = I_0 + I_s(t) + g_{Na,s} \cdot m_{\infty}^2 \cdot (1 - n_s) \cdot (V_{Na} - V_s) + g_{NaP,s} \cdot m_{\infty,NaP,s}^2 \cdot (V_{Na} - V_s) + g_{Dr,s} \cdot n_s^2 \cdot (V_K - V_s) + \frac{g_c}{\kappa} \cdot (V_d - V_s) + g_{leak} \cdot (V_1 - V_s), \quad (1)$$

$$C_m \frac{dV_d}{dt} = g_{Na,d} \cdot m_d^2 \cdot h_d \cdot (V_{Na} - V_d) + g_{NaP,d} \cdot m_{\infty,NaP,d}^2 \cdot (V_{Na} - V_d) + g_{Dr,d} \cdot n_d^2 \cdot p_d \cdot (V_K - V_d) + \frac{g_c}{1 - \kappa} \cdot (V_s - V_d) + g_{leak} \cdot (V_1 - V_d). \quad (2)$$

The parameter g is a maximal conductance (g_{max} , mS/cm²), whereas m and s are activation parameters, and h , n , and p are inactivation parameters. As described previously (Doiron et al., 2002), the activation of spiking Na⁺ in the soma, $m_{\infty,s}$, instantaneously tracks the membrane voltage, V_s , and its inactivation, h_s , is modeled as $h_s \approx 1 - n_s$, where n_s is the activation parameter for $I_{Dr,s}$ described by:

$$\frac{dn_s}{dt} = \frac{n_{\infty}(V_s) - n_s}{\tau_{n,s}}. \quad (3)$$

To simplify the ghostbuster model, the activation of dendritic spiking Na⁺ was originally slaved to the dendritic voltage as described for $I_{Na,s}$. This resulted in burst inter-spike intervals (ISIs) that were <3 msec, which were rarely seen experimentally. Immunocytochemical and electrophysiological data suggest that spiking Na⁺ channels are confined to active zones along the proximal apical dendrite. This type of channel distribution can lead to delays in depolarizing after potential (DAP) production that are not accounted for in the ghostbuster, resulting from the single compartment dendrite. Thus, we describe the activation of spiking Na⁺ in the dendrite, m_d , by:

$$\frac{dm_d}{dt} = \frac{m_{\infty,d}(V_d) - m_d}{\tau_{m,d}}. \quad (4)$$

This resulted in doublet ISIs that were 4–6 msec and closer to experimental values (5–8 msec).

The inactivation of spiking Na⁺ h_d in the dendrite, as well as the

activation n_d and the slow inactivation p_d of the dendritic K^+ currents, are modeled as described by Doiron et al. (2002):

$$\frac{dh_d}{dt} = \frac{h_{\infty,d}(V_d) - h_d}{\tau_{h,d}}, \quad (5)$$

$$\frac{dn_d}{dt} = \frac{n_{\infty,d}(V_d) - n_d}{\tau_{n,d}}, \quad (6)$$

$$\frac{dp_d}{dt} = \frac{p_{\infty,d}(V_d) - p_d}{\tau_{p,d}}. \quad (7)$$

Persistent sodium (NaP) currents may modulate bursting (Doiron et al., 2003b). Thus, for completeness and a better fit to experimental data, we incorporate NaP currents in both the soma and the dendrite as described by Doiron et al. (2001). This increased the excitability of model neuron and lowered bias current required to quantitatively reproduce *in vitro* results. For simplicity, we slave the activation of NaP, m_{NaP} , to the membrane voltage in both the soma and the dendrite.

The voltage dependence of each activation and inactivation parameter is described by steady state conductance curves as presented for the activation of somatic spiking Na^+ , m_{∞} :

$$m_{\infty}(V_s) = \frac{1}{1 + e^{-(V_s - V_{1/2})/k}}.$$

The values for g_{max} , τ , $V_{1/2}$, and k for each current are given in Table 1. Other parameter values are as follows: the ratio of somatic to total area, $\kappa = 0.6$; the reversal potentials, $V_{Na} = 40$ mV, $V_K = -88.5$ mV, $V_{leak} = -70$ mV; membrane capacitance, $C_m = 1 \mu F/cm^2$, $g_{leak} = 0.22$ mS/cm², $g_c = 0.21$ mS/cm². In some simulations, $g_{Na,d}$ was set to zero to remove active dendritic spiking and prevent bursting. This does not affect NaP conductance.

Equations 1–7 were integrated in time via a fourth-order Runge–Kutta approximation scheme with a time step of 2.5 μ sec. All simulations were 200 sec of simulation time.

Data analysis. *In vivo*, *in vitro*, and model results were analyzed using IgorPro and MATLAB (Mathworks, Natick, MA). The first second of the spike train was not analyzed to exclude transients resulting from the onset of the stimulus. We determined ISI criteria to identify burst ISIs on the basis of the valleys of a bimodal ISI histogram (10 msec; *in vitro*, model neuron) and a previously used criterion for deviations from Poisson statistics (Bastian and Nguyenkim, 2001) (12 ± 4 msec; *in vivo*). Generally, ISIs < 10 msec are in agreement with backpropagation-dependent burst ISIs reported *in vitro* (Noonan et al., 2003). Using the ISI criteria, the spike trains were separated into three different spike trains for analysis. The first, denoted “all spikes,” was the raw spike train that contained the timing of all spikes. The second contained the timing of only the first spike of the bursts, denoted “burst events.” The final was denoted “isolated spikes” and contained only the timing of spikes that were not burst spikes. The coherence $C(f)$ between the three individual spike trains and the stimulus was calculated according to $C(f) = [P_{ss}(f)]^2 / [P_{ss}(f)P_{xx}(f)]$, where P_{ss} and P_{xx} are the power (P) spectrums of the stimulus and the spike train, respectively, and P_{ss} is the cross-

spectrum between the stimulus and the spike train. Coherence takes a value between 0 and 1 and is a measure of the strength of the neural response to a stimulus at a given stimulus frequency (f).

Burst- and spike-triggered averages. We determined 400 msec regions of the stimulus centered on each isolated spike or burst event for the *in vivo* and *in vitro* spike trains to be the spike-triggered average (STA) and burst-triggered average (BTA), respectively. To assess the specificity of spikes or bursts, we reinjected 0.5 Hz trains of STAs or BTAs with an offset current of +0.1 nA into a small number of pyramidal cells *in vitro*.

Feature detection. We used the feature extraction technique described in detail by Metzner et al. (1998). Briefly, the spike train and stimuli were binned at $\Delta t = 0.5$ msec, so that no more than one spike occurred per bin. For each bin, the stimulus vector was determined as the stimulus waveform preceded the bin ($t - \Delta t$; t). These vectors were then grouped as distributions according to whether a spike occurred, $P(s/\lambda = 1_{all})$, a spike occurred and it was an isolated spike, $P(s/\lambda = 1_{isol})$, a spike occurred and it was the first spike of a burst, $P(s/\lambda = 1_{burst})$, or a spike did not occur (null), $P(s/\lambda = 0)$. The means (m_i), which are the spike, isolated spike, burst-, and null-triggered averages, respectively, as well as the covariances (Σ_i) of the distributions, were estimated. Fisher’s linear discriminant function was used to determine optimal feature vector (f) to maximize the SNR.

$$\frac{1}{2} (\Sigma_0 + \Sigma_1) \cdot f = m_1 - m_0,$$

$$SNR = \frac{\left[f^T \cdot (m_1 - m_0) \right]^2}{f^T \cdot \left(\frac{1}{2} \Sigma_0 + \frac{1}{2} \Sigma_1 \right) \cdot f}.$$

The probability of correct detection, P_D (correctly classifying a stimulus vector as eliciting or not eliciting a spike), and the probability of false alarm, P_{FA} (misclassifying a stimulus vector as eliciting or not eliciting a spike), can then be determined. The separations between each of the distributions $P(f^T \cdot s/\lambda = 1_{all})$, $P(f^T \cdot s/\lambda = 1_{burst})$, $P(f^T \cdot s/\lambda = 1_{isol})$, and null distribution $P(f^T \cdot s/\lambda = 0)$ were assessed using these techniques, and the receiver operating characteristic (ROC) curve (P_D vs P_{FA}) was plotted for each case.

