Discharge Profiles of Ventral Tegmental Area GABA Neurons during Movement, Anesthesia, and the Sleep–Wake Cycle

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Although mesolimbic dopamine (DA) transmission has been implicated in behavioral and cortical arousal, DA neurons in the ventral tegmental area (VTA) and substantia nigra pars compacta (SNc) are not significantly modulated by anesthetics or the sleep–wake cycle. However, VTA and SN non-DA neurons evince increased firing rates during active wakefulness (AW) and rapid eye movement (REM) sleep, relative to quiet wakefulness. Here we describe the effects of movement, select anesthetics, and the sleep–wake cycle on the activity of a homogeneous population of VTA GABA-containing neurons during normal sleep and after 24 hr sleep deprivation. In freely behaving rats, VTA GABA neurons were relatively fast firing (29 ± 6 Hz during AW), nonbursting neurons that exhibited markedly increased activity during the onset of discrete movements. Adequate anesthesia produced by administration of chloral hydrate, ketamine, or halothane significantly reduced VTA GABA neuron firing rate and converted their activity into phasic 0.5–2.0 sec ON/OFF periods. VTA GABA neuron firing rate decreased 53% during slow-wave sleep (SWS) and increased 79% during REM, relative to AW; however, the discharging was not synchronous with electrocortical α wave activity during AW, δ wave activity during SWS, or γ wave activity during REM. During deprived SWS, there was a direct correlation between increased VTA GABA neuron slowing and increased δ wave power. These findings indicate that the discharging of VTA GABA neurons correlates with psychomotor behavior and that these neurons may be an integral part of the extrathalamic cortical activating system.

Key words: ventral tegmental area; anesthesia; slow-wave sleep; rapid eye movement sleep; sleep deprivation; GABA; cortical activation

The ventral tegmental area (VTA) is the source of dopamine (DA)-containing neurons that project to structures in the ventral striatum, hypothalamus, and prefrontal association cortex, known collectively as the mesocorticolimbic DA system. This neural circuit has been implicated in mediating several motivated behaviors (for review, see Mogenson, 1987; Wise and Rompre, 1989). In this context, midbrain DA neurons in the VTA and substantia nigra pars compacta (SNc) respond to alerting, activating, and reward-related stimuli (Trulson and Preussler, 1984; Schultz, 1986; Freeman and Bunney, 1987; Schultz et al., 1993). Although mesolimbic DA transmission has been implicated in behavioral (for review, see Kalivas et al., 1993) and electrocortical (Radulovacki et al., 1979; Kropf et al., 1989; Kropf and Kuschinsky, 1991; Sebban et al., 1999a,b) activation, the firing rate of DA neurons in the VTA and SNc is not significantly modulated by the sleep–wake cycle or anesthetics (Miller et al., 1983; Steinfels et al., 1983). However, VTA and SNc non-DA neurons evince increased firing rates during active wakefulness (AW) and rapid eye movement (REM) sleep, relative to quiet wakefulness (OW) (Miller et al., 1983).

Although some progress has been made in elucidating the role of DA neurons in arousal and reinforcement, relatively less is known regarding the role of midbrain non-DA neurons in these behaviors. Midbrain neurons that are negative for tyrosine hydroxylase staining lie in close proximity to tyrosine hydroxylase-positive DA neurons. It has been suggested that these non-DA neurons are GABAergic neurons (Nagai et al., 1983; Otterson and Storm-Mathisen, 1984; Mugnaini and Oertel, 1985). GABA-mediated responses have been implicated in the modulation of the sleep–wake cycle (Nishikawa and Scatton, 1985). Increases in GABA release during slow-wave sleep (SWS) have been observed in the posterior hypothalamus (Nitz and Siegel, 1997), an area implicated in the regulation of behavioral arousal (Szymusiak and McGinty, 1986). Microinjection of GABA agonists into the posterior hypothalamus produces hypersomnia in the cat (Lin et al., 1989). Significantly, GABAergic neurons projecting to the posterior hypothalamus arise in the VTA and SNc (Ford et al., 1995). GABAergic neurons likely play a critical role in the modulation of DA mesocorticolimbic neurotransmission, which has recently been implicated in the control of REM sleep in the canine model of narcolepsy–cataplexy (Nishino and Mignot, 1997).

