Turning a negative into a positive: ascending GABAergic control of cortical activation and arousal

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Gamma-aminobutyric acid (GABA) is the main inhibitory neurotransmitter in the brain. Recent technological advances have illuminated the role of GABAergic neurons in control of cortical arousal and sleep. Sleep-promoting GABAergic neurons in the preoptic hypothalamus are well-known. Less well-appreciated are GABAergic projection neurons in the brainstem, midbrain, hypothalamus, and basal forebrain, which paradoxically promote arousal and fast electroencephalographic (EEG) rhythms. Thus, GABA is not purely a sleep-promoting neurotransmitter. GABAergic projection neurons in the brainstem nucleus incertus and ventral tegmental nucleus of Gudden promote theta (4–8 Hz) rhythms. Ventral tegmental area GABAergic neurons, neighboring midbrain dopamine neurons, project to the frontal cortex and nucleus accumbens. They discharge faster during cortical arousal and regulate reward. Thalamic reticular nucleus GABAergic neurons initiate sleep spindles in non-REM sleep. In addition, however, during wakefulness, they tonically regulate the activity of thalamocortical neurons. Other GABAergic inputs to the thalamus arising in the globus pallidus pars interna, substantia nigra pars reticulata, zona incerta, and basal forebrain regulate motor activity, arousal, attention, and sensory transmission. Several subpopulations of cortically projecting GABAergic neurons in the basal forebrain project to the thalamus and neocortex and preferentially promote cortical gamma-band (30–80 Hz) activity and wakefulness. Unlike sleep-active GABAergic neurons, these ascending GABAergic neurons are fast-firing neurons which disinhibit and synchronize the activity of their forebrain targets, promoting the fast EEG rhythms typical of conscious states. They are prominent targets of GABAergic hypnotic agents. Understanding the properties of ascending GABAergic neurons may lead to novel treatments for diseases involving disorders of cortical activation and wakefulness.

Keywords: wakefulness, gamma rhythm, theta rhythm, EEG, arousal, hypnotics

Introduction

Our current understanding of the brain and the mechanisms involved in switching between different behavioral states is based on the investigational tools available to researchers. Thus, the development of histochemical and immunohistochemical methods to identify monoaminergic and cholinergic neurons, together with biochemical methods to study their metabolism, led to the concentration of neuropsychiatric research on these neurotransmitter systems in the latter
part of the 20th century (1). Similarly, in the sleep-wake field, much research focused on the role of the monoaminergic and cholinergic neurotransmitter systems (2), leading to an influential theory on the mechanisms which control switching between non-REM and REM sleep (3, 4). By contrast, much less attention was paid to the control of the sleep-wake cycle by the more prevalent gamma-aminobutyric acid (GABA)ergic and glutamatergic systems due to their later discovery and the lack of tools to manipulate and record their activity.

The sleep-promoting effects of allosteric agonists of the GABA<sub>Α</sub> receptor are well-established (5). However, we have only recently discovered the location and properties of GABAergic neurons controlling the sleep-wake cycle. The sleep-promoting action of GABAergic neurons located in the preoptic hypothalamus (6–8) is now well-known and accepted (9). More recently, other groups of sleep-promoting GABAergic neurons in the lateral hypothalamus (melanin-concentrating hormone neurons) and brainstem [parafacial zone; (10)] have been identified. It is less well-appreciated that there are several groups of fast-firing, subcortical GABAergic neurons with ascending projections, which promote wakefulness and cortical activation. Given that GABA is normally an inhibitory neurotransmitter in the adult brain, this promotion of cortical activation may seem paradoxical. However, the important concept to be grasped is that these ascending GABAergic neurons do not exert a tonic inhibition of their cortical and thalamic targets, but rather, they sculpt the pattern of activity of their targets in favor of the fast oscillations of firing typical of brain-activated states. Here, we summarize our current knowledge of these neurons, which recent evidence suggests may be critical in promoting cortical activation during wakefulness and REM sleep.

We first describe the technological advances that have allowed investigation of these GABAergic cell populations, followed by a review of the properties and likely functions of each group of GABAergic neurons.