Statistics. The statistical significances of the results were assessed using paired t tests.

Results

Pyramidal cell burst responses to noise stimuli applied *in vivo* and *in vitro*

The stimulus consisted of broadband Gaussian noise (0–60 Hz) presented as AMs of electrosensory input *in vivo* and intracellular current injection *in vitro*. The mean firing rates in response to the stimulus were similar *in vitro* (18 ± 4 Hz) and *in vivo* (22 ± 13 Hz), and the spike trains were qualitatively similar in that they consisted of isolated spikes as well as high-frequency clusters of two to four spikes (Fig. 1A1,B1, arrowheads). The majority of ISI histograms both *in vivo* and *in vitro* had initial sharp peaks at the shortest intervals (Fig. 1A2,B2) and could not be fit by a single exponential ($\chi^2 > 500$); thus the spike train statistics are non-Poissonian, suggestive of burst discharge.

Based on the ISI histograms (Fig. 1A2,B2), an ISI criterion was established to identify spikes belonging to bursts (Turner et al., 1994, 1996; Debusk et al., 1997; Metzner et al., 1998; Reinagel et al., 1999). A criterion ISI of 10 msec was chosen for *in vitro* data. Because *in vivo* pyramidal cells show a wide range of firing frequencies, criterion ISIs were individually chosen for each cell (see Materials and Methods). The mean ISI criterion *in vivo* was 12 ± 4 msec. The burst fraction, defined as the total number of spikes contained within bursts divided by the total number of spikes, was determined for each cell, and the means were 0.49 ± 0.16 *in*

Table 1. Model parameter values

Current	g_{max} (mS/cm ²)	$V_{1/2}$ (mV)	k (mV)	τ (msec)
$I_{Na,s}[m_{\infty,s}(V_s)]$	55	−40	3	NA
$I_{NaP,s}[m_{\infty,s}(V_s)]$	0.14	−58.5	6	NA
$I_{Dr,s}[n_s(V_s)]$	20	−40	3	0.39
$I_{Na,d}[m_d(V_d)/h_r(V_d)]$	3.1	−40/−52	5/−5	0.3/4.0
$I_{NaP,d}[m_{\infty,d}(V_d)]$	0.1	−58.5	6	NA
$I_{Dr,d}[n_d(V_d)/p_d(V_d)]$	35	−40/−65	5/−6	6/15

Both somatic (s) and dendritic (d) compartments contain the essential spiking currents; fast inward Na^+ ($I_{Na,s}/I_{Na,d}$) and outward delayed rectifying (Dr) K^+ ($I_{Dr,s}/I_{Dr,d}$) as well as persistent Na^+ ($I_{NaP,s}/I_{NaP,d}$). The parameters for the steady state conductance curves for each current are the maximum conductance (g_{max}), half-deactivation voltage ($V_{1/2}$), deactivation time constant (τ), and the deactivation slope (k). In the case of two entries under $V_{1/2}$, τ , and k , the first value corresponds to the activation parameter and the second value to the deactivation parameter. NA, Not applicable.

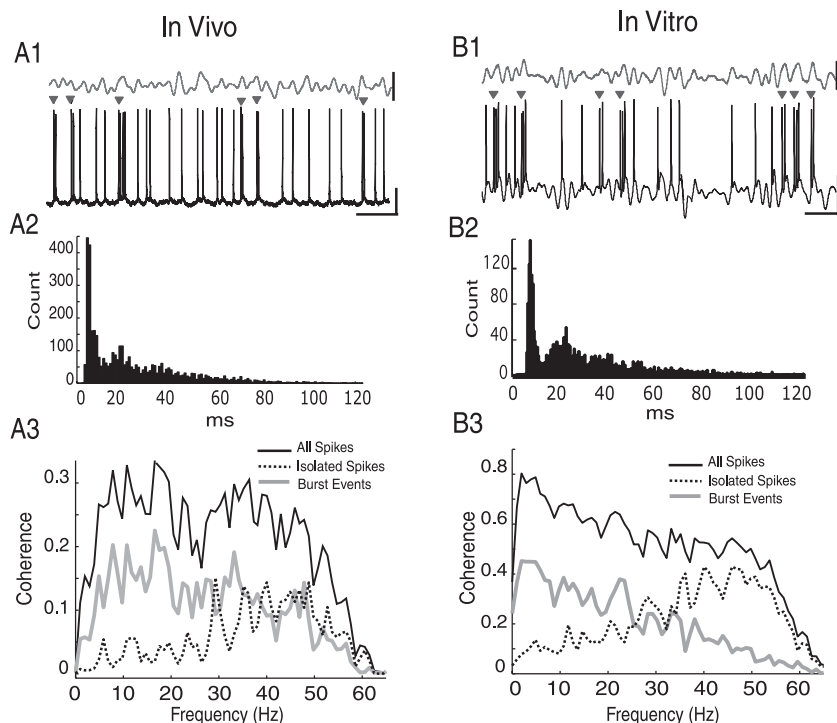


Figure 1. ELL pyramidal cell responses to broadband (0–60 Hz) Gaussian noise stimuli. *A*, A pyramidal cell response *in vivo* to a localized application of the broadband stimulus at the skin (firing rate, 40 Hz; burst fraction, 0.49). *B*, A pyramidal response *in vitro* to direct current injection of the broadband stimulus (firing rate, 33 Hz; burst fraction, 0.27). *A1*, *B1*, Spike trains in response to broadband stimuli (gray) consisted of isolated spikes and high-frequency bursts of spikes (arrowheads). *A2*, *B2*, ISI histograms with a significant peak in the 3–10 msec range that corresponds to the high-frequency spike events. *A3*, *B3*, Coherence plots between the stimulus and the neural response for the full spike train (black solid line) and the partitioned spike trains: burst events (gray line) and isolated spikes (dashed line). Calibration: (in *A*) stimulus, 0.18 mV/cm; spike train, 20 mV; 100 msec. Calibration: (in *B*) stimulus, 0.5 nA; spike train, 20 mV; 100 msec.

vivo and 0.22 ± 0.12 *in vitro*. The burst criterion was then used to separate the full spike train into two component trains: bursts, which consisted of the timing of the first spike in a burst event, regardless of the number of spikes in a burst, and isolated spikes, which consisted of the timing of the remaining single spikes.

respectively. In contrast, isolated spikes are on average less selective ($C_{low} \approx C_{high}$; *in vivo*, $p > 0.5$; *in vitro*, $p > 0.5$) with $C_{low}:C_{high}$ ratios of 0.7–1.0. We also investigated the frequency selectivity of bursts by injecting low-frequency (0–20 Hz) or high-

The coherence between each of the event trains (all spikes, burst events, and isolated spikes) and the stimulus was calculated (Fig. 1*A3*,*B3*). The mean coherence values between 0 and 20 Hz (C_{low}) and 40 and 60 Hz (C_{high}) were calculated to assess the frequency selectivity of the event trains. These and other measured values are reported in Table 2, and the results of statistical analysis are presented in the text. Previous *in vivo* studies (Chacron et al., 2003) have demonstrated that pyramidal cells give low-pass responses on stimulation of their receptive field center, and this was confirmed in the sample reported here ($C_{low} = 0.24 \pm 0.12$; $C_{high} = 0.10 \pm 0.097$; $p < 0.01$). Analysis of the *in vitro* full spike trains corroborated that the pyramidal cells were also low pass (*in vitro*, $C_{low} = 0.47 \pm 0.081$; $C_{high} = 0.308 \pm 0.08$; $p < 0.01$). The low-pass response can be appreciated in the ratios of $C_{low}:C_{high}$, which were 2.4 and 1.5, *in vivo* and *in vitro*, respectively.

Bursts and isolated spikes code for different signal components

Pyramidal cells were capable of responding to the entire stimulus frequency range with bursts or isolated spikes. However, both *in vivo* and *in vitro*, burst responses were significantly biased toward low-frequency stimulus events ($C_{low} > C_{high}$; *in vivo* and *in vitro*, $p < 0.01$) because the $C_{low}:C_{high}$ ratio *in vivo* and *in vitro* was 4.7 and 3,

Table 2. Summary of measured values *in vivo*, *in vitro*, and the model

Condition	<i>In Vivo</i>		<i>In Vitro</i>				Model	
	0–60 Hz	0–60 Hz	0–20 Hz	40–60 Hz	0–60 Hz		0–60 Hz	
					Control	TTX	ADM	PDM
Firing rate (spikes/sec)	22 ± 12.9	18 ± 4	16 ± 3	16 ± 2	18	18	32 Hz	33 Hz
Burst fraction	0.49 ± 0.16	0.22 ± 0.12	0.32 ± 0.19	0.008 ± 0.005	0.16	0.08	0.24	0.064
<i>n</i>	33	24	6	6	3	3	NA	NA
All spikes								
C_{low}	0.24 ± 0.12	0.47 ± 0.08	0.51 ± 0.09	NA	0.5	0.5	0.67	0.72
C_{high}	0.10 ± 0.10	0.31 ± 0.08	NA	0.60 ± 0.10	0.32	0.34	0.38	0.39
Burst events								
C_{low}	0.13 ± 0.08	0.13 ± 0.07	0.15 ± 0.07	NA	0.19	0.12	0.27	0.14
C_{high}	0.03 ± 0.03	0.04 ± 0.02	NA	0.02 ± 0.01	0.05	0.03	0.16	0.07
Isolated spikes								
C_{low}	0.08 ± 0.04	0.28 ± 0.08	0.42 ± 0.05	NA	0.25	0.33	0.19	0.47
C_{high}	0.08 ± 0.08	0.30 ± 0.14	NA	0.54 ± 0.13	0.29	0.33	0.25	0.319
BTA								
P_{low}	0.041 ± 0.055	0.20 ± 0.06						
P_{high}	0.009 ± 0.054	0.05 ± 0.01						
STA								
P_{low}	0.021 ± 0.054	0.06 ± 0.03						
P_{high}	0.010 ± 0.016	0.04 ± 0.01						

Stimuli were Gaussian noise ranging from 0 to 60, 0 to 20, and 40 to 60 Hz and were applied *in vivo*, *in vitro*, or in the model as indicated. The mean firing rate and burst fraction during stimulation are reported. In some cases, the stimulus was applied in the presence or absence of TTX *in vitro* or in the presence of active dendritic backpropagation (ADM) or the absence of backpropagation (PDM). Coherence between the stimuli and the full spike trains, burst trains, and isolated spike trains were measured. C_{low} and C_{high} correspond to the mean values of the coherence measured in the 0–20 and 40–60 Hz ranges, respectively. The power spectrums of the BTA and STA were obtained. P_{low} and P_{high} correspond to the mean power measured in the 0–20 and 40–60 Hz ranges, respectively. NA, Not applicable.

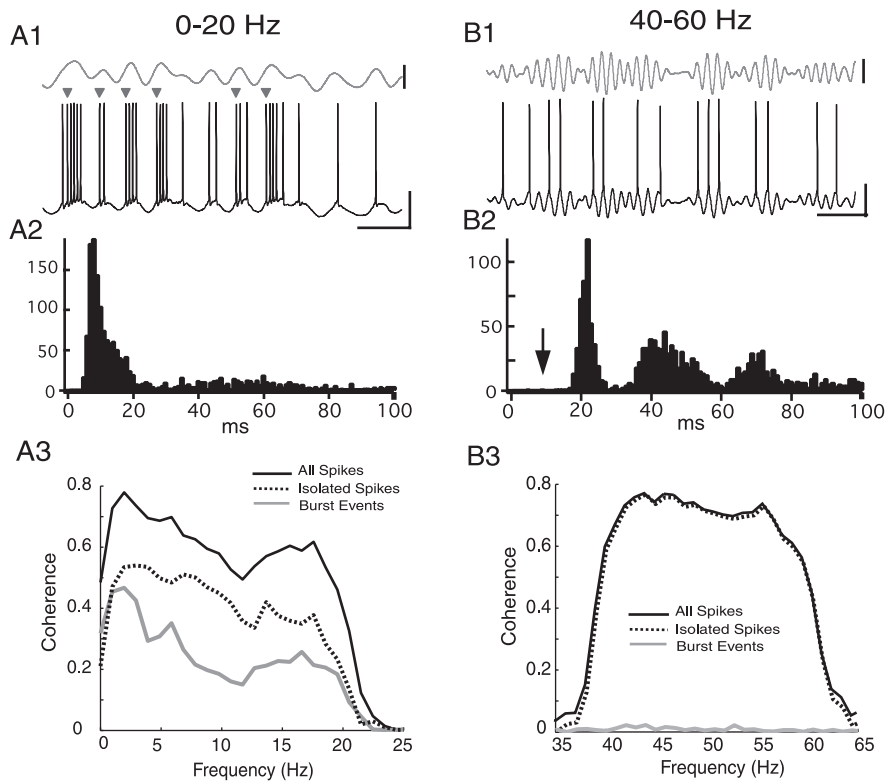


Figure 2. Pyramidal cell responses *in vitro* to low-frequency (0–20 Hz) versus high-frequency (40–60 Hz) Gaussian noise stimuli. *A1*, The pyramidal cell response (black) to direct current injection of a 0–20 Hz noise stimulus (gray) consisted of single spikes and high-frequency bursts of two to eight spikes (arrowheads). Firing rate, 20 Hz; burst fraction, 0.49. *A2*, ISI histogram is strongly skewed toward short 3–10 msec ISIs. *A3*, Coherence between the low-frequency stimulus and the neural response for the full spike train (black solid line) and the partitioned spike trains: burst events (gray line) and isolated spikes (dashed line). *B1*, The pyramidal cell response (black) to direct current injection of a 40–60 Hz noise stimulus (gray) consisted primarily of single spikes. Firing rate, 20 Hz; burst fraction, 0.00. *B2*, The ISI histogram is multimodal but lacks short 3–10 msec ISIs (arrow). *B3*, Coherence between the high-frequency stimulus and the neural response for the full spike train (black solid line) and the partitioned spike trains: burst events (gray line) and isolated spikes (dashed line). Calibration: stimulus, 0.5 nA; spike train, 20 mV; 100 msec.

frequency (40–60 Hz) noise *in vitro* (Fig. 2*A,B*, respectively). During LF stimulation, pyramidal cells responded with both bursts (2 ± 1 bursts/sec; burst fraction, 0.32 ± 0.19) and isolated spikes (10 ± 3 spikes/sec), whereas HF stimulation yielded almost exclusively isolated spikes (16 ± 2 Hz; burst fraction, 0.008 ± 0.005). These differential responses are apparent in the ISI histograms where LF stimulation resulted in bursts (Fig. 2*A1*, arrowheads) consisting of two to eight spikes corresponding to a peak at 3–10 msec (Fig. 2*A2*) that were absent during HF stimulation (Fig. 2*B2*, arrow). During LF stimulation, both bursts and isolated spikes were coherent with stimulus, although isolated spikes had higher coherence values (bursts: C_{low} , 0.15 ± 0.07 ; isolated spikes: C_{low} , 0.42 ± 0.05) (Fig. 2*A3*). In addition, isolated spikes were coherent with the HF stimulus (C_{high} , 0.54 ± 0.13) (Fig. 2*B3*). These results show that the isolated spikes can be coherent with either LF or HF stimuli and are therefore capable of stimulus estimation over the entire frequency range. Although bursts are selectively elicited by LF stimuli, the lower burst rate results in lower coherence values than those of isolated spikes. Thus, even at low frequencies, isolated spikes outperform bursts for stimulus estimation.

To characterize the signals that evoke bursts versus single spikes, we computed the BTA and STA (Fig. 3*A*). Both *in vivo* and *in vitro*, the power spectra of the BTA and STA are qualitatively

similar to the coherence curves for burst events and isolated spikes, respectively (Fig. 3*B*). The BTA has power predominantly at low frequencies ($P_{low} > P_{high}$; *in vivo*, $p = 6.8 \times 10^{-4}$; *in vitro*, $p = 1 \times 10^{-9}$), whereas the STA has significant power over the entire frequency range ($P_{low} \approx P_{high}$; *in vivo*, $p > 0.1$; *in vitro*, $p > 0.1$).

We then explored the dependence of bursts on the amplitude and temporal characteristics of the BTA *in vitro*. As expected, reinjection of BTA and STA waveforms resulted in bursts ($81 \pm 8.5\%$ of presentations) and spikes ($94 \pm 2\%$), respectively (Fig. 3*C*). The BTA was of significantly greater amplitude ($p < 0.01$) and half-width ($p < 0.01$) than the STA. In some systems, the BTA and STA are scaled versions of each other (Gabbiani et al., 1996; Reinagel et al., 1999), suggesting that increasing the stimulus amplitude would be sufficient to induce bursting. However, scaling the STA to the height of the BTA produced bursts in only $10 \pm 3\%$ of stimulus presentations. Alternatively, we doubled the SD of the 0–60 Hz stimulus. The spectral properties of the BTA and STA were conserved (data not shown), but the amplitude of the STA was increased to 0.6 nA, a value that surpasses the amplitude of the BTA under control conditions (0.45 nA). This very strong stimulus increased low-frequency burst coherence but had a minimal effect on burst coherence with high frequencies (ΔC_{high} , 0.037 ± 0.002 ; $\Delta C_{low} > \Delta C_{high}$; $p < 0.01$; $n = 14$). These results demonstrate that bursting is dependent on the temporal

characteristics of the stimulus as well as its amplitude; this conclusion is consistent with theoretical studies of electrosensory pyramidal cell burst dynamics (Laing and Longtin, 2003).

Bursts in ELL pyramidal cells therefore appear to selectively signal low-frequency components of sensory input, whereas isolated spikes are able to encode the entire frequency range. The bursts analyzed above were empirically defined from the ISI histogram, and we next explore their relationship to the dynamically defined ghostburst mechanism.

Backpropagation-dependent bursting

ELL pyramidal cells have a well characterized burst mechanism that is dependent on the active backpropagation of somatic spikes along the proximal apical dendrite (Lemon and Turner, 2000; Doiron et al., 2001, 2002). We used constant current injections (0.5–1.0 nA) *in vitro* to verify that backpropagation-dependent bursting can be elicited in all neurons studied. Figure 4 shows dendritic (*A*) and somatic (*B*) recordings *in vitro* in response to a constant depolarizing current injection. Somatic spikes backpropagate along the dendrite and broaden as the burst evolves. This is reflected at the soma as an increasing DAP (Fig. 4*B*, inset). Finally, DAP potentiation reaches the threshold for high-frequency doublet generation (Fig. 4*B*, arrow). However, the

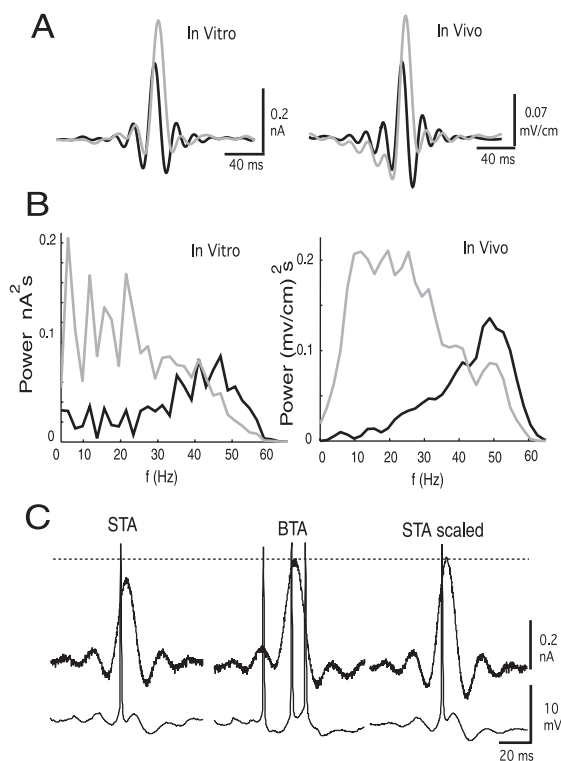


Figure 3. Burst- and spike-triggered averages are distinct stimulus features. *A*, BTA (gray) and STA (black) *in vitro* (left) and *in vivo* (right) were of significantly different height and duration (generated on the basis of the responses to 0–60 Hz stimulation of the cells presented in Fig. 1). *B*, The power spectrum of the BTA (gray) was significantly low pass, and that of the STA (black) was broadband to high pass both *in vitro* and *in vivo*. *C*, Rejection of the STA or BTA *in vitro* produced single spikes and doublets, respectively. Scaling the STA to the amplitude of the BTA did not elicit doublets in the majority of presentations.

doublet ISI falls within the dendritic spike refractory period halting the backpropagation of the second action potential of the doublet (Fig. 4*A*, asterisk), consequently terminating the burst. The doublet ISIs constitute a substantial peak in the 3–10 msec range of the ISI histogram (Turner et al., 1994; Doiron et al., 2003b). In addition, 0–60 Hz stimulation *in vitro* yields ISI histograms with a prominent peak in the 3–10 msec range (Fig. 1*B2*) and results in dendritic failures (Fig. 4*D*). Likewise, ISI histograms with a peak in a similar time range are observed in response to 0–60 Hz stimulation of a subset of pyramidal cells with high spontaneous firing frequencies (Fig. 1*A2*). Bursting and dendritic failure can also be elicited by step increases in EOD amplitude in some pyramidal cells *in vivo* (Fig. 4*C*). Thus, we propose that the high-frequency doublets present during 0–60 Hz stimulation are attributable to the backpropagation-dependent burst mechanism *in vitro*. This same mechanism is likely to underlie bursting in at least a subset of cells *in vivo*; however, additional studies are needed to determine whether the same mechanism gives rise to bursting in pyramidal cells with lower spontaneous firing frequencies (Bastian and Nguyenkim, 2001).

It has been demonstrated that the backpropagation of somatic action potentials along the dendrite is dependent on voltage-dependent sodium (Na^+) conductances (Turner et al., 1994; Doiron et al., 2003b). We focally applied TTX to the proximal apical dendrite of pyramidal cells *in vitro*. As demonstrated previously (Turner et al., 1994) this eliminated bursting induced by constant current (data not shown). TTX application also greatly reduced the number of high-frequency doublets elicited during

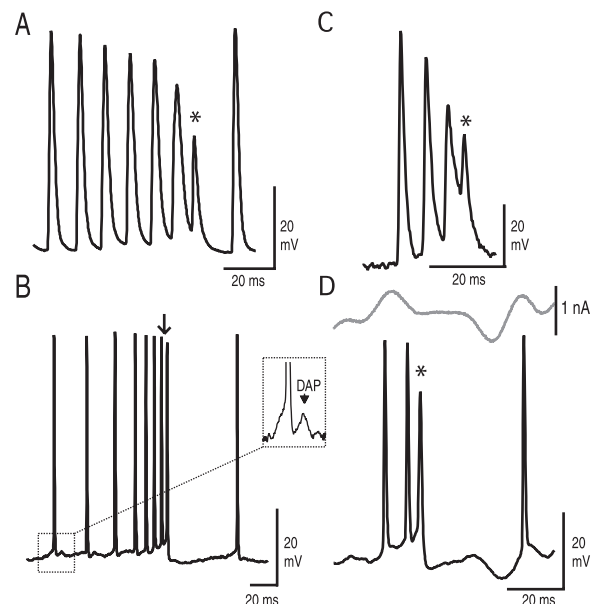


Figure 4. Evidence for backpropagation-dependent bursting *in vitro* and *in vivo*. *A*, Dendritic recording of a failure (*) of spike backpropagation elicited by step depolarization *in vitro*. *B*, Somatic recording of a backpropagation-dependent burst elicited by step current *in vitro*. Note the DAP resulting from a dendritic spike (inset) that leads to high-frequency doublet generation (arrow). *C*, Dendritic failure in response to a step change in the EOD amplitude *in vivo*. *D*, Dendritic failure in response to broadband noise stimulation (gray) *in vitro*.

0–60 Hz noise injection (Fig. 5*A*), and the ISI histogram became unimodal (Fig. 5*B*); note that the average firing rate did not change significantly. These results show that the bursts seen during broadband *in vitro* current injection are attributable to active dendritic processes.

Because bursts are low-frequency coherent, we hypothesized that preventing bursts during broadband current injection would decrease the low-frequency coherence of the full spike train. During TTX application, there was an expected reduction in burst coherence that coincided with an increase in low-frequency coherence for isolated spikes (Fig. 5*C,D*). However, surprisingly, there was not a substantial change in full spike train coherence (Fig. 5*C,D*) ($n = 3$). Contrary to our hypothesis, these results suggest that only the first spike of a burst is important for low-frequency coherence, and that bursts do not appear to code additional information about the stimulus.

Single spikes effectively code low-frequency stimuli

TTX applied to the apical dendrites eventually diffuses to the soma and blocks spiking; thus, there was only a limited time for data acquisition during which recordings were stationary and the somatic Na^+ was not affected. Therefore, we also investigated the role of backpropagation-dependent bursts in stimulus encoding using a two-compartment reduction of the detailed ionic model of an ELL pyramidal cell (Doiron et al., 2001, 2002). The model parameters and stimulation protocols (see Materials and Methods) were adapted so that the firing rate and burst fractions reported for the model were comparable with those presented *in vitro* (Fig. 1*B*).

Injection of 0–60 Hz Gaussian noise into the model neuron yielded similar results to those seen *in vivo* and *in vitro*. Spike trains at the soma contained both single spikes and bursts (Fig. 6*A1*, arrowheads). The majority of bursts were high-frequency doublets, which corresponded to failures in the dendrite (Fig.

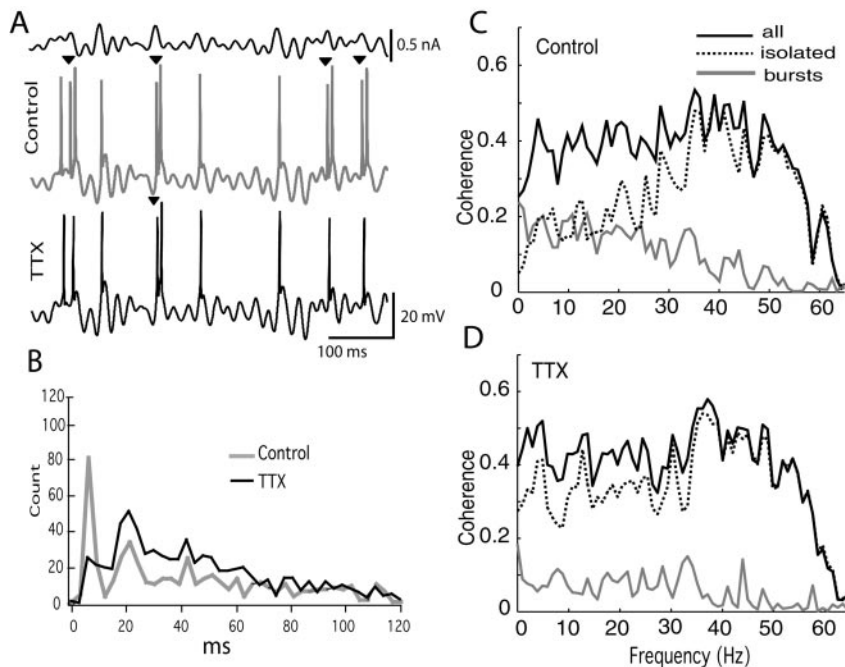


Figure 5. TTX blocks bursting but does not affect overall low-frequency coherence. *A*, Pyramidal cell response to a broadband stimulus (top trace) before (gray) and after (black) application of TTX to the apical dendrite. The number of bursts (arrowheads) is greatly reduced by TTX, which is reflected in the decreased peak in the 3–10 msec range of the ISI histogram (*B*), control (gray), and TTX (black). *C*, Coherence of bursts (gray), isolated spikes (black dashed line), and all spikes (black solid line) with the stimulus before application of TTX. *D*, In the presence of TTX, burst coherence decreased (gray), whereas isolated spike coherence (black dashed line) increased. The coherence of the full spike train did not change significantly (black solid line).

6A1, asterisks). The ISI histogram was bimodal with peaks at 3–5 and 20 msec (Fig. 6*B*, thin gray line). An ISI criterion of 6 msec, which corresponds to the trough of the ISI histogram, was chosen to distinguish burst spikes. Based on this criterion, the coherence between each of the event trains (all spikes, burst events, and isolated spikes) and the stimulus were calculated as described above. Similar to pyramidal cells *in vivo* and *in vitro*, the model neuron was low pass, and bursts were low-frequency coherent, whereas single spikes had a broadband response (Table 2; Fig. 6*C*, compare with Fig. 1).

We mimicked TTX blockade of active dendritic processes and prevented burst discharge by setting the dendritic, spiking Na^+ conductance ($g_{\text{max,Na,d}}$) to zero, effectively making the dendrite passive. From this point on, this will be referred to as the passive dendrite model (PDM), whereas the model with active dendritic processes intact will be referred to as the active dendrite model (ADM). The injection of a 0–60 Hz stimulus into the PDM results in an ISI histogram that still contains high-frequency ISIs (8–10 msec), but the bimodality and the highest frequency doublet ISIs (<6 msec) are lost (Fig. 6*A2,B*, thick black line). As with experimental blockade, there was a large reduction in burst fraction and low-frequency burst coherence (Table 2) (data not shown). Because burst events are reduced to single spikes, there is a substantial increase in the low-frequency coherence of isolated spikes (Fig. 6*D*, dashed lines). Consistent with experimental data, there was no substantial change in the coherence of the full spike train (Fig. 6*D*, solid lines). Thus, removing active backpropagation may result in a loss or delay of the second spike of the doublet, but this appears to be inconsequential, because the majority of information appears to be coded in the first spike of a burst.

Bursts and isolated spikes implement different coding strategies

Although single spikes provide better estimates of the detailed time course of the stimulus, it has been suggested that bursts may be more reliable indicators of specific stimulus features (Lisman, 1997). We used the Fisher linear discriminant (Gabbiani et al., 1996; Metzner et al., 1998) to evaluate the effectiveness of feature detection by burst events versus single spikes. Burst events were distinguished from single spikes on the basis of the previously described ISI criteria, and the following events were identified: all spikes (bursts plus isolated spikes), burst events, isolated spikes, and the null distribution (no bursts or isolated spikes). Stimulus features that corresponded to the 50 msec stimulus epoch that preceded each event were identified and then grouped into distributions according to the event elicited. An optimal feature was determined for each event distribution that maximized the separation between that event distribution and the null distribution. Each individual stimulus feature was then compared with the optimal feature and, on the basis of the degree of similarity, a prediction was made as to whether or not that event (single spike, burst, or null) would occur. The probability of correctly predicting an event versus

the probability of false alarm is plotted as the ROC curve.

Figure 7 shows the feature extraction of bursts versus single spikes in response to 0–60 Hz stimulation in representative pyramidal cells *in vivo* (Fig. 7*A*) and *in vitro* (Fig. 7*B*) as well as to 0–20 and 40–60 Hz stimulation *in vitro* (Fig. 7*C,D*). The optimal features (solid lines) detected by all spikes (black), bursts (thick gray), and isolated spikes (dashed) in response to each stimulus are presented in Figure 7*A2, B2, C2*, and *D2*. The distributions arising from the projection of the stimulus features onto the optimal features are shown in Figure 7*A3, B3, C3*, and *D3*; a reduced overlap between the event distributions (bursts, gray; isolated spikes, dashed), and the null distribution (black line) indicates better feature detection.

In response to 0–60 Hz stimulation, both *in vivo* and *in vitro*, bursts have higher SNR values (Fig. 7*A1,B1*, insets) than the full ($\text{SNR}_B > \text{SNR}_F, p < 0.01$) and isolated ($\text{SNR}_B > \text{SNR}_I, p < 0.01$) spike trains. In addition, isolated spikes have lower SNR values than the full spike train ($\text{SNR}_F > \text{SNR}_I, p < 0.01$). The SNRs *in vitro* are much higher than *in vivo* because of the much lower rate of spontaneous activity *in vitro* compared with *in vivo*. Comparable results to *in vitro* responses were also obtained for the same stimulus in the model neuron (data not shown). The optimal features are derived from the BTA (low pass) and STA (broadband), which differ in their frequency content (Fig. 3). In addition, bursts are coherent with the low-frequency content of the stimulus. For these reasons, we propose that bursts are detectors of low-frequency stimulus features. The situation for isolated spikes is more complex (see below).

Because the optimal features detected by bursts and isolated spikes during 0–60 Hz stimulation differ, we cannot directly assess the discrimination performance of bursts or single spikes

when they are coding similar frequency ranges. Thus, we directly compared the feature detection capability of bursts and isolated spikes for narrow bands of LF (0–20 Hz) and HF (40–60 Hz) stimuli. In response to LF stimuli, bursts have better SNRs than isolated spikes (SNR_B , 3.76 ± 0.23 ; SNR_I , 2.36 ± 0.26 ; $p < 0.05$) (Fig. 7C1, inset). The optimal feature detected by bursts is a scaled version of the isolated spike feature with the burst feature having a higher amplitude threshold for bursting (Fig. 7C2). Thus, bursts are more selective for their stimulus features than single spikes. Conversely, the lower threshold for isolated spikes leads to tonic firing that tracks the stimulus whenever the membrane potential is above threshold. This improves coherence and therefore stimulus estimation (Fig. 2) but reduces the difference between the optimal feature and the null feature, which hinders feature detection.

Alternatively, when presented an HF stimulus (40–60 Hz), isolated spikes have high SNR values (Fig. 7D1, inset). Although the lack of bursts prevents a statistical comparison between isolated spikes and bursts, in four of six cells, the SNR for isolated spikes was greater than for the full spike train, suggesting that rare instances of bursting may actually compromise high-frequency stimulus detection.

These results demonstrate that bursts, although not necessary for stimulus estimation, offer a significant advantage over isolated spikes for low frequency feature detection. Conversely, isolated spikes are better estimators of low-frequency stimuli. At high frequencies, isolated spikes could be considered as either feature detectors or stimulus estimators depending on the downstream decoding mechanism.

Burst dynamics are required for feature detection

The implication that bursts are more reliable feature detectors than isolated spikes is based on the hypothesis that sequential activation at a synapse by two or more spikes separated by short ISIs is more effective than when the spikes are separated by long ISIs (Lisman, 1997). This raises the question of whether short ISIs that are a consequence of biophysical burst mechanism provide an advantage for information transfer as compared with short ISIs that are not the result of a burst mechanism. We again used the ghostbuster model to investigate whether bursting dynamics are required to generate high-frequency multispike events that are selective for low frequencies. In Figure 8, three scenarios are presented in which we attempted to recreate the low-frequency selectivity and the feature extraction in the model neuron with a PDM to match that of the ADM. As shown previously, removing burst dynamics shifted the ISI histogram from bimodal (Fig. 8A1) to unimodal (Fig. 8A2–A4) distributions. Because HF events are no longer well defined, we identified subsets of HF events on the basis of ISI criteria (Fig. 8A2–A4, colored vertical

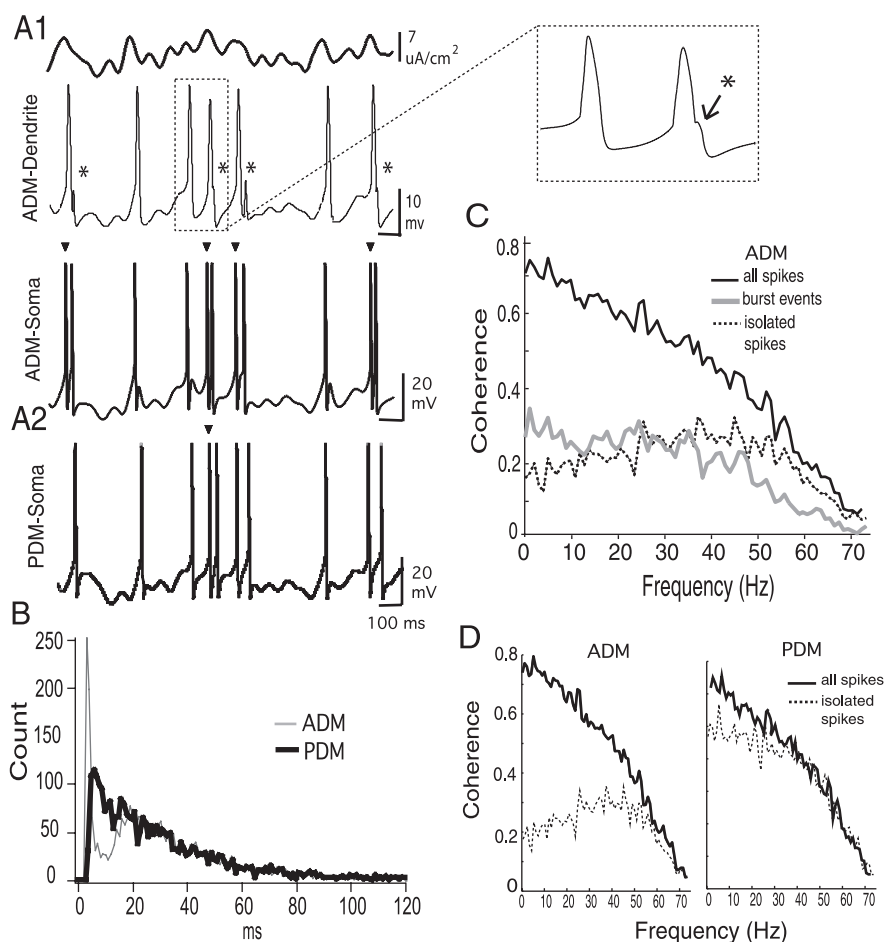


Figure 6. The model pyramidal cell recreates *in vitro* responses to broadband stimuli. *A*, Model response to injection of the broadband stimulus (top trace). *A1*, *A2*, Two cases are presented, one in which the model neuron has an active dendrite (ADM; *A1*) and one in which the dendritic Na^+ conductance is set to zero (PDM; *A2*). The ADM had a firing rate of 32 Hz, and somatic spike trains contained isolated spikes and high-frequency bursts (arrowheads; burst fraction, 0.24) that corresponded to dendritic failures (asterisks; enlargement). The PDM had a firing rate of 33 Hz but a burst fraction of 0.03. *B*, ISI histogram of the ADM (bimodal; thin gray line) with a significant peak in the 3–5 msec range that corresponded to burst events and the PDM (unimodal; thick black line). *C*, Coherence plots between the stimulus and the ADM response for the full spike train (black solid line) and the partitioned spike trains: burst events (thick gray line) and isolated spikes (black dashed line). *D*, Coherence of the full spike train (solid line) and isolated spikes (dashed line) with the stimulus in the ADM (left) and PDM (right). Calibration: $7 \mu\text{A}/\text{cm}^2$, 20 mV, 100 msec (unless otherwise noted).

lines) that resulted in the same HF event rate (5 Hz) or fraction (0.24) as indicated by the bimodal histogram of the ADM. In the first scenario, we changed the definition of an HF event from an $\text{ISI} < 6$ msec to an $\text{ISI} < 10$ msec, such that the mean firing rate and the HF event rate and fraction were the same as the ADM. The HF events were low pass (Fig. 8B; ADM, black; PDM 10 msec ISI, blue) but were poor feature detectors compared with true ADM bursts (Fig. 8C; control, black; 10 msec ISI, blue).

To ensure that reduced feature detection in the PDM was not attributable to the loss of ISIs < 6 msec, in the second and third scenarios we increased the offset current, I_0 , injected into the PDM and used 6 msec criterion to distinguish HF events (Fig. 8A3, A4). In the second scenario, the fraction of HF events with ISIs < 6 msec was comparable with the burst fraction in the ADM. The PDM required 18 times more current to generate the same fraction of ISIs < 6 msec as the ADM, which increased the mean firing rate to 80 Hz and the HF event rate to 15 Hz. This resulted in broadband HF event coherence (Borst and Haag, 2001) but very poor feature detection (Fig. 8B, C, red lines).

In the final scenario, the rate of ISI events < 6 msec was com-

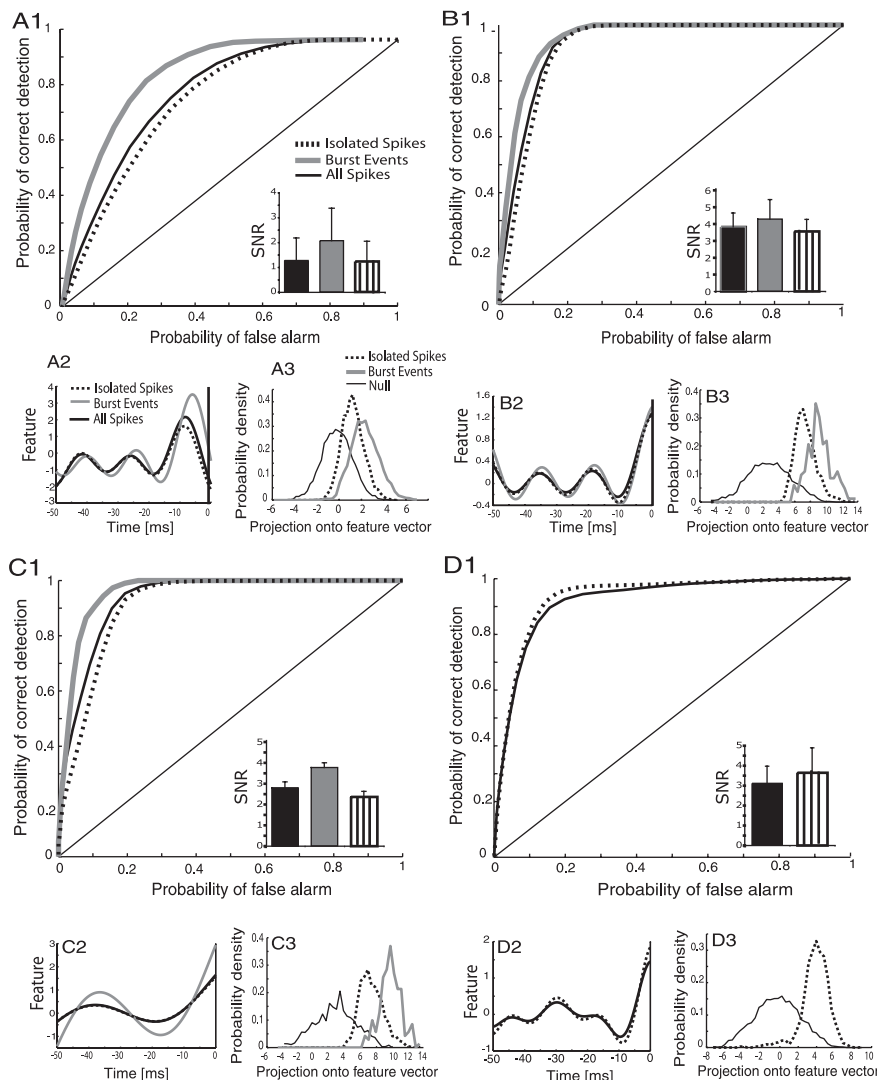


Figure 7. Feature extraction *in vivo* and *in vitro*. *A*, Analysis of spike trains in response to a broadband stimulus *in vivo*. *B*, Analysis of spike trains in response to a broadband stimulus *in vitro*. *C*, Analysis of spike trains in response to LF stimulation (0–20 Hz) *in vitro*. *D*, Analysis of spike trains in response to HF stimulation (40–60 Hz) *in vitro*. *A1–D1*, ROC curves for the partitioned spike trains: all spikes (black solid line), bursts (thick gray line), and isolated spikes (black dashed line). Inset, The mean SNR of all spikes (black solid line), bursts (gray line), and isolated spikes (black striped line). *A2–D2*, The optimal features detected by all spikes (black solid line), bursts (gray line), and isolated spikes (black dashed line). Note that extended structure of these features (less than -20 msec) may be attributable to the narrow-band nature of the stimulus set. This may induce long-range correlations in the stimulus waveforms that might not be causally related to spikes and not eliminated by spike-triggered averaging. *A3–D3*, Projection onto the optimal features, bursts (gray line), isolated spikes (black dashed line), and the null feature (black thin line).

parable with the burst rate in the ADM. Again, to produce an HF event rate of 5 Hz, the passive neuron required five times the current, resulting in a higher mean firing rate (50 Hz) than the ADM. These events were low-frequency coherent but also failed to recover the same degree of feature extraction as true bursts (Fig. 8*B, C*, green lines).

These results demonstrate that neurons with passive dendrites can generate HF events (ISIs < 6 or < 10 msec) that are coherent with low frequencies. However, in each instance, the optimal feature detected by the HF event is of lower amplitude at the time of the event ($t = 0$) compared with the ADM (Fig. 8*D*, black). This decreases the distinction between the distribution of features detected by HF events and the null distribution. When stimulus features are projected onto these optimal features, there is less distinction between the HF event distributions and the null dis-

tribution (Fig. 8*E*) leading to a higher probability of false alarm. The end result is a reduced ability to detect specific stimulus features, even when the high-frequency event rate is conserved. Thus, back-propagation-dependent pyramidal cell bursts are not only low-frequency selective, but the burst mechanism enables pyramidal cells to generate a distinct stream of high-frequency burst events that demonstrate superior feature extraction capability over nonburst HF events.

Discussion

We demonstrated both *in vivo* and *in vitro* that in response to broadband stimuli, ELL pyramidal cells generate a spike train that consists of two parallel information streams. Specifically, ELL pyramidal cells selectively detect low-frequency features of broadband Gaussian stimuli with spike bursts, whereas single spikes simultaneously estimate the entire stimulus frequency range.

Interestingly, the low-frequency coherence of pyramidal cell responses was not dependent on burst dynamics, because the coherence between full spike train and the stimulus was unaffected by the dendritic application of TTX *in vitro* and by removing dendritic Na^+ conductance in the ghostbuster model. Under these conditions, a low-frequency event triggers a somatic spike, which is coherent with the stimulus. However, this spike fails to back-propagate, and additional burst spikes do not occur. This demonstrates that the low-frequency coherence with the stimulus is carried in the timing of the first spike of the burst (Kepecs and Lisman, 2003). Because coherence is directly related to the linear estimation of a signal from a spike train (Gabbiani and Koch, 1998; Roddey et al., 2000), we conclude that bursts are not necessary for this estimate.

We quantified the selectivity of bursting neurons using a feature extraction technique described previously (Gabbiani et al., 1996; Metzner et al., 1998). Although single spikes are sufficient to estimate low-frequency input, our *in vitro* results demonstrate that bursts are substantially better detectors of these signals. Although this corroborates previous *in vivo* results (Gabbiani et al., 1996; Metzner et al., 1998), we expand on these studies by showing that feature detection by bursts is low-frequency specific and, at least in some cells, dependent on the backpropagation-dependent burst mechanism.

Benefits of burst dynamics in neural coding

The analysis of feature extraction by the model neuron with intact burst dynamics revealed that bursts were superior feature detectors even when compared with similar high-frequency spike events produced by the model neuron with a passive dendrite. Moreover, the passive dendrite model required substantially

higher input current than the bursting model neuron to generate comparable numbers of high-frequency spike sequences, which significantly increased the mean firing rate. Thus, whereas coherence and stimulus estimation are enhanced by increases in firing rate (Borst and Haag, 2001), the reduced feature selectivity of the nonbursting neuron shows that these high-frequency events cannot adequately substitute for bursts when they are used as feature detectors.

Furthermore, it has been argued that neural codes should be sparse and implement low firing rates to be efficient (Rolls and Tovee, 1995; Vinje and Gallant, 2000). Burst dynamics may allow for such codes, because they permit low firing rates while having a significant proportion of distinct high-frequency burst events. We therefore propose that pyramidal cells may use high-frequency bursts to selectively detect and efficiently code for the low-frequency components of stimuli.

Decoding pyramidal cell output

For bursts to be an important component of a neural code, an appropriate postsynaptic decoding mechanism is required. For instance, a presynaptic burst ISI may match a resonance timescale of a postsynaptic neuron and gate information transfer (Izhikevich et al., 2003). Alternatively, bursts in conjunction with short-term synaptic plasticity could improve information transfer at unreliable synapses (Lisman, 1997). ELL pyramidal cell output is decoded in the torus semicircularis (TS) of the midbrain, and both synaptic depression and facilitation may contribute to the selective extraction of low-frequency features. Fortune and Rose (1997) demonstrated that different TS cells in a related electric fish are tuned for various stimulus–frequency ranges (low pass, broadband, and high pass). The output of the TS is conveyed to brain regions selectively responsible for electrolocation (low frequency, optic tectum) or electrocommunication (low to high frequencies, nucleus electrosensorius) (Bastian, 1982; Heiligenberg et al., 1991; Bell and Maler, 2004). A number of TS neurons temporally filters electrosensory stimuli resulting in low-pass (3–8 Hz), broadband (0–30 Hz), and high-pass cells (>10 Hz) (Fortune and Rose, 1997). In the case of low-pass cells, responses to stimulus frequencies >10 Hz are attenuated by synaptic depression. Interestingly, this depression can be overcome because these cells also show marked facilitation to high frequency (~10 msec ISI) presynaptic activity (Fortune and Rose, 2001). In this study, we have shown that ELL pyramidal cells convert low-frequency sensory input to high-frequency bursts. Thus, these bursts provide the high-frequency events that can be reliably detected by TS low-pass neurons. ELL pyramidal cells can therefore simultaneously code for both low-frequency stimulus components via bursts as well as high frequency components via isolated spikes. Different TS cell populations (low pass, broad-

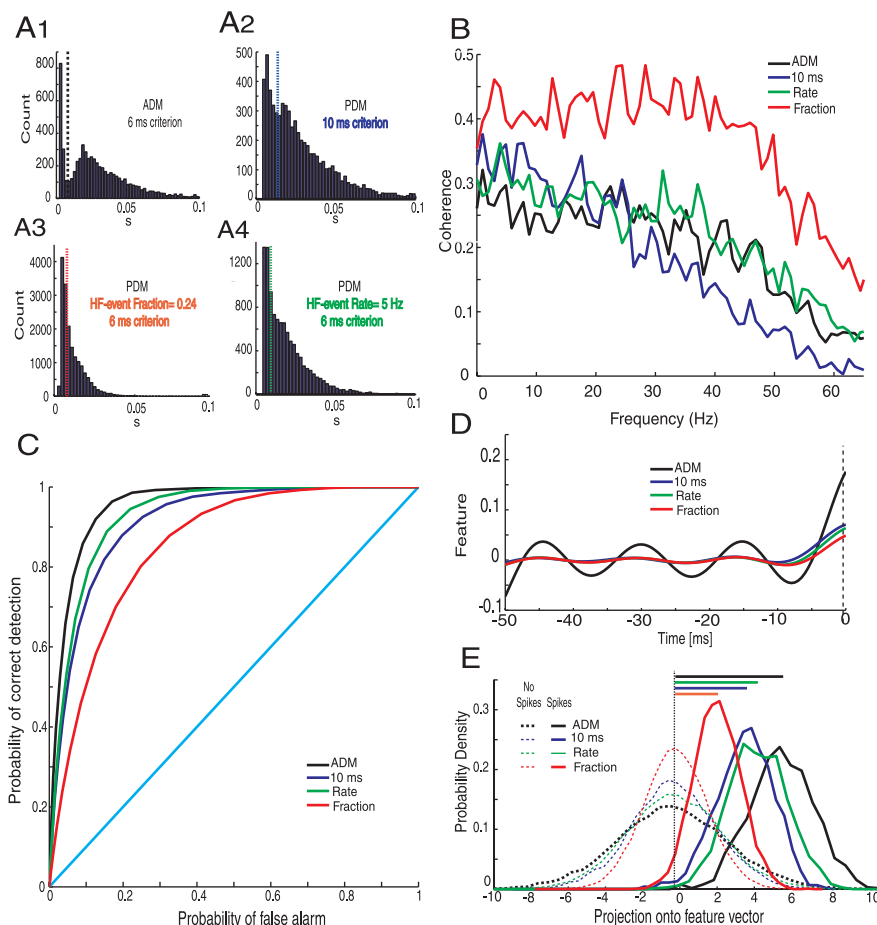


Figure 8. Burst dynamics are required for feature detection. *A*, ISI histograms in response to broadband stimulation of the model neuron with an ADM or a PDM. *A1*, Bimodal histogram of the ADM; black dashed line indicates the 6 msec ISI criterion for bursts (see Materials and Methods); firing rate, 33 Hz. *A2*, Unimodal histogram of the PDM; blue line indicates 10 msec ISI criterion that conserves the fraction (0.24) and rate (5 Hz) of HF events seen in the ADM; firing rate, 33 Hz. *A3*, PDM histogram in response to an increase in I_0 that increased the firing rate to 80 Hz but conserved the fraction of <6 msec HF events generated by the ADM (red line). *A4*, PDM histogram in response to an increase in I_0 that increased the firing rate to 50 Hz and conserved the rate of <6 msec HF events generated by the ADM (green line). *B*, Burst (ADM) or HF event (PDM) coherence with the broadband stimulus (colors as determined and indicated in *A*). *C*, ROC curves of bursts (ADM; black) or HF events (PDM; colors as in *A*). *D*, Optimal features detected by bursts or HF events in each scenario. *E*, Feature projections in each scenario onto the optimal feature (solid lines) as compared with the null feature (dashed lines).

band, or high pass) can then selectively decode information related to low-frequency signals emanating from prey versus high-frequency communication signals.

The ELL and the thalamus as sensory gateways

The parallels between the ELL-toral connections and thalamocortical pathways have been reviewed recently (Berman and Maler, 1999; Krahe and Gabbiani, 2004). Thalamic bursting has been both biophysically characterized and modeled (Lu et al., 1992; McCormick and Huguenard, 1992; Mukherjee and Kaplan, 1995; Smith et al., 2000). In lateral geniculate (LGN) relay cells, bursts have been shown to exhibit frequency tuning (Mukherjee and Kaplan, 1995; Smith et al., 2000; Fortune and Rose 2001) and enhance detectability at stimulus onset (Sherman, 2001; Martinez-Conde et al., 2002). Furthermore, it has been suggested that depending on the specific LGN-cortical layer projection, bursts and single spikes could present distinct streams of information transfer for the same stimulus features (Reinagel et al., 1999) and that feedback from the cortex could switch coding

modes between signal detection and stimulus estimation (Sherman, 2001).

Likewise, ELL pyramidal cell bursting is regulated through several dendritic and somatic ionic conductances (Noonan et al., 2003) and could conceivably be modulated by feedback input to the apical dendrites (Berman and Maler, 1999; Bastian and Nguyenkim, 2001). Recent studies have shown that ELL pyramidal cell responses to sensory input depend on both the frequency content and spatial extent of the stimulus (Bastian et al., 2002; Chacron et al., 2003; Doiron et al., 2003a) and that feedback is differentially activated by spatially localized versus spatially extended stimuli (Doiron et al., 2003a; Bastian et al., 2004). Thus, not only could feedback to the ELL switch coding strategies from feature detection by bursts to stimulus estimation by single spikes as proposed for the thalamus, but it could also lead to the differential coding of distinct behaviorally relevant low-frequency (prey like) versus high-frequency (communication) stimuli.

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