We have recently characterized, in anesthetized rats, a homogeneous population of VTA non-DA neurons that contain GABA, connect to DA neurons, and project to corticolimbic structures (Steffensen et al., 1997, 1998). They were distinguished electrophysiologically from DA neurons by their rapid-firing, nonbursting activity, short-duration action potentials, EPSP-dependent spontaneous spikes, and lack of spike accommodation to depolarizing current pulses. To evaluate the potential role of VTA GABA neurons in cortical arousal and psychomotor behavior, we studied the discharge profiles of these neurons during the induction and maintenance of adequate anesthesia, during elec-
trocortical rhythmic activity, and during the sleep–wake cycle in normal and sleep-deprived unrestrained rats.

Preliminary results have been published previously (Steffensen et al., 1996; Lee et al., 1997).

MATERIALS AND METHODS

Animal care. Nineteen male Sprague Dawley rats (Charles River Laboratory, Hollister, CA) weighing 300–500 gm were housed individually with ad libitum access to food and water and were maintained on a reverse 12 hr light/dark cycle (off at 10:00 A.M., on at 10 P.M.). Animal care, maintenance, and experimental procedures were in accordance with the Scripps Research Institute Animal Research Committee (IACUC approved; Animal Welfare Assurance no. A3194-01).

Microwire electrode implantation surgery and single-unit recording. Rats were anesthetized with sodium pentobarbital (50 mg/kg, i.p.) for microwire implantation surgery. Eight stainless steel Teflon-insulated micro-wires (50–62 µm) were assembled in a single bundle (diameter of spayed microwires tip is 0.75 mm; NB Labs, Denison, TX) were connected to a pin on one or two strip connectors. Microwire bundles were lowered into the VTA [−5.6 to −6.2 mm anteroposterior, 0.7–1.0 mm mediolateral, and 7.8 mm from the cortical surface (Paxinos and Watson, 1986)] (Nagai et al., 1983; Otterson and Storm-Mathisen, 1984; Mugnaini and Oertel, 1985). EEG leads (120 µm) were connected to screws implanted in the cranium over the head of the caudate nucleus and left parietal and frontal cortices; however, only retrosplinal to contralateral frontal recordings from electrodes located over the retrosplinal, parietal, and frontal cortices were monitored but no SWS was allowed. After a minimum of 24 hr of sleep deprivation, each rat was again connected to the recording apparatus, and EEG and single-unit activity were recorded simultaneously in 6 of the 19 rats during a sleep–wake cycle before and after 24 hr of sleep deprivation. The last sleep–wake cycle before sleep deprivation was recorded between 8:00 A.M. and 12 P.M. during the reverse 12 hr light/dark cycle (off at 10:00 A.M., on at 10:00 P.M.). After an episode of normal REM sleep, each rat was awakened and housed together with the other rats in a 3 × 3 × 3 foot, open-field box. Their activity was monitored continuously. They were constantly handled and exposed to novel objects and alerting stimuli during the 24 hr period of sleep deprivation. Every 2–3 hr, each rat was connected to the recording apparatus, and EEG and single-unit activity were monitored but no SWS was allowed. After a minimum of 24 hr of sleep deprivation, each rat was again connected to the recording apparatus and allowed to sleep. Care was taken to record the deprivation sleep between 8:00 A.M. and 12 P.M.

Histology. At the termination of the chronic recordings, electrolytic lesions (≤ 3 mA; 10–15 sec; Stimulator S88 and Isolator Unit PSIU 6, Grass Instrument, Quincy, MA) were passed through the recording electrode during deep anesthesia to verify its location in the VTA region. The animals were subsequently administered a lethal dose of halothane anesthesia or pentobarbital, and the brains were removed and preserved in 10% formalin. The brains were frozen and sectioned in a cryostat into 50 µm slices for inspection of the lesion site.