**Novel Technological Tools Have Allowed Us to Identify, Record from, and Selectively Manipulate the Activity of GABAergic Neurons Controlling the Sleep-Wake Cycle**

In the last two decades, rapid advances in technology have allowed us for the first time to selectively identify and manipulate the activity of GABAergic neurons involved in control of the sleep-wake cycle. These techniques are essential since the GABAergic neurons involved in these processes are often interspersed among other neurons utilizing different neurotransmitters. Immunohistochemical staining for the enzyme which synthesizes GABA, glutamic acid decarboxylase (GAD), or staining for GABA itself, allowed the precise study of the distribution of GABAergic neurons in sleep-wake controlling regions of the brainstem, hypothalamus, and basal forebrain (BP) (6, 11–16). When combined with Fos immunohistochemistry to stain the nuclei of neurons which were recently active (i.e., expressed this immediate-early gene product), these immunohistochemical techniques proved invaluable in revealing the location of sleep- or wake-active GABAergic neurons (13–15).

While useful in determining the location of wake-active GABAergic neurons, Fos immunohistochemistry cannot reveal their firing rate or pattern, due to its limited temporal resolution. For this, electrical recordings in intact animals are essential. Heroic juxtacellular unit recordings in anesthetized and awake rats allowed characterization of the discharge of identified GABAergic neurons across the sleep-wake cycle for the first time (17–20). In this technique, recordings of extracellular action potentials from individual neurons are performed with glass micropipettes containing neurobiotin, which enters the recorded neuron, allowing subsequent immunohistochemical identification using co-staining for GAD or other markers such as parvalbumin (PV), a calcium binding protein expressed in subsets of fast-firing GABAergic neurons in many brain areas (21–24).

Extracellular unit recordings in vivo are essential for establishing the normal firing patterns of GABAergic neurons and correlating them with the electroencephalographic (EEG) activity and behavior. However, they have a limited ability to reveal the underlying cellular mechanisms, i.e., the ion channels and neurotransmitter receptors which are the targets of most pharmacological agents used clinically to modulate brain activity. To determine these cellular mechanisms, intracellular recordings of identified GABAergic neurons are required. The technically demanding GAD stain proved difficult to perform post hoc after in vitro electrophysiological recordings. Thus, to determine the intrinsic membrane properties of GABAergic neurons controlling the sleep-wake cycle, our laboratory took advantage of genetically modified mice expressing green fluorescent protein (GFP) under the control of the Gad1 (GAD67) promoter. Several different types of mice expressing GFP or other fluorescent markers under the control of part or all of the GAD promoter are now available, but the best validated are the GAD67-GFP knock-in mice generated by Yuchio Yanagawa and colleagues (25). In our laboratory, we confirmed that the sleep-wake cycle and cortical rhythms are normal in these mice (26, 27) and that GFP selectively labels essentially all GABAergic neurons in the BF and brainstem (24, 28). GFP also delineates other known GABAergic neuronal groups controlling the sleep-wake cycle, e.g., the circadian pacemaker in the suprachiasmatic nucleus and histaminergic neurons in the tuberomammillary nucleus of the hypothalamus (Figure 1) (29). A great advantage of these mice is that in vitro recordings can target GABAergic neurons online, prior to recording. Furthermore, retrograde tracing studies of the projections of these neurons do not require the use of GAD staining (29). Other genetically modified mice expressing a red fluorescent marker (tdTomato) in PV neurons also proved useful in characterizing the properties of PV-containing, cortically projecting, BF GABAergic neurons (24, 30, 31).

While recording the activity of GABAergic neurons in vivo and determining their properties in vitro provided important clues to their function and potential ways to pharmacologically manipulate their activity, tests of their functional role require selective stimulation and inhibition experiments. The development of mice expressing the bacterial enzyme Cre recombinase under the control of the GAD, vesicular GABA transporter (vGAT), or PV promoters further advanced our knowledge concerning the role of GABAergic neurons. Use of these mice allowed selective optogenetic (33) and designer receptor exclusively targeted by designer drugs (DREADD)
FIGURE 1 | GAD67-GFP knock-in mice delineate the location of GABAergic neurons involved in cortical arousal and sleep-wake control (24, 25, 28, 34). The location of select nuclei described in this review is illustrated, moving in a caudal to rostral direction through the mouse brain (A–H). Numbers in the bottom right corner indicate the location of these coronal sections with respect to Bregma (in mm). Green (GFP) fluorescence is present in cell bodies and fibers of GABAergic neurons. (A) Nucleus incertus (NI) GABAergic neurons involved in theta-rhythm generation are located near the midline of the central gray, above the medial longitudinal fasciculus (mlf), and ventromedial to the densely packed GABAergic neurons in the dorsal tegmental nucleus of Gudden (DTg). DTg neurons are involved in signaling head direction and project to the lateral mammillary body. (B) Ventral tegmental nucleus of Gudden (VTg) GABAergic neurons are clustered ventral to the mlf and the dorsal raphe (DR) nucleus. They innervate glutamatergic neurons in the medial mammillary body (see Figure 2). (C) GABAergic cell bodies and fibers delineate the substantia nigra pars reticulata (SNr), located lateral to the medial lemniscus (ml) and medial reticular nucleus (mRt). SNr GABAergic neurons represent the main output of the basal ganglia in rodents and tonically inhibit the motor thalamus and the centromedian-parafascicular nucleus (CM-Pf). (D) Tubero-mammillary (TMN) histamine neurons located lateral to the lateral mammillary nucleus (LM) also express GABAergic markers and may release GABA. (E) Thalamic reticular nucleus (TRN) GABAergic neurons surround and inhibit almost all thalamic relay nuclei. More ventrally and medially are the zona incerta (ZI) GABAergic neurons, which project to higher-order thalamic nuclei and the neocortex. The internal capsule (ic) separates these nuclei from GABAergic neurons in the globus pallidus (GP). (F) The master circadian pacemakers in the suprachiasmatic nucleus (SCN) of the hypothalamus are GABAergic. 3V = 3rd ventricle. (G) Caudal/intermediate nuclei of the basal forebrain contain many large-sized GABAergic neurons which project to the neocortex and regulate gamma oscillations and wakefulness. acp, anterior commissure, posterior part; HDB, horizontal limb of the diagonal band; MCPO, magnocellular preoptic nucleus; SI, substantia innominata; VP, ventral pallidum. (H) Rostral nuclei of the basal forebrain, the medial septum (MS), and vertical limb of the diagonal band (VDB) contain GABAergic septohippocampal neurons regulating hippocampal theta and gamma rhythms. Scale bars: (A,B,D,F) 0.25 mm; (C,G,H) 0.5 mm; (E) 0.75 mm.
approaches (34). In these techniques, Cre recombinase-dependent (double-floxed), viral vector mediated transduction allows the introduction of excitatory and inhibitory channel/pumps, which can be activated by light (optogenetics) or modified G-protein coupled cholinergic receptors which are activated by a normally inert drug, clozapine-N-oxide (DREADD technique) into selected groups of neurons, i.e., GABAergic or PV neurons containing Cre Recombinase. Work currently underway using these approaches (35–37) has provided important evidence supporting a role of BF GABAergic/PV neurons in the promotion of wakefulness and cortical gamma rhythms.

In the following, we describe what is currently known about subcortical, ascending GABAergic projection neurons which are active during wakefulness (and rapid-eye-movement sleep) and regulate wakefulness, as well as the higher frequency rhythms typical of brain-activated states. We do not cover GABAergic neurons which are primarily sleep-active (10, 20, 38–40) or those involved in the descending control of muscle tone (41, 42). We also do not cover GABAergic interneurons in the hippocampus and neocortex in detail, although they are extremely important in generating fast EEG rhythms (43–45) and are a target of ascending GABAergic projections (46–48). We begin in the brainstem and continue rostrally along the dorsal and ventral pathways of the ascending reticular activating system (ARAS), which include the thalamus and BF as their final nodes.

Brainstem GABAergic Neurons Controlling Theta Rhythms

The brainstem has a relatively small number of ascending, wake/REM-active GABAergic neurons (49). However, a large number of locally projecting, GABAergic interneurons surround and control the activity of ascending brainstem cholinergic, monoaminergic, and glutamatergic systems involved in sleep-wake control (13, 14, 18, 28, 49). In this section, we focus on two brainstem nuclei with GABAergic projection neurons, the nucleus incertus (NI) and the ventral tegmental nucleus of Gudden (VTg) (Figures 1A,B), both of which are involved in control of theta (4–8 Hz) rhythms which dominate the cortical EEG in rodents during active wakefulness and REM sleep (50), and are important in synchronizing the activity of brain areas involved in spatial navigation and memory formation (Figure 2).

Nucleus Incertus

Several lines of evidence suggest that NI GABAergic neurons containing the neuropeptide relaxin-3 are involved in control of theta rhythms. Anatomical studies revealed that GABAergic, relaxin-3 positive neurons in the NI project to regions involved in theta rhythm generation such as the supramammillary nucleus, medial septum/diagonal band (MS/DB), and the hippocampus (52–55) (Figure 2A). Other projections target intralaminar thalamus, hypothalamus, and amygdala. Relaxin-3 positive fibers in the MS/DB contact the cholinergic and GABAergic/PV neurons projecting to the hippocampus, which are known to act as pacemakers for theta rhythm (56). Ultrastructural analysis indicates that relaxin-3 positive terminals form symmetrical contacts typical of GABAergic synapses (56). Conversely, NI receives return GABAergic projections from the septum and horizontal limb of the diagonal band (57) (Figure 2A).

The anatomical projections of NI neurons led to direct physiological tests of a role for NI in control of theta rhythm. Electrical stimulation of the NI in urethane-anesthetized rats induced theta rhythm in the hippocampus (58). Conversely, electrolytic lesions or pharmacological inhibition of the NI abolished theta rhythm evoked by stimulation of the neighboring pontine reticular formation (PnO) in urethane anesthetized animals (58).

How might NI neurons facilitate hippocampal theta rhythms? Initial single-unit recordings revealed that the majority of NI neurons did not fire rhythmically or fired rhythmically at frequencies (13–25 Hz) higher than theta rhythm frequencies (58), indicating that they are unlikely to be involved in converting tonic reticular input into phasic theta frequency firing. However, more recent findings suggest that there are two populations of NI neurons, a relaxin-3 positive subpopulation which is excited by corticotropin-releasing factor (CRF) and exhibits strong phase-locked firing with the ascending phase of hippocampal theta oscillations, and a relaxin-3 negative subpopulation which is unaffected by CRF and whose firing is not phase-locked with hippocampal theta (59). Taking together the anatomical and physiological findings, it seems plausible that the GABAergic control of arousal 

![Image](www.frontiersin.org)
VTg

Another ascending brainstem projection involved in theta generation involves the VTg region originally described by the German psychiatrist, Bernhard von Gudden (61). VTg consists of two small aggregations of GABAergic neurons located on either side of the midline between the dorsal and median raphe (28, 62–64) (Figure 2B). In vitro recordings in both urethane-anesthetized (65) and freely moving (66, 67) rats revealed that VTg neurons fire long-lasting bursts of action potentials at high-frequencies which are phase-locked with hippocampal theta rhythms. Our in vitro recordings from identified GABAergic VTg neurons in GAD67-GFP knock-in mice suggest that these long-lasting bursts are due to a low-threshold calcium spike and subsequent activation of a long-lasting calcium-activated sodium conductance (68, 69). Like other ascending GABAergic neurons, VTg neurons are very fast-firing in vitro (maximal firing rate >200 Hz) and have narrow action potentials with brief afterhyperpolarizations (68, 69). They also exhibit strong afterhyperpolarization-activated cation currents (H-currents), which are often present in rhythmically active neurons.

Unlike NI neurons, VTg neurons do not have projections to the structures most commonly linked to theta-rhythm generation, the medial septum, supramammillary nucleus, or hippocampus. Instead, the VTg shares strong reciprocal connections with glutamatergic neurons of the medial mammillary body (MMB, Figure 2B) (63, 70–72). A parallel GABAergic ascending projection system involved in transmitting head-direction information arises in the dorsal tegmental nucleus and projects to the lateral mammillary nucleus (73). As with the VTg, single-unit recordings from the MMB have reported neurons which fire rhythmic bursts in synchrony with hippocampal theta rhythm (74, 75), although these bursts are shorter in duration and occur at a different phase with respect to hippocampal theta rhythms. Like VTg neurons, MMB neurons have low-threshold calcium spikes, which underlie the theta bursts recorded in vivo (68, 76).

The intrinsic membrane properties, neurotransmitter phenotype, and reciprocal connections of VTg and MMB neurons suggest a mechanism which may generate their theta activity (68): bursts of action potentials in glutamatergic MMB neurons lead to depolarization of VTg neurons and activation of low-threshold calcium channels; calcium influx into VTg neurons activates a calcium-activated cation conductance which prolongs the burst and increases the number of action potentials; long-lasting bursts in VTg neurons lead to a strong GABAergic inhibition of MMB neurons, de-inactivating the low-threshold calcium channels in MMB neurons, which are then activated once the VTg-mediated hyperpolarization subsides, restarting the cycle. Strong afterhyperpolarization-activated cation currents in VTg neurons help maintain the rhythmic firing by providing a depolarizing influence during the intraburst interval (68). Descending inputs to the MMB from the subiculum may act to synchronize theta rhythms in the hippocampus with those in the MMB-VTg circuit (74, 77, 78).

The major ascending output of the MMB innervates the medial and ventral parts of the anterior thalamus via axons which ascend within the mammillothalamic tract. Theta burst neurons have been recorded in the anterior thalamus, in particular in the ventral part targeted by the MMB (79, 80). The anterior thalamus in turn projects to the cingulate cortex and presubiculum, which is connected with the entorhinal cortex, one of the two main afferent inputs to the hippocampus. Thus, the VTg-MMB system may act as a theta rhythm generating system for the neural circuit described by Papez (51, 65, 79, 81, 82) (Figure 2B). Damage to the MMB—thalamus pathway in Korsakoff’s syndrome or via stroke results in diencephalic amnesia (73, 83). Similarly, neurotoxic lesions of the VTg in animals impair memory formation and learning (84).

Summary

Two ascending GABAergic brainstem nuclei, the NI and VTg, play key roles in the generation of theta rhythms during waking and REM sleep (Figure 2). The NI promotes theta generation in the supramammillary-medial septal–hippocampal system involved in formation of spatial maps of the environment, whereas the VTg-MMB system promotes theta rhythms in the anterior thalamus head-direction system (85). Integration of information from these two systems allows an animal to represent its position within the environment, a key requirement for spatial and episodic memory formation/retrieval during waking and REM sleep (86).

Ventral Tegmental Area GABAergic Neurons Involved in Arousal and Reward

In addition to the well-known dopaminergic neurons involved in reward and addiction, the midbrain ventral tegmental area (VTA) nucleus contains a substantial percentage (~20–35%) of GABAergic neurons (87–89). Important recent studies suggest that VTA GABAergic projection neurons, VTA GABAergic interneurons, and GABAergic inputs from the rostromedial tegmental nucleus (RMTg) are all involved in reward and reward-related arousal processes. VTA GABAergic projection neurons target widespread forebrain targets including the BF/preoptic area, amygdala, mediodorsal thalamus, and lateral hypothalamus, as well as arousal nuclei of the brainstem such as the dorsal raphe and deep mesencephalic nuclei (90). Sparser but functionally important projections target the prefrontal cortex (PFC) (87, 90, 91) and nucleus accumbens (NAcc) (90, 92, 93). A portion of the VTA GABAergic projection to PFC synapses onto neocortical GABAergic interneurons of unknown subtype (87), whereas the GABAergic VTA projection to NAcc preferentially targets cholinergic interneurons (94), suggesting that VTA GABAergic projection neurons indirectly modulate the activity of principal neurons in these structures.

VTA GABAergic Neurons are Fast-Firing and Electrically Coupled

In vivo extracellular recordings in halothane-anesthetized rats revealed that GABAergic VTA neurons exhibit a rapid (~19 Hz), cluster-type discharge pattern, short-duration action potentials,
and lack of accommodation of firing during prolonged depolarizations (93). They were activated antidromically and orthodromically by stimulation of the internal capsule, confirming that they are projection neurons and were inhibited by electrical stimulation of the NAcc. Intracellular, sharp-electrode recordings in vivo revealed they had relatively depolarized resting membrane potentials (~62 mV) and small action potentials (68 mV). Post hoc staining for neurobiotin and GABA confirmed that they were GABAergic and not dopaminergic (93). The expression of connexin-36 mRNA and protein, inhibition of prolonged discharges induced by high-frequency stimulation of the internal capsule by gap junction blockers, and the presence of neurobiotin dye coupling all suggest that VTA GABAergic neurons are electrically coupled, facilitating synchronization of their activity (95–97). In vitro recordings in GAD67-GFP knock-in mice indicated that VTA GABAergic neurons can be distinguished from dopamine neurons by their narrow action potentials and lack of an A-type potassium current (89).

The Discharge of VTA GABAergic Neurons is Associated with Arousal

In unrestrained, unanesthetized rats, VTA GABAergic neurons discharged even more rapidly than in anesthetized animals (29 ± 6 Hz during active waking), and their discharge was markedly elevated during the onset of movement and prior to brain stimulation reward, suggesting a role in arousal/attention (98, 99). Firing rates during movement could reach as high as 100–200 Hz for 10–20 s (98). Consistent with the initial report (93), the discharge of VTA GABAergic neurons was strongly suppressed by deep chloral hydrate, ketamine, or halothane anesthesia, and was reduced during non-REM sleep (98). During REM-sleep, VTA GABAergic neurons increased their mean firing rate beyond that observed during active waking to ~52 Hz. Furthermore, during REM sleep enhanced firing was correlated with EEG gamma band activity. However, to date, a role for VTA GABAergic neurons in controlling particular EEG frequency bands has not been tested using selective excitation and inhibition experiments.

VTA GABAergic Neurons are Excited by Wake/Arousal Promoting Neurotransmitters

Whole-cell recordings from two groups of non-dopaminergic, putative GABAergic VTA neurons demonstrated that they are strongly excited by the wake-promoting orexin/hypocretin peptides (100, 101). Putative VTA GABAergic neurons are also excited by other arousal/stress-related neuromodulators such as histamine, CCK, and substance P (101, 102). These studies did not distinguish effects on projection neurons from those on local GABAergic interneurons, an important future direction.

Role of VTA GABAergic Neurons in Reward Processes

Recent work has elaborated on the role of VTA GABAergic neurons in reward-related processes (Figure 3A). This work suggests that motivationally important information from the PFC is conveyed to VTA GABAergic neurons which target NAcc. Anatomical studies showed that PFC projections to VTA selectively target GABAergic neurons which project to the NAcc, but not those projecting back to the PFC (103). In vivo recordings demonstrated a task-dependent increase in coherence at gamma band frequencies between PFC and VTA during a working memory task, likely reflecting transmission of information from PFC to fast-firing VTA GABAergic neurons (104). The projection of VTA GABAergic neurons to the NAcc selectively targets and inhibits cholinergic NAcc interneurons (94). Activation of the VTA GABA→NAcc pathway caused a pause in the discharge of cholinergic interneurons, resembling that previously observed in animals learning stimulus-outcome associations. Furthermore, optogenetic activation of this pathway enhanced discrimination of a motivationally important stimulus paired with an aversive outcome (94).

VTA GABAergic Interneurons Control Dopaminergic activity

Many drugs of abuse, such as opiates (105) and benzodiazepines (106), inhibit VTA GABAergic interneurons, leading to increased dopamine release via disinhibition. Conversely, optogenetic activation of VTA GABAergic neurons in vivo disrupts reward consummatory behavior (107). VTA GABAergic neurons, likely interneurons, identified using optogenetic activation, inhibited neighboring dopaminergic neurons, and encoded the presence of expected rewards (108). Disruption of this coding by drugs of abuse likely contributes to mechanisms of addiction.

The GABAergic Neurons of the RMTg Convey Aversive Information to VTA Dopaminergic Neurons

The posterior part of the VTA and its extension into the tegmental region of the reticular formation (collectively known as the posterior VTA or RMTg nucleus) contains a population of ascending GABAergic neurons which project to and regulate the activity of more rostrally located VTA dopaminergic neurons (109, 110). Unlike VTA GABAergic neurons, however, RMTg neurons do not have substantial projections to the forebrain (109). Recent studies implicate these GABAergic neurons in aversive behavior and reward prediction through a basal ganglia→lateral habenula→RMTg circuit (111, 112) (Figure 3B). Lateral habenula neurons are excited by stimuli that indicate the absence of a reward (113), through an excitatory pathway arising in the basal ganglia (112), leading to excitation of RMTg GABAergic neurons and subsequent inhibition of reward-related midbrain dopaminergic neurons and pedunculopontine neurons (111, 114). Like VTA GABAergic interneurons, RMTg neurons are inhibited by μ-opioids and cannabinoids (115). Further experiments suggested that the RMTg may be the most important site for disinhibitory effects of these compounds on dopaminergic activity (116).

Summary

Ventral tegmental area GABAergic projection neurons promote reward and reward-related arousal through their mesoaccumbens and mesocortical projections, which notably target interneurons in both structures. Conversely, VTA GABAergic interneurons and RMTg GABAergic inputs, encoding expected reward or the absence of rewarding stimuli, exert a tonic inhibitory effect on VTA dopaminergic neurons (Figure 3).
GABAergic control of arousal

 Ascending Hypothalamic GABAergic Systems Controlling the Thalamus and Neocortex

Retrograde tracing studies identified four major hypothalamic systems projecting to the cerebral cortex (117). Of these four systems, three subsets of neurons in the tuberal part of the lateral hypothalamic contain GABA: GABAergic neurons in the ventral part of the zona incerta (ZI), melanin concentrating hormone (MCH) neurons, and tuberomammillary histaminergic neurons. MCH neurons are sleep-active and sleep-promoting and are therefore not considered further in this section. Although histaminergic neurons promote arousal (118) and they contain the biosynthetic machinery for GABAergic transmission (119), the functional role is poorly understood (120). Therefore, we concentrate in this section on ZI GABAergic projection neurons.

Glutamic acid decarboxylase immunostaining coupled with retrograde tracing confirmed the presence of a group of GABAergic projection neurons in the ZI (121). Many of these neurons may also contain the neuropeptide α-melanocyte stimulating hormone (122, 123), or a closely related peptide (124). GABAergic ZI neurons project heavily to the thalamus (125) as well as to the neocortex (121). Immunostaining also revealed a population of PV-containing neurons in this same area, suggesting that as in other areas, these GABAergic neurons are likely to be fast-firing. In vitro recordings from the ventral ZI region revealed that most neurons discharged spontaneously at high rates (9.3 Hz median firing rate) and also exhibited rhythmic firing (126). In the same study, in vivo recordings in urethane-anesthetized animals revealed firing at 3–4 Hz. However, in a different study under light ketamine or urethane anesthesia, ventral ZI neurons discharged much more rapidly, with mean rate of 26 Hz (127). The role of ZI projections to the cortex has still not been explored in any detail, but recent studies suggest an important regulation of higher order sensory nuclei of the thalamus (see next section).

GABAergic Neurons Controlling Thalamocortical Activity during Wakefulness

The midline and intralaminar thalamic nuclei represent the final node of the dorsal portion of the reticular activating system (9). These nuclei, and the primary thalamic sensory relay nuclei, are under strong inhibitory control from GABAergic neurons in several different regions. The most prominent, widespread, and well-known GABAergic input arises from the thalamic reticular nucleus (TRN). A more restricted input to the midline centromedian-parafascicular nucleus (CM-Pf) and motor thalamus (ventrolateral and ventromedial nuclei) originates from the output of the basal ganglia, the globus pallidus, pars interna (GPI) and the substantia nigra, pars reticulata (SNr) (128–130). Other GABAergic inputs to the midline thalamus arise in the ZI (125) and BF (131). In general, all of these GABAergic inputs maintain a high rate of tonic inhibition in the thalamic relay nuclei, which is likely important in suppressing unimportant information and unnecessary motor activity. However, in situations requiring high attention and responses to important situations, this tonic input is transiently suppressed allowing enhanced arousal/attention (disinhibition of higher-order nuclei) and sensorimotor transmission (disinhibition of first-order sensory and motor nuclei).

Thalamic Reticular Nucleus

In addition to being crucially involved in generating sleep spindles during non-REM sleep (132, 133), TRN neurons maintain a high rate of tonic firing during wakefulness (134–136), which serves to prevent unimportant sensory information being transmitted through the thalamic relay nuclei to the cortex. TRN neurons receive excitatory inputs from noradrenergic and serotonergic neurons which maintain this high firing rate during wakefulness (137). Reduced activity of sensory-related TRN neurons occurs prior to correct performance in attention tasks (138), leading to
disinhibition of relay neurons (139). The origin of this reduced activity of TRN neurons is unclear. TRN receives inhibitory GABAergic/PV projections from globus pallidus (140), substantia nigra pars reticulata (141), and BF GABAergic/PV neurons (142, 143). Input from BF GABAergic/PV neurons is particularly interesting, considering the strong inputs to these BF neurons from PFC regions involved in processing novelty (144–146) and their strongly state-dependent firing (see next section). Other experiments have suggested that TRN may be involved in gamma-band oscillations through entrainment of the activity of thalamocortical neurons (147–149).

Basal Ganglia GABAergic Input to the Thalamus

The two main GABAergic output nuclei of the basal ganglia are the GPi and SNr. GPi and SNr project to thalamic motor output nuclei and to the centromedian-parafascicular nucleus (CM-Pf), which is one of the “non-specific” thalamic nuclei regulating the level of arousal through widespread, diffuse projections to the cortex (150). CM-Pf also has a prominent projection to the striatum, enhancing activity in cortico-basal ganglia-thalamocortical circuits. Electrical or pharmacological activation of CM-Pf in rodents enhances behavioral arousal and recovery from anesthesia, supporting a role in control of consciousness (151–154). Furthermore, changes in CM-Pf activity precede loss of consciousness caused by anesthetics or transitions into sleep (155). Thus, GABAergic control of CM-Pf by the basal ganglia output nuclei is likely to be extremely important in control of arousal and consciousness.

Neurons in GPi and SNr express very high levels of the α1 subunit of the GABA_A receptor, the target of the hypnotic, zolpidem (156–159) in rodents and in humans (160). Interestingly, in brain damaged patients, it has been hypothesized that the activity of these neurons is pathologically enhanced, resulting in an inhibition of thalamic and pedunculopontine neurons and loss of consciousness (161). Thus, suppression of their activity by zolpidem may result in a paradoxical arousing effect in some patients (162–166).

ZI GABAergic Input to the Thalamus

In addition to direct projections to the cortex, ZI GABAergic projection neurons innervate the thalamus, superior colliculus, and brainstem (125, 167). Thalamic projections of ZI GABAergic neurons are particularly strong to higher-order nuclei, where they effect a tonic inhibition of sensory transmission (125, 167). In the rodent, ZI neurons receive a strong input from trigeminal axons transmitting sensory information from the whiskers, leading to excitation of ZI GABAergic neurons and preventing excitation of the posterior thalamic group neurons (127). This feed-forward inhibition can be overcome by excitation of the motor cortex and activation of an intra- INCIDENTAL GABAergic circuit (168). Many other cortical areas also converge on ZI GABAergic neurons, which through their widespread cortical, thalamic, and brainstem projections are in a position to globally modulate brain arousal (169). Cholinergic stimulation in vitro or in vivo in anesthetized rats inhibited the firing of ventral ZI neurons, suggesting that high arousal states involving increased acetylcholine release lead to suppression of ZI neuronal firing and promotion of sensory transmission (126).

Summary

GABAergic projection neurons in the TRN, GPi, SNr, and ZI act to tonically inhibit various thalamic relay neurons during wakefulness. Suppression of their firing by cortical or subcortical inputs is a powerful mechanism to increase thalamic activity and thereby increase attention, arousal, sensory processing, and motor activity in a context-dependent manner.

Cortically Projecting BF GABAergic Neurons Promote Cortical Activation and Wakefulness

The BF represents the final node of the ventral portion of the brainstem ARAS. Rostral BF neurons in the medial septum and vertical limb of the diagonal band project to the hippocampal formation, whereas intermediate and caudally located BF neurons in the horizontal limb of the diagonal band, magnocellular preoptic area, ventral pallidum, substantia innominata, and nucleus basalis project to the neocortex, as well as to the thalamus, lateral hypothalamus, and brainstem (11, 12, 16, 46, 47, 170). Many studies have focused on the arousing effects of cortically projecting cholinergic neurons in this region (171), which are among the first to degenerate in Alzheimer’s disease (172). However, recent studies suggest that neighboring GABAergic projection neurons and interneurons may be equally important in cortical activation. Juxtaglomerular labeling experiments in vivo identified a significant minority of BF GABAergