Burst and Tonic Response Modes in Thalamic Neurons During Sleep and Wakefulness

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Weyand, Theodore G., Michael Boudreaux, and William Guido. Burst and tonic response modes in thalamic neurons during sleep and wakefulness. J Neurophysiol 85: 1107–1118, 2001. Thalamic neurons can exhibit two distinct firing modes: tonic and burst. In the lateral geniculate nucleus (LGN), the tonic mode appears as a relatively faithful relay of visual information from retina to cortex. The function of the burst mode is less understood. Its prevalence during slow-wave sleep (SWS) and linkage to synchronous cortical electroencephalogram (EEG) suggest that it has an important role during this form of sleep. Although not nearly as common, bursting can also occur during wakefulness. The goal of this study was to identify conditions that affect burst probability, and to compare burst incidence during sleeping and waking. LGN neurons are extraordinarily heterogeneous in the degree to which they burst, during both sleeping and waking. Some LGN neurons never burst under any conditions during wakefulness, and several never burst during slow-wave sleep. During wakefulness, <1% of action potentials were associated with bursting, whereas during sleep this fraction jumps to 18%. Although bursting was most common during slow-wave sleep, more than 50% of the bursting originated from 14% of the LGN cells. Bursting during sleep was largely restricted to episodes lasting 1–5 s, with ~47% of these episodes being rhythmic and in the delta frequency range (0.5–4 Hz). In wakefulness, although visual stimulation accounted for the greatest number of bursts, it was still a small fraction of the total response (4%, 742 bursts/17,744 cycles in 93 cells). We identified two variables that appeared to influence burst probability: size of the visual stimuli used to elicit responses and behavioral state. Increased stimulus size increased burst probability. We attribute this to the increased influence large stimuli have on a cell’s inhibitory mechanisms. As with sleep, a large fraction of bursting originated from a small number of cells. During visual stimulation, 50% of bursting was generated by 9% of neurons. Increased vigilance was negatively correlated with burst probability. Visual stimuli presented during active fixation (i.e., when the animal must fixate on an overt fixation point) were less likely to produce bursting, than when the same visual stimuli were presented but no fixation point present ("passive" fixation). Such observations suggest that even brief departures from attentive states can hyperpolarize neurons sufficiently to de-activate the burst mechanism. Our results provide a new view of the temporal structure of bursting during slow-wave sleep; one that supports episodic rhythmic activity in the intact animal. In addition, because bursting could be tied to specific conditions within wakefulness, we suggest that bursting has a specific function within that state.

INTRODUCTION

Owing to similarities in receptive field structure, the lateral geniculate nucleus (LGN) appears to largely function as a relay of visual information between the retina and visual cortex. However, it is not a simple relay. Anatomical studies indicate that <20% of afferent synapses are of retinal origin (see Sherman and Guillery 1996, for recent review). These other inputs likely provide a gain mechanism for controlling the fidelity by which retinal inputs are transferred to cortex (Livingstone and Hubel 1981; Swadlow and Weyand 1985; see reviews by McCormick and Bal 1997; Sherman 1996; Singer 1977) or synchronize activity to boost the salience of spatially contiguous edges (Sillito et al. 1994). Bolstered by observations from intracellular records, several groups (e.g., McCormick and Feener 1990) have additionally promoted the idea that the LGN operates as a switch. In the tonic or "on" mode, retinal inputs are available for transfer to cortex, with the exact gain being a function of the current potency and sign of nonretinal afferents. In the "burst" or "off" mode, the membrane potential is predominantly hyperpolarized, but occasional (or even periodic) depolarization triggers a powerful low-threshold calcium conductance (I_t) that results in bursts of action potentials. This burst mode is correlated with slow-wave sleep (SWS). Whereas the function of the tonic mode is transparent, the function of the burst mode is obscure. Steriade et al. (1993) recently speculated that because the burst mode is associated with initiating and maintaining slow-wave sleep, its ability to drive the cortex into oscillations may function to correct some ionic imbalances caused by wakefulness. Although not nearly as frequent, bursting can also occur within wakefulness (Guido and Weyand 1995; McCarley et al. 1983; Ramcharan et al. 2000). Such observations are important, as they raise the possibility that bursting is used as a distinct signal in sensory processing (cf., Guido and Weyand 1995; Guido et al. 1995; Sherman 1996). The conditions under which bursting occurs during wakefulness are not well-delineated. Identifying these conditions was a major goal of the current study. A second goal was to better delineate the incidence of bursting during slow-wave sleep. Several recent and influential reviews have portrayed thalamic neurons as “disconnected” from their sensory inputs during slow-wave sleep, whereupon such neurons become oscillatory as a result of an interplay of their intrinsic conductances (McCormick and Bal 1997; Steriade et al. 1993). Such portrayals are a caricature of the one quantitative study of the temporal structure of LGN activity during sleep and waking (McCarley et al. 1983). A reassess-
ment of activity during slow-wave sleep might help to clarify this important issue.

**METHODS**

**Initial surgery**

All procedures were approved by the Institutional Care and Use Committees at Louisiana State University Medical Center, and the general methods have been described previously (Malpeli et al. 1992; Weyand and GaKa 1998). Briefly, cats underwent at least two sterile surgical procedures. In the first surgery, we cemented an aluminum crown to the skull to fix the head during subsequent behavioral and recording sessions and attached an insulated wire loop to the sclera of one eye (to allow us to determine gaze using the magnetic search coil technique) (Robinson 1962). The anesthetized cat was placed in a stereotaxic frame, and the fascia and muscle retracted to expose the skull. An aluminum crown was custom-fitted to the contours of the skull and affixed to the skull using stainless steel rod and bone cement. A Teflon-insulated coil was then attached to the sclera of one eye similar to the methods described by Judge et al. (1980). The leads from this coil were fed under the skin and crown, and terminated by soldering them to subminiature connectors. A fiberglass cover was attached to the crown to protect the connectors and microelectrode drive when the animal was not in the testing apparatus. The cat was removed from the stereotaxic frame and returned to its home cage.

Following training (described in **Training**), the cat was subjected to a second sterile surgery in which we implanted stimulating electrodes in visual cortex and a swiveling base for holding a microelectrode drive (Malpeli et al. 1992). For this surgery, the anesthetized cat was again placed in a stereotaxic frame, and holes were drilled through the cement and bone to expose visual cortex and the cortex overlying the LGN. Six to eight stimulating electrodes were placed individually into the lateral gyrus (1–3 mm apart and at depths of 3–5 mm) through an intact dura. The electrodes were cemented into place and the hole sealed with dental acrylic. A microelectrode base with a 9-mm-long cannula was inserted through the hole over the LGN. A protective stylus filled the cannula until we were ready to begin recording sessions. The hole over the LGN was then sealed with dental cement, and the cat removed from the stereotaxic frame and returned to its home cage. To obtain electroencephalographic (EEG) records, we either used a pair of unused stimulating electrodes, or, in one case, we inserted a staggered pair (2-mm vertical separation) of platinum-insulated wires into the contralateral cortex with the upper tip flush with the cortical surface.

**Training**

Following at least 1 wk to recover from initial surgery, food deprivation was begun and 24 h later behavioral testing initiated. The cat was placed in a loose-fitting canvas bag and the head fixed to a Plexiglas frame. The cat faced a dimly lit (0.5 cd/m²) screen on which we could project images such as bars, square-wave gratings (contrast 0.4) and a 0.2° spot produced by a low-power laser that was dimmed with a 3.0 N.D. filter. The cat was trained to look at this spot, and if it jumped to a new location, make a saccade to reacquire fixation on the spot. The bars and gratings were used to probe the excitability of the cell under study and were never behaviorally relevant. The laser spot was always behaviorally relevant.

**Testing/recording**

Following at least 1 wk to recover from surgery to implant stimulating electrodes, testing was initiated by placing the cat in the apparatus and replacing the protective stylus with a tungsten-in-glass microelectrode (~1.0 MΩ at 1 kHz). Signals from the microelectrode were amplified (×10,000), filtered (0.3–8 KHz, 24 dB/octave), and fed to an oscilloscope and audio monitor. EEG signals were also amplified (×10,000), filtered (1–60 Hz, 24 dB/octave), and passed to an oscilloscope and an A/D (A-D) converter that digitized the signals at 250 Hz. Horizontal and vertical eye position signals were amplified, filtered, and passed to an A-D converter that digitized these signals at 250 Hz. All data acquisition, control of behavioral testing, and stimulus display were under computer control. Single neurons were isolated on-line using a voltage discriminator, whose output (pulses) was fed to the computer. These pulses were then time-stamped (0.1-ms resolution) and put into a data file that also included a record of eye position, EEG, and stimulus status. For more than 30 recording sessions, the eye position, EEG signals, and unit activity were also passed to a VCR that digitized data at 22.5 KHz. These records were used for off-line analysis and for producing figures of analog traces.

For most sessions, we used electrical stimulation of visual cortex to help determine the position of the electrode in the brain. Electrical stimuli consisted of 0.1-ms monophasic pulses of varying intensity. Electrical activation was attempted for nearly all isolated neurons. Antidromic activation was inferred by invariant latency and verified by the test of impulse collision (Bishop et al. 1962). Some neurons (especially those dorsal to the LGN) were activated synaptically rather than antidromically.

**Definition of bursts**

Because we recorded extracellularly and filtered out activity <300 Hz, we could not directly observe changes in membrane potential associated with activation of the low-threshold calcium conductance (Iₜ). Lu et al. (1992) showed that when two or more action potentials separated by <4 ms (250 Hz) are preceded by at least 100 ms of quiescence, the probability of an underlying Iₜ is better than 0.99. These criteria correspond to what has been used in previous studies of bursting in anesthetized, paralyzed (Guido et al. 1995; Lu et al. 1993), and awake, behaving cat (Guido and Weyand 1995). Although we feel these criteria to be sufficient, our results from activity patterns during sleep indicate that they are overly conservative (cf., Lu et al. 1993; Ramacharan et al. 2000). Therefore in presenting our results, we have sometimes adopted more liberal criteria of two spikes in 6.66 ms (150 Hz) preceded by 50 ms of quiescence. Adapting one criterion or the other obviously alters the quantitative aspects of burst incidence, but does not alter the major conclusions of the study. In each figure and analysis, we indicate the criteria used: conservative (4 ms or less interspike interval preceded by 100 ms or more of quiescence) or liberal (6.66 ms or less interspike interval preceded by 50 ms or more of quiescence). Two-dimensional interspike interval plots (joint-interval histograms, JIHs) (McCarley et al. 1983) shown in Figs. 2 and 6 are particularly useful for viewing differences in burst incidence using liberal or conservative criteria. In Fig. 2, we have inscribed two “boxes” in the bottom right corner showing the differences between using conservative (smaller box) or liberal (larger box) criteria. For this representative example, changing criteria had essentially no effect in wakefulness, but increased the number of bursts during slow-wave sleep by ~25%.

**Definition of sleep**

Slow-wave sleep was defined by the presence of dominant cortical EEG spectra below 8 Hz. Although delta waves (0.5–4 Hz) were the most impressive EEG event during slow-wave sleep (e.g., Fig. 1), these waves were interspersed with episodes of faster activity that we also included as slow-wave sleep when accompanied by closed eyes and slow or absent eye movements. Rapid eye movement (REM) sleep was not observed, probably because we never deliberately sleep-deprived our animals, or perhaps because the testing situation was not particularly conducive to sleeping. In addition, sleep spindles (large-amplitude 10- to 14-Hz cortical oscillations) were not commonly observed probably because our EEG electrodes were above
RESULTS

We recorded from 148 LGN neurons in three cats. Among these, 109 were identified in layer A, 21 in A1, and 9 in C. In addition, we also encountered nine fibers above the LGN, four of which could be anterogradely activated from visual cortex. We assumed that the cell bodies of these fibers were in the LGN, although we cannot be sure of the layer. Because of eye movements in the awake animal, we made no attempt to classify neurons as X, Y, or W. Twenty-two neurons were identified using antidromic identification. Based on latency, we had a clear recording bias for Y cells (mean latency, 0.91 ms; $\sigma = 0.61$ ms; range, 0.4–3.0 ms; 15/22 latencies <1.2 ms) (So and Shapley 1979).

Bursts during sleep and waking

Figure 1 presents analog traces of an LGN neuron and concomitant cortical EEG during sleep (top) and wakefulness (bottom), illustrating the burst and tonic modes commonly associated with these two states. As the name implies, the burst mode is characterized by periods of quiescence interspersed with high-frequency “bursts” of action potentials. Other studies have demonstrated that these bursts are sodium spikes (action potentials) riding a slower depolarization attributed to the activation of the low-threshold calcium current $I_T$. The top records show expanded views of three bursts (conservative criteria). Bursting during sleep tends to be episodic or rhythmic, and we describe some quantitative aspects below. During “tonic” episodes as illustrated in the bottom traces, activity can be remarkably regular, often varying with ambient illumination level. For the traces shown, 95% of the activity was at 50 Hz or less. Figure 2 shows the temporal distribution of a different LGN neuron over a prolonged period (10 min) during which the cat drifts in and out of sleep. Figure 2A shows the distribution in raster format with the first spike in a burst indicated by a large dot, whereas Fig. 2B shows the spike distribution as JIHs (McCarley et al. 1983) during each epoch of sleep (left) and wakefulness (right). The JIHs sharply contrast the temporal structure of activity during sleep and wakefulness. During wakefulness, activity is remarkably regular (tonic) and a central distribution is obvious. In contrast, during sleep the regularity is disrupted (burst mode) and two tails form. The tail in the bottom right of each JIH is the cardinal spike of a burst, while the well-defined tail to bottom left indicates the shortest interspike interval (highest frequency) achieved within a burst. Figure 2C shows the sleep/wake cycles collapsed into two JIHs and includes insets to illustrate the differences in burst incidence using the two sets of criteria used in this study. Dots within the smaller inset represent the cardinal spike of bursts (and hence, the number of bursts) using the conservative criteria, whereas dots within the larger inset represent the cardinal spikes of bursts using the liberal criteria. Although these figures illustrate the general temporal structure of many LGN neurons during sleep and wakefulness, the data presented be-
low serve to emphasize that Figs. 1 and 2 are not necessarily representative. In fact, bursting during sleep is not ubiquitous, and tonic activity during wakefulness could be interrupted to include bursting.

Prevalence of bursting during SWS: how common is bursting during sleep?

Although bursting was common during sleep (0.51 bursts/s vs. 0.06 burst/s during waking, liberal criteria), the heterogeneity among LGN neurons in the degree to which they burst was striking. Figure 3A presents the incidence of bursting among 57 LGN neurons during SWS. For this figure, bursting during each 5-s epoch of SWS is coded as a shade of gray, with black being zero bursts observed and white being five bursts or more. Figure 3B presents the same data as in Fig. 3A, but now plotted as frequency of bursting during SWS. One-half of the cells failed to have a burst rate exceeding 0.5 bursts/s, and eight never burst. Another way of appreciating the heterogeneity in bursting among LGN neurons is that if we assume that these 57 neurons are representative, 50% of the bursting at any time is generated by 14% of the neurons.

As with burst frequency, the fraction of action potentials associated with bursting during sleep was appreciable, but variable. During some sleep epochs, >70% of spikes were associated with bursting, whereas in several cells, bursting never occurred. Overall, we estimate 18% of spikes during slow-wave sleep were burst related. However, because we could not directly observe the $I_T$ events, we are confident this is an underestimate (see DISCUSSION).

Rhythmic bursting during SWS: how rhythmic is bursting during sleep?

Rhythmic bursting is believed to play a prominent role in promoting the low-frequency EEG observed during SWS (e.g., McCormick and Bal 1997; Steriade et al. 1993). Although we showed that bursting is not ubiquitous among LGN neurons during SWS (Fig. 3), we did note that the presence of one burst increased the probability of observing another ($P < 0.005$; based on comparing the observed median burst interval with 999 simulations in each of 17 LGN neurons). Figure 4A shows the episodic nature of bursting (liberal criteria) in one LGN neuron over a 16-min period during which the cat was mostly in SWS. Figure 4B shows the distribution of interburst intervals for 17 LGN neurons for which we collected continuous records (10–50 min) for various intervals of SWS. It indicates that ~95% of the bursts occurred within 5 s of one another. To analyze potential underlying rhythmicity of such episodes, we identified “burst bouts” (see METHODS), i.e., periods during which bursting was common. If bursting was rhythmic, then the standard deviation of interburst intervals within a bout should approach zero. Figure 4C shows two examples of autocorrelograms of two burst bouts, one highly rhythmic (Fig. 4C, top) and one arrhythmic (Fig. 4C, bottom). Figure 4D shows the distribution of probabilities (determined using simulations; see METHODS) for 475 burst bouts collected from
A

B

8822 intervals
17 cells

C

10 bursts
p<0.001

11 bursts
p<0.153

D

E

475 bouts
mean = 2.49 Hz
s.d. = 0.88
delta range (1-4 Hz) = 93%
continuous records in the 17 LGN neurons. Using a criterion of $P < 0.05$, 47% of these bouts were rhythmic. Figure 4E shows the distribution of interburst intervals, showing that the vast majority of intervals were concentrated within the delta (i.e., 0.5–4 Hz) range. Although interburst intervals were within a frequency range appropriate for driving cortical delta rhythms, we failed to observe any consistent phase relationships between occipital EEG and concomitant thalamic bursting. This may, of course, be attributable to local variations in the occipital EEG (cf., McCarley et al. 1983).

**Bursting during visual stimulation: variability among neurons**

The majority of LGN neurons responded reliably and vigorously to visual stimuli of the appropriate polarity presented to the receptive field center. For most of these, bursting was not associated with the visually driven response. Figure 5A shows the visually driven response of an LGN neuron in tonic mode to a flashing stimulus placed over the receptive field, whereas Fig. 5B shows the visually driven response of a different LGN neuron whose initial response was often a high-frequency burst of action potentials (indicated by asterisks). Figure 5C shows the burst probabilities for 93 LGN neurons evaluated under different conditions of visual stimulation. This figure emphasizes the heterogeneity among LGN neurons. For 33/93 neurons (35%), we were unable to elicit any bursting, and as indicated, there was significant variability in bursting among the remaining 60 neurons. As with bursting during sleep, a minority of the neurons contributed most of the bursts. We found that 50% of the visually driven bursts originated from 9% (8/93) of neurons. Finally, although it would not seem unreasonable that neurons that burst prolifically during visual stimulation would also burst extensively during sleep, such a relationship was not observed.

**Bursting during visual stimulation: stimulus size**

The probability of observing a burst as part of the visual response could be increased by increasing stimulus size. Figure 6A shows the interspike interval histograms generated from single trials in which a 4° stimulus (top) or a 40° stimulus (bottom) are used to elicit the response. Clearly, bursting was evident with the large stimulus and not with the small stimulus. Figure 6B shows JIHs for the responses when we used the 4° stimulus (left) or the 40° stimulus (right). Probability of bursting for the small stimulus was 0.01 bursts/cycle and 0.37 bursts/cycle for the large stimulus. Figure 6C shows the probability of bursting we observed for each of four different stimulus sizes on this neuron. Table 1 shows that the results obtained for the single cell shown in Fig. 6 extend to a sample of 16 LGN neurons in which stimulus size was parametrically manipulated. For this larger sample, burst probability increased with stimulus size ($ \chi^2 = 229$, $P < 0.001$, 2 d.f.).

**Bursting related to state**

The probability of bursting during visual stimulation was greatest for the initial stimulus cycle. Figure 7A shows a single trial in which the initial, but not subsequent stimulus cycles elicited a burst. Figure 7B shows the ordinal distribution of 319 bursts from 67 cells across the first 5 cycles of visual stimulation in each trial. Overall, if a visual stimulus was going to elicit a burst, the burst would most likely occur to the first stimulus presented in a trial ($ \chi^2 = 17.01$, $P < 0.005$, 4 d.f.). One interpretation for this observation would be that the animal’s attention increased as the visual stimuli were presented. The task demanded considerable attention in the sense that the cat must inhibit the prepotent reflex of looking at a newly presented stimulus in the field. In some trials, it is conceivable that, although the cat was “on target,” the cat was less attentive and the cell relatively hyperpolarized.

Further evidence for a negative relationship between attention and bursting emerged when we analyzed bursting probability under “active” versus “passive” conditions. Figure 8 presents traces showing spike frequency during active (i.e., fixation spot present, Fig. 8A) and passive (i.e., no fixation spot present, Fig. 8B) conditions. As indicated by asterisks, bursts were present during six of the eight cycles shown under passive, but absent under active conditions. This cell was extensively analyzed under both active and passive conditions. During passive conditions, bursting occurred in 79/862 cycles (9.2%), whereas only one burst (1/118 cycles, 0.8%) was observed under active conditions. This observation in the single cell could be extended to our overall sample, bursting was more common under active than passive conditions (Table 2. $\chi^2 = 109$, $P < 0.001$, 1 d.f., 65 neurons passive only, 14 neurons active only, 15 neurons active and passive).

Finally, the least-likely period during which a neuron would exhibit bursting appeared to be active fixation, with or without an overt fixation point. We analyzed 28 LGN neurons during which the cat maintained fixation on the dimmed laser point placed at the center of an otherwise blank screen. In some trials, this fixation point was extinguished for 1 s and then returned. Compared to an epoch of “spontaneous” activity taken immediately prior to trial onset, burst probability during fixation was nominally decreased (0.046/s vs. 0.038 s, n.s., conservative criteria). However, for each cell, the probability

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**Fig. 4.** Bursting (liberal criteria) during sleep tended to be rhythmic. A: a 16-min continuous record of an LGN neuron indicating when bursting occurred (dots). For this record, the cat was mostly in slow-wave sleep, and bursting was only associated with sleep. Bursting tended to cluster in time. B: compiled interval histogram showing 8,822 interburst intervals observed in 17 neurons that were continuously monitored. It shows that most interburst intervals were 1 s or less. C: examples of autocorrelograms generated from a rhythmic burst-bout (top) and a poorly rhythmic burst-bout (bottom). The even spacing of the interburst intervals in the top autocorrelogram indicates rhythmicity. The results of the Monte Carlo simulations are plotted as probabilities for each burst-bout. For the top, the standard deviation of the observed interburst intervals was smaller than any of the simulations, and hence, $P < 0.001$. For the bottom, 152 of the 999 simulations had standard deviations less than the observed standard deviation of interburst intervals, and hence, $P < 0.153$. D: histogram of the probabilities obtained from analyzing 475 burst bouts. Forty-seven percent (223) of the burst bouts showed a probability of 0.05 or less, indicating that nearly ½ of all burst bouts during sleep are rhythmic, in a statistical sense. E: histogram plotting the mean frequency of each of the 475 burst bouts. The average frequency of these burst bouts was 2.49 Hz (i.e., average interburst interval was ~400 ms). As indicated, 93% of the interburst intervals were within the delta frequency range (0.5–4 Hz).
of observing a burst during the spontaneous epoch was greater than during fixation ($\chi^2 = 9.51, P < 0.01, 2 \text{ d.f.}$), reinforcing the view that membrane potential is depolarized during “attentive” states.

**Discussion**

This study makes several contributions to our understanding of the burst/tonic dichotomy of LGN activity during sleeping and waking. First, the heterogeneity among LGN neurons in the degree to which they burst during either sleep or wakefulness had not been fully appreciated. Second, we were able to identify specific conditions within wakefulness in which burst probability increased dramatically. These conditions include presenting stimuli more likely to influence inhibitory surround mechanisms, as well as manipulating behavioral contingencies. The former observations reinforce the potency of the inhibitory surround in shaping visual response. The latter indicates that “wakefulness” can include shifts in membrane potential sufficient to de-inactivate the $I_T$ burst mechanism. Finally, our method of analyzing potential rhythmic bursting indicates that most LGN neurons burst rhythmically during sleep, if only for a few seconds.

**FIG. 5.** Tonic and burst mode during visual stimulation. A: tonic response of an LGN neuron when a 7° square was flashed over the receptive field centered 11° to the left of gaze, and on the horizon (bars indicate light part of stimulus cycle). B, top: different LGN neuron that showed burst responses (indicated by asterisks), when a rectangle 14° wide by 12° high was flashed over the receptive field centered 15° left and 8° up from gaze. Bottom: expanded traces of the 3 bursts. C: distribution of burst probabilities for 93 LGN neurons during visual stimulation. For each neuron, we divided the number of bursts associated with presentation of a visual stimulus by the total number of cycles. This figure represents the burst probabilities calculated from 11,212 visual responses. LGN neurons varied widely in the degree to which bursting represented part of the visual response.

**What is the structure of the thalamocortical network during sleep?**

Several recent papers (e.g., see review by McCormick and Bal 1997) have promoted the idea that rhythmic bursting represents the dominant form of LGN activity during slow-wave sleep. Rhythmic bursting was the exception, rather than the rule, in the previous study of activity in the cat’s LGN (McCarley et al. 1983) (5 of 26 neurons). The recent study by Ramcharan et al. (2000) did not observe rhythmic bursting in the LGN of the sleeping monkey. We found that LGN bursting to be episodic, and rhythmic nearly half of the time. Although this seems to be a much different conclusion than the other studies, we strongly suspect that we were analyzing nearly identical spike trains. The difference is the method of analysis. McCarley et al. (1983) collapsed data across long epochs. Such treatment will necessarily dilute momentary rhythmicity that is interrupted by other arrhythmic epochs. The question McCarley et al. (1983) were addressing was whether there were overall trends in rhythmic behavior. The analysis Ramcharan et al. (2000) employed is straightforward: construct fast Fourier transforms (FFTs) from autocorrelograms and determine the statistical reliability of the FFT signal. Again, the problem is that they were analyzing long epochs that would dilute brief
epochs of rhythmic behavior. Using an analysis that tests for rhythmicity over brief epochs, we found most LGN cells burst rhythmically, if only for a few seconds. Thus our observations support the idea that slow-wave sleep is a period of episodic disconnections from sensory inputs, allowing intrinsic conductances to manifest brief episodes of rhythmic activity (cf., McCormick and Bal 1997; Steriade et al. 1993). Finally, we know our measurements are conservative. Because we depend on extracellular spikes to imply an underlying $I_T$ event, we have restricted our analysis to episodes of two spikes or more to identify bursting. In reality, there are $I_T$ events that generate one or no spikes. Thus underlying rhythmic activity is almost certainly even more common.

We observed significant heterogeneity among LGN neurons in the degree to which they burst, whether during sleeping or waking. Although one way of interpreting our results is that during any given second in slow-wave sleep the probability of observing a burst in an LGN neuron is just under 28%, such descriptions undermine the heterogeneity of the sample. Perhaps most telling is that 50% of the bursting appears to be produced by 14% of the neurons. McCarley et al. (1983) also noted heterogeneity in bursting among LGN neurons. They found a rather large fraction (37%) that did not burst during slow-wave sleep (vs. 17% for our large sample, 57 neurons).

The functional significance of this heterogeneity is unknown. However, large populations of synchronous rhythmic bursting would not be desirable as such activity could easily incite epileptiform activity (cf., Steriade et al. 1993).

Variability in bursting among LGN neurons during sleep and wakefulness indicates that either LGN neurons vary in the degree to which the $I_T$ channel is expressed, and/or, there are significant differences in intrinsic circuitry. Calcium imaging and whole cell in vitro studies suggest the $I_T$ channel and its kinetics are homogenous among relay cells (Coulter et al.
Qualitative and quantitative differences exist among relay cell afferents. Besides obvious differences in X, Y, or W retinal afferents, there is evidence suggesting that intrinsic, brain stem, and corticothalamic afferents vary in their density and distribution (Friedlander et al. 1981; Murphy and Sillito 1996; Weber et al. 1989; Wilson et al. 1984). These differences in circuits offer an obvious avenue for heterogeneity. For example, while global variables such as sleep and waking might shift membrane polarity across large populations of neurons, local variations in cortical afferent activity could alter the probability of the specific neuron recorded expressing a burst. Significant differences also exist between thalamic nuclei in the degree to which bursting is rhythmic during sleep. This point is explicit in the recent paper by Ramcharan et al. (2000). Despite an inability to observe rhythmic bursting in the monkey’s LGN, rhythmic bursting was striking in the nearby ventrobasal complex. The observations by Ramcharan et al. (2000) in the ventrobasal complex appear consistent with the analyses offered by Steriade and his colleagues (Domich et al. 1986; Glenn and Steriade 1982) of highly rhythmic bursting in the cat’s ventral lateral, ventral medial, and central lateral nuclei during slow-wave sleep.

**Sensory inputs, vigilance influence I_T mechanism during wakefulness**

Activation of the $I_T$ mechanism requires that the membrane potential be hyperpolarized for a period of 50–100 ms. This period allows the channel to “de-inactivate,” such that subsequent depolarization will alter channel configuration allowing calcium entry. In agreement with previous studies, bursting is much less common during wakefulness than slow-wave sleep (Hubel 1960; Livingstone and Hubel 1981; McCarley et al. 1983; Ramcharan et al. 2000). The observation that bursting during wakefulness occurs at all may be somewhat surprising since the membrane potential is largely depolarized during this period (promoting the tonic mode). However, variations in state is only one contributor to altering conductances to alter membrane potential. The receptive fields of LGN neurons have an antagonistic center-surround organization in which stimuli of one polarity excite one portion of the field and inhibit the other. In the awake animal, this inhibition is capable of de-inactivating the $I_T$ channel, despite the prominence of a tonic mode. Coenen and Vendrik (1972) show analog traces in which an obvious burst is evident following the removal of inhibition by extinguishing a light presented to the center of an off-center LGN neuron. Bursting could be promoted during wakefulness by providing large stimuli that activated both center and surround mechanisms, by manipulating level of alertness, or, as previously documented, encouraging eye movements (Guido and Weyand 1995; Lee and Malpeli 1998). We interpret our results to simply indicate that during wakefulness membrane potential is dynamic, and each of the conditions that promoted bursting are conditions that would predictably hyperpolarize the membrane potential. Whereas membrane potential is generally more depolarized during wakefulness than during sleep (Hirsch et al. 1983), we were able to relatively easily produce bursts in many LGN neurons, indicating that the membrane could be hyperpolarized for extended periods. Each of the conditions under which we observed bursting are conditions when the membrane potential could be expected to be hyperpolarized.

Increasing the size of the visual stimulus increased burst probability. Given the center-surround organization of LGN neurons, increasing stimulus size should have a greater influence on the surround than smaller stimuli. For some LGN neurons, whole field illumination sufficiently hyperpolarizes the LGN neuron to de-inactivate the $I_T$ channel and the occasional retinal input elicits a burst. In Fig. 6 we showed a more quantitative analysis of how stimulus size affects burst proba-

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<td>0.033</td>
<td>0.055</td>
<td>0.036</td>
<td>0.061</td>
<td>0.091</td>
<td>0.108</td>
<td>0.148</td>
</tr>
</tbody>
</table>
ability on a different LGN neuron. It is not obvious why large visual stimuli were more efficacious than small stimuli at eliciting bursts since our stimuli were sufficiently large in nearly all cases to include both the center and surround. One possibility is that the larger stimuli activated (inhibited) additional circuitry that lies beyond the classical receptive fields (e.g., McIlwain 1964), additionally hyperpolarizing the LGN neuron to deinactivate the $I_T$ channel. Whatever the mechanism, the potency of large stimuli (over smaller stimuli) in eliciting bursting has been observed previously in the LGN of the awake, paralyzed cat (Coenen and Vendrik 1972; their Fig. 2). Hyperpolarization associated with the large stimulus, but not the smaller stimulus, would be sufficient to de-inactivate the $I_T$ channel that elicits bursting on the next cycle. Had we better control of gaze, it would have been interesting to determine how effective stimuli restricted to the surround such as annuli would have been in promoting bursting. Finally, although we could promote bursting by using large stimuli, it is worth emphasizing that large stimuli provided no guarantee of observing burst responses. One-third of neurons never burst during visual stimulation, and 50% of visually driven bursting originated from 9% of cells analyzed. This again serves to underscore the variability in response characteristics among LGN neurons.

State affects burst incidence because state affects membrane potential. This has been demonstrated directly by Hirsch et al. (1983), and indirectly by a number of investigators. Again, the records of Coenen and Vendrik (1972) are instructive. Following extinguishing a spot to the center of an ON LGN neuron, bursts could be observed in the “drowsy” state, but not in the “awake” state (Coenen and Vendrik 1972; Fig. 5). Such an observation indicates that the membrane potential can become sufficiently hyperpolarized to de-inactivate the $I_T$ channel. Our own results, whether based on increased visually driven bursts associated with passive versus active gaze, or decreased burst probability associated with fixation, argues that state changes, as well as surround antagonism are capable of de-inactivating the $I_T$ channel. Several investigators have commented on the increased activity and/or transfer ratio associated with alertness or arousal (Coenen and Vendrik 1972; Livingstone and Hubel 1981; Sakakura 1968; Swadlow and Weyand 1985; also reviewed by Sherman and Guillery 1996; Singer 1977). What is the significance of bursting during wakefulness?

Having shown that bursting can occur during wakefulness and that such observations would be consistent with the known circuitry of the LGN, the question arises as to the functional significance of bursting in the awake animal. If bursting were restricted to slow-wave sleep, it and the supporting $I_T$ channel could be viewed as an adaptation for establishing slow-wave sleep in cortex. In wakefulness, bursting is such a departure from a linear response; it is not clear what it means as a visual signal. One attractive idea is that bursting has little to do with analyzing visual detail, but instead serves as a “wake-up call” to cortex, switching the LGN from burst to tonic mode (Guido and Weyand 1995; Guido et al. 1995; Sherman 1996). As the animal’s attention wanes, the neuron hyperpolarizes, deinactivating the $I_T$ channel. Subsequent visual stimuli would then activate the channel, sending a high-frequency volley of action potentials to visual cortex. This volley would be sufficiently powerful to drive corticothalamic neurons, which would then depolarize and switch LGN neurons to tonic mode via activation of metabotropic glutamate receptors (Godwin et al. 1996). In contrast to the burst mode, activity in the tonic mode is more linearly related to stimulus attributes (Guido et al. 1995; McCormick and Feeser 1990). Although attractive, our present results indicate that this idea is too limited. It does not explain, for example, the functional significance of bursting associated with saccades, nor with large visual stimuli. Further, as appealing and intuitive as the idea that bursting is more potent than single spike activity in activating cortical circuits (e.g., Lisman

### Table 2. Burst probability under passive and active gaze

<table>
<thead>
<tr>
<th></th>
<th>Total</th>
<th>Passive</th>
<th>Active</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bursts</td>
<td>742</td>
<td>423</td>
<td>319</td>
</tr>
<tr>
<td>Cycles</td>
<td>17,744</td>
<td>6,869</td>
<td>10,875</td>
</tr>
<tr>
<td>Probability, bursts/cycle</td>
<td>0.042</td>
<td>0.062</td>
<td>0.029</td>
</tr>
</tbody>
</table>

Number of neurons in Passive is 29 and in Active is 80.
1997; Reinarag et al. 1999), empirical evidence in a thalamocortical system is lacking.

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Murphy PC and Sillito AM. Functional morphology of the feedback pathway from area 17 of the cat visual cortex to the lateral geniculate nucleus. J Neurosci 16: 1180–1192, 1996.