RESULTS

Extracellular electrophysiological characterization of VTA GABA neurons

We have previously described the electrophysiological, neurochemical, and ultrastructural characteristics of VTA GABA neurons in anesthetized (Steffensen et al., 1998) and freely behaving rats (Gallegos et al., 1999). In brief, VTA GABA neurons recorded in halothane-anesthetized rats represent a homogeneous population of phasic (only when anesthetized; see below), rapid-firing, nonbursting, short duration (<500 µsec) action potential neurons that connect to VTA DA neurons and receive excitatory input from the cortex and hippocampus. The most distinguishing feature of VTA GABA neurons recorded in halothane-anesthetized rats was their uninterrupted phasic activity characterized by alternating 0.5–2.0 sec ON/OFF periods (Fig. 1B) (Steffensen et al., 1998). In freely behaving rats, VTA GABA neurons do not exhibit phasic activity (Fig. 1C) (Gallegos et al., 1999). They can be classified as VTA GABA neurons based on their spiking characteristics and by response to afferent input. As in anesthetized rats, VTA GABA neurons are relatively rapid-firing neurons. The range of firing rates of all VTA GABA neurons recorded in this study during AW ranged from 4 to 65
Hz, with a mean of 28.7 ± 5.6 Hz (n = 25). This was significantly higher (p < 0.05) than the mean firing rate of 19 Hz reported previously for VTA GABA neurons recorded in halothane-anesthetized rats (Steffensen et al., 1998). Similar to VTA GABA neurons recorded in anesthetized rats, VTA GABA neurons were characterized by nonbursting, short-duration (<500 μsec) spikes (Fig. 1A). Spike characteristics, as well as anatomical localization to the VTA, were the primary criteria used to classify the neurons as VTA GABA neurons. In addition to the primary spiking criteria (i.e., initial negative-going spike waveforms, <500 μsec spike duration, nonbursting, relatively fast firing), we also established secondary criteria based on their response to afferent input. VTA GABA neurons were identified as such by at least one of the following stimulation criteria: multiple spiking after high-frequency stimulation of the internal capsule (IC); dual-latency spiking after single stimulation of the fimbria/fornix (f/f); or inhibition of spontaneous activity by single stimulation of the nucleus accumbens (NAcc). VTA GABA neurons were consistently driven orthodromically or antidromically, or both, by single stimulation of the IC. Short trains of high-frequency IC stimulation (10 pulses at 200 Hz) elicited multiple spike discharges that occurred with latencies nearly an order of magnitude greater than their single-spike antidromic or orthodromic latency of 2–3 msec (Fig. 1D). We have previously demonstrated that IC-stimulated multiple spiking is blocked by systemic MK-801 or in situ micro-electrophoretic application of APV, indicating that the IC-stimulated input is mediated by NMDA receptors (Steffensen et al., 1998). VTA GABA neuron spikes are also elicited orthodromically by fimbria/fornix stimulation at dual latencies (mean latency = 6.2 ± 1.1 msec and 22 ± 2.3 msec; n = 7) (Fig. 1E). Finally, these neurons could also be identified in the freely behaving rat by the inhibition of their spontaneous activity after stimulation of the NAcc (Fig. 1F) (mean duration of inhibition = 82 ± 7 msec; n = 6). All neurons classified as VTA GABA neurons met the criteria for spike characteristics and either were driven by IC or f/f stimulation or inhibited by NAcc stimulation.

**Effects of movement and anesthetics on VTA GABA neuron spontaneous activity**

We observed that the firing rate of VTA GABA neurons was phasically modulated by diverse forms of motor activity. The type of movement was not quantitatively examined; however, marked accelerations in firing rate were associated with the onset of certain movements such as head orienting or forelimb movement or transitions from SWS to AW. On the other hand, little variation in firing rate was observed with transitions to or during sustained locomotor activity. During phasic motor activity the firing rate of each neuron increased dramatically. The rate meter in Figure 2A depicts the firing rate of three VTA GABA neurons recorded...
simultaneously in the same rat during AW/QW and during the induction of anesthesia by chloral hydrate. During movement the firing rate of each neuron increased dramatically. The mean increase was 85 ± 6% (n = 14). Spontaneous firing rates often eclipsed 100 Hz for 10–20 sec. Systemic administration of 200 mg/kg chloral hydrate markedly decreased the firing rates of the three VTA GABA neurons recorded simultaneously from microwire electrodes chronically implanted in a freely behaving rat during movement and the induction of anesthesia produced by chloral hydrate. Although the firing rates of the three VTA GABA neurons differed, all were characterized by marked increases in firing during movement (horizontal bars). Chloral hydrate (200 mg/kg) suppressed the firing of two VTA GABA neurons and modestly decreased the firing of the remaining neuron in this rat. Adequate anesthesia was determined by the lack of reflex response to tail pinch. B. Compared with saline, adequate anesthesia produced by intraperitoneal chloral hydrate or ketamine (150 mg/kg) and exposure to halothane vapor (1% in 4 × 4 × 10 inch Plexiglas box) significantly decreased VTA GABA neuron firing rate by 86, 62, and 45%, respectively. Asterisks indicate significance level p < 0.05. C, D, These rate meter records show 5.0 sec epochs of the instantaneous firing rates of the same three simultaneously recorded VTA GABA neurons before (C) and after (D) the induction of chloral hydrate anesthesia. Note that before anesthesia the firing rate of each spike is relatively regular even when sampled at 100 msec time bins. After adequate anesthesia, the regular firing rate is transformed into pronounced phasic (ON/OFF) activity. Indeed, in one cell there are paroxysms of increased firing during the ON periods of phasic activity; however, when averaged over 2 min there is a definite slowing of firing rate, likely caused by the suppression of firing during the OFF period of phasic activity. Often, the ON period of phasic firing during anesthesia is characterized by an initial acceleration of firing that likely gives rise to the transients, followed by an adaptation and then an abrupt OFF period.

VTA GABA neuronal activity during the sleep–wake cycle

Active wakefulness was recognized by low-voltage, desynchronized EEG activity, increased EMG activity, locomotor activity, upright posture, open eyes, and responsiveness to sound or touch (Fig. 3). SWS was characterized by the presence of high-voltage, synchronized EEG activity, recumbent posture, closed eyes, and diminished EMG activity. Rapid eye movement sleep was characterized by low-voltage desynchronized EEG, continued behavioral signs of sleep, and a decrease in EMG activity to the level of background noise. The discriminated unit activity of a relatively slow VTA GABA neuron recorded simultaneously with the EEG and EMG activity is also shown. The discharging of this VTA GABA neuron is modulated by the stage of sleep. Figure 4 shows the firing rate of a more typical, rapidly firing VTA GABA neuron where background noise. The discriminated unit activity of a relatively slow VTA GABA neuron recorded simultaneously with the EEG and EMG activity is also shown. The discharging of this VTA GABA neuron is modulated by the stage of sleep. Figure 4 shows the firing rate of a more typical, rapidly firing VTA GABA neuron during multiple sleep–wake cycles over >3 hr. The firing rate was modulated by movement during AW, was regular during SWS, and was consistently elevated during REM episodes. Figure 5 shows the firing rates of all 25 VTA GABA neurons studied during the sleep–wake cycle. To summarize, VTA GABA neuron firing rate decreased significantly (p = 0.0012; F = 13.475) during SWS and increased significantly (p = 0.042; F = 4.602) during REM sleep, relative to AW (Fig. 5B).
Correlations between VTA GABA neuronal activity and electrocortical activity

The decrease in VTA GABA neuron firing during SWS and the increase during REM relative to AW was not accompanied by synchronized rhythmic activity. In other words, they did not exhibit instantaneous or rhythmic firing (bimodal distribution of interspike intervals) at the same frequency as retrosplenic electrocortical δ (1–4 Hz) activity during SWS (mean SWS firing rate = 12.9 ± 2.6 Hz; n = 25) or α (8–18 Hz) activity during AW. However, their instantaneous and average firing rate were within the broad frequency range of γ activity (30–58 Hz) during REM (mean REM firing rate = 37.9 ± 5.6 Hz; n = 25). Despite the marked slowing of VTA GABA neurons during SWS, the instantaneous firing rate of VTA GABA neurons was rarely correlated with δ activity, the predominant EEG frequency of the retrosplenic cortex. With the possible exception of the correlation between instantaneous firing rate and γ activity during REM, VTA GABA neuron unit activity, as determined by inspection of the first-order interval spike histograms or autocorrelograms, showed no rhythmic activity in association with α activity during AW or δ activity during SWS. Nonetheless, to more closely examine the possibility that the unit discharge activity might be correlated with electrocortical activity, we performed STA of VTA GABA neuron discharges during AW, SWS, and REM sleep. As shown in Figure 6, there appeared to be little correlation between unit firing and retrosplenic EEG activity.

Correlations between VTA GABA neuronal activity and electrocortical activity after sleep deprivation

Sleep deprivation produces an increase in δ wave power during deprived sleep relative to normal sleep (Rosenberg et al., 1976; Borbely et al., 1981; Lancel et al., 1991). Because VTA GABA neuron firing rate decreased during SWS and increased during REM relative to AW, we sought to determine whether VTA GABA neuron activity correlated with the changes in EEG power produced by sleep deprivation. Figure 7 shows the simultaneous δ and γ band activity associated with the firing rate of two VTA GABA neurons recorded from two separate sleep-deprived rats. There is a marked increase in δ activity during SWS and a mild increase in γ activity during REM, but not AW. As during normal sleep, deprived sleep VTA GABA neuron firing appears to be activity dependent during AW, low during SWS, and enhanced during REM sleep. Figure 8 summarizes the effects of deprived sleep on EEG band power and VTA GABA neuron firing rate, as well as the correlation between the changes that occurred in EEG band power versus the changes that occurred in VTA GABA neuron firing rate during AW, SWS, and REM. Compared with the last episode of sleep before deprivation (Fig. 8A), deprived-sleep α activity during AW increased significantly (p = 0.013; F_{(2,11)} = 14.183; mean normal sleep 8–18 Hz power = 0.22 ± 0.02 mV²/Hz), δ activity during SWS increased significantly (p = 0.004; F_{(2,11)} = 35.807; mean normal sleep 1–4 Hz power = 3.9 ± 0.33 mV²/Hz), and γ activity during REM was not significantly affected (p = 0.253; F_{(2,11)} = 1.664; mean REM sleep 30–58 Hz power = 0.07 ± 0.009 mV²/Hz). The firing rate of VTA GABA neurons was averaged during the same epochs corresponding to the EEG analysis above. Compared with the last episode of sleep before deprivation, deprived-sleep VTA GABA neuron firing rate (Fig. 8B) was not significantly (p = 0.956; F_{(2,33)} = 0.003) affected during AW (mean normal sleep AW firing rate = 34.2 ± 7.8 Hz), decreased significantly (p = 0.011; F_{(2,33)} = 8.277) during SWS (mean normal sleep SWS firing rate = 16.1 ± 3.4 Hz), but was not significantly (p = 0.102; F_{(2,33)} = 3.009) affected during REM (mean normal sleep REM firing rate = 43.4 ± 7.3 Hz). Figure 8C summarizes the relationship between the change in VTA GABA neuron firing rate and the change in α, δ, and γ power corresponding to deprived versus normal AW α activity, SWS δ activity, and REM γ activity, respectively. Each point represents the average change in firing.
rate of all VTA GABA neurons recorded in a particular rat (one point per rat) and the average change of power for each of the bands. There was a mild correlation between the increase in δ wave power and the decrease in VTA GABA neuron firing rate during SWS ($r = 0.658; p < 0.05$).

VTA GABA neuron firing rate was also correlated with changes in total EEG rms voltage. Compared with the last episode of sleep before deprivation, deprived-sleep EEG rms during AW increased significantly ($p = 0.0025; F_{(2,11)} = 31.23$; mean normal AW rms $V = 0.34 \pm 0.02$ mV), during SWS increased significantly ($p = 0.00001; F_{(2,11)} = 325.456$; mean normal SWS rms $V = 0.53 \pm 0.02$ mV), and during REM increased significantly ($p = 0.004; F_{(2,11)} = 24.270$; mean normal REM rms $V = 0.4 \pm 0.03$ mV). Similar to δ wave activity and VTA GABA neuron slowing, there was a mild correlation between the increase in rms voltage and the decrease in VTA GABA neuron firing rate during SWS ($r = 0.696; p < 0.05$).

**DISCUSSION**

The most distinguishing feature of VTA GABA neuron spontaneous activity recorded in halothane-anesthetized rats was their uninterrupted phasic activity characterized by alternating 0.5–2.0 sec ON/OFF periods (Steffensen et al., 1998). In freely behaving rats, phasic activity was not observed, and the firing rate, on average, was greater than in halothane-anesthetized rats (i.e., $33 \pm 5$ Hz vs $19 \pm 2$ Hz). Although not quantified in this study, the firing rate of these neurons was modulated during movement, often associated with the initiation of certain head or forelimb movements or onset of waking, but not during sustained locomotor activity, because VTA GABA neuron firing rate is not modulated during traverse of a 5 foot runway for reward (R. A. Gallegos, S. C. Steffensen, J. R. Criado, R.-S. Lee, and S. J. Henriksen, unpublished observation). We have observed VTA GABA neuron spontaneous firing rates exceeding 100 Hz during specific motor behaviors or during REM sleep, a rate that is consistent with their short refractory period and lack of spike accommodation (Steffensen et al., 1998).

Although general anesthetics do not significantly affect the spontaneous firing rate of midbrain DA neurons, they reduce their characteristic bursting activity and alter their sensitivity to DA receptor agonists and drugs of abuse (Bunney et al., 1973a,b; Mereu et al., 1984; Kelland et al., 1990). In contrast, the firing rate of VTA GABA neurons was reduced significantly by the three anesthetics quantified in this study and abolished by others not quantified, including the fast-acting and slow-acting barbiturates (S. C. Steffensen, R.-S. Lee, and S. J. Henriksen, unpublished observation). VTA GABA neuron firing rate was depressed most by chloral hydrate, then ketamine, and then halothane. All of these anesthetics produced adequate anesthesia, as determined by
the lack of reflex response to tail pinch. Adequate anesthesia not only depressed VTA GABA neuron firing rate but induced pronounced phasic ON/OFF activity similar to that reported previously in rats maintained on halothane (Steffensen et al., 1998). These results demonstrate that VTA GABA neurons are especially sensitive to anesthetics and that anesthetics induce a pattern of discharge activity that differs significantly from that during SWS, wherein the discharge activity of VTA GABA neurons was also slow, but regular, and nonphasic.

The activity of VTA GABA neurons was studied during the normal sleep–wake cycle to evaluate their relationship to cortical arousal. Relative to AW, VTA GABA neuron firing rate decreased 53% during SWS. Although VTA GABA neuron unit discharge slowed during both SWS and anesthesia, we could not distinguish whether the decreased rate resulted from reduced afferent input or from intrinsic decreases in the excitability of VTA GABA neurons. However, because we have demonstrated previously that the firing rate of VTA GABA neurons is highly dependent on excitatory synaptic input from NMDA receptor-mediated excitatory afferents (Steffensen et al., 1998), it is likely that the slowing results, at least in part, from diminished glutamatergic input. VTA GABA neuron unit discharge increased 79% during REM sleep, a state characterized by an inhibition of EMG activity and decreased responsiveness to external stimuli (Wu et al., 1989). This observation indicates that it is possible for the discharge of VTA GABA neurons to increase independently of motor activity or sensory input. Furthermore, changes in gross locomotor activity exhibited little correlation with VTA GABA neuron firing rate, providing further evidence that their activity does not merely reflect changes in motor output.

Although the rate or pattern of firing of midbrain DA neurons appears to be unaltered during the sleep–wake cycle (Miller et al., 1983; Steinfels et al., 1983), it has been demonstrated that non-DA neurons in the substantia nigra reticulata (SNr) and VTA evince increased firing rates during REM compared with SWS and in AW compared with QW (Miller et al., 1983). However, there was no significant difference in the firing rate of VTA or SNr non-DA neurons during REM sleep stage compared with AW (Miller et al., 1983). In contrast, here we report a significant increase in the firing rate of VTA GABA neurons during REM sleep compared with AW. The activity of this homogeneous population of VTA GABA neurons is modulated differentially during the sleep–wake cycle and preferentially during REM

Figure 5. Summary of VTA GABA neuron spontaneous firing rate during slow-wave sleep and REM sleep compared with active wakefulness. A, Firing rates for VTA GABA neurons were sampled in 10 sec epochs randomly during all states, and a minimum of 12 epochs was taken for each state for determinations of firing rate. All of the 25 VTA GABA neurons that were studied demonstrated decreased firing rates during SWS relative to AW. Most of the 25 VTA GABA neurons that were studied showed increased firing rates during REM relative to AW. B, This graph summarizes the firing rates of VTA GABA neurons during SWS and REM compared with AW. VTA GABA neuron mean firing rate was significantly decreased during SWS and significantly increased during REM sleep, relative to AW. Asterisks indicate significance level $p < 0.05$.

Figure 6. Lack of correlation between VTA GABA neuron spike discharge and EEG activity. The three traces show spike-triggered averaging of unit-EEG cross-correlation for a single neuronal potential recorded in a representation rat across sleep–wake states. VTA unit spikes (100 individual events) were used to average the pre-unit and post-unit discharge EEG activity during each of the three states, AW, SWS, and REM. Regardless of state, there was no correlation between unit activity and EEG activity. The dashed zero-line is the time of occurrence of VTA GABA neuron spike.
sleep, when motor responses are “paralyzed,” suggesting that they do not subserve motor behaviors per se but are involved in psychomotor-related events underlying cortical arousal.

We explored a possible causal relationship between VTA GABA neuron firing and cortical activation by correlating unit activity with EEG spectral band activity during AW, SWS, and REM sleep. VTA GABA neuron spiking was not rhythmically synchronized with δ, α, or γ activity during AW, SWS, or REM, respectively. However, the instantaneous and average firing rates of VTA GABA neurons were correlated temporally with γ activity during REM, indicating a link between unit activity and retrosplenial EEG activity. Whether VTA GABA neuron activity contributed to or just reflected the cortical rhythm was beyond the scope of this study. Such determinations likely require in situ pharmacological or experimental manipulations of neuronal activity.

δ (1–4 Hz) wave EEG activity is a function of previous waking. During SWS, δ activity is maximal at the beginning of the sleep period and declines progressively during the sleep period (Rosenberg et al., 1976; Borbely et al., 1981; Lancel et al., 1991). After sleep deprivation, δ activity is enhanced, especially in the first part of deprived sleep (Rosenberg et al., 1976; Borbely et al., 1981; Tobler and Borbely, 1986; Lancel et al., 1991). Indeed, in humans and rats, the rate of rise and peak response of δ activity during SWS increases after sleep deprivation (Trachsel et al., 1989; Dijk et al., 1990). We found that δ activity during SWS increased nearly threefold in deprived sleep versus normal sleep. Concomitant with the increase in δ activity was a decrease in VTA GABA neuron firing rate. In fact, there was a mild correlation between the degree of increase in δ activity and the degree of slowing of VTA GABA neuron activity during the progressive increase in δ wave activity (seen best during the first episode of SWS in B). Temporal correlations can be drawn between EEG spectral band activity, designation of sleep state, and VTA GABA neuron activity. EEG spectral band power was determined at 4.0 sec epochs.

Figure 7. Cortical electroencephalographic spectral band activity and VTA GABA neuron firing rate. A, B, EEG spectral band activity and VTA GABA neuron firing rate are shown for two sleep-deprived rats. In both rats, retrosplenial δ wave activity (1–4 Hz) was greatest during SWS relative to AW and REM and increased progressively during each SWS episode. Notwithstanding the relative amplitude and signal-to-noise ratio, retrosplenial γ wave activity (30–58 Hz) was of greatest amplitude during REM sleep, relative to AW and SWS. VTA GABA neuron firing rate was directly correlated with the EEG spectral band activity, being low when δ wave activity was high and high when γ activity was high. Note that there is a small yet progressive slowing of VTA GABA neuron activity during the progressive increase in δ wave activity (seen best during the first episode of SWS in B). Temporal correlations can be drawn between EEG spectral band activity, designation of sleep state, and VTA GABA neuron activity. EEG spectral band power was determined at 4.0 sec epochs.

Concomitant with the increase in δ activity was a decrease in VTA GABA neuron firing rate. In fact, there was a mild correlation between the degree of increase in δ activity and the degree of slowing of VTA GABA neuron activity, suggesting a link between VTA GABA neuron activity and cortical arousal (Fig. 8). Enhanced θ wave power has also been observed during REM sleep in deprived rats (Borbely et al., 1984; Tobler and Borbely, 1986). It was hypothesized that, similar to the regulation of SWS, REM recovery results from an increase in both duration and intensity of θ; however, more recent studies have failed to find consistent elevations in θ activity during REM recovery (Lancel et al., 1992). γ wave (30–58 Hz) activity and θ activity covary across the sleep–wake cycle, being high during AW and REM and low during SWS or QW (Maloney et al., 1997). It also reflects cortical arousal, independent of motor activity, attaining maximal levels during REM, when EMG activity is minimal. It was proposed that the covariation of γ and θ activity across states and behaviors suggests that a common system may modulate these fast and slow EEG rhythms and that such modulation, potentially emanating from the basal forebrain (Maloney et al., 1997), could predominate during certain states or behaviors, such as during REM sleep. We did not observe a significant correlation between the degree of increase of γ activity and the degree of increase of VTA GABA neuron firing rate after sleep deprivation, likely because of the lack of significant change in γ activity during deprived REM.

Early stimulation (Moruzzi and Magoun, 1949) and lesion
This graph summarizes the effects of 24 hr sleep deprivation on EEG spectral power during the sleep–wake cycle. Although α wave activity (α: 8–18 Hz) during AW and δ (δ: 1–4 Hz) wave activity during SWS were significantly increased during deprived sleep relative to normal sleep, γ (γ: 30–58 Hz) wave activity during REM sleep was not significantly altered (expressed as percentage deprived vs normal sleep). Asterisks equal significance level p < 0.05. B, This graph summarizes the effects of 24 hr sleep deprivation on VTA GABA neuron activity during the sleep–wake cycle. Although there was no significant difference in VTA GABA neuron firing rate during deprived AW α wave activity in deprived rats, there was a significant slowing during deprived SWS δ wave activity relative to normal sleep. Asterisk equals significance level p < 0.05. C, This graph plots the change in VTA GABA neuron firing rate on the abscissa versus the change in EEG spectral band power for each state of the sleep–wake cycle in deprived sleep versus normal sleep. Note that VTA GABA neuron slowing increases as a function of increased δ wave activity during SWS. There is no clear relationship between the change in VTA GABA neuron firing and the change in α or γ activity during either AW or REM sleep, respectively. Each point within each category represents a different rat.

(Bach-Y-Rita et al., 1966) studies have implicated the midbrain reticular formation, including the VTA, in the electrocortical and behavioral activation that characterize wakefulness. However, studies involving more selective lesions of the reticular formation have revealed a dissociation between behavioral and electrocortical activation (Feldman and Waller, 1962; Jones et al., 1973), indicating that distinct subareas of the rostral brainstem core underlie their respective mechanisms. Cholinergic neurons in the basal forebrain serve as the extrathalamic relay from the reticular formation to the cerebral cortex and have been shown to be critically involved in the regulation of cortical activity and behavioral state (Krnjevic and Phillips, 1963; Jones, 1993). Recently, it has been demonstrated that corticopetal cholinergic and GABAergic neurons in the basal forebrain fire rhythmically or are correlated with cortical EEG activity (Duque et al., 2000; Manns et al., 2000). Although it remains to be definitively established whether basal forebrain neuronal activity is contributory to or reflective of cortical activity, these findings provide strong evidence for a role for basal forebrain neurons in regulating extrathalamic cortical activation.

Although VTA GABA neuron firing was directly correlated with the sleep–wake cycle, there was no evidence of specific activity preceding or lagging each state or of synchronous activity in association with the cortical EEG. The mere concurrence of VTA GABA neuronal activity with cortical activation is not enough to establish causal or mechanistic connections between neuronal activity and electrocortical or behavioral activation. However, VTA GABA neurons may still be important regulators or switches of extrathalamic electrocortical or behavioral activation. VTA GABA neurons, including their projections and their inputs, similar to the role of SNr or SNC GABA neurons in regulating motor output, are in a critical position to modulate DA psychomotor output as integrators of convergent information from sensory, cortical, and limbic areas. The tonic glutamatergic input that regulates the firing of VTA GABA neurons may function in a manner similar to the role played by subthalamic inputs to SNr GABAergic neurons in mediating SNr inhibition of SNC DA neurons (Tepper et al., 1995). Alternatively, by virtue of their widespread axonal distribution and their wide dynamic range, VTA GABA neurons may be involved, independent of DA neurons, in the reticular activating system for extrathalamic regulation of cortical activity.

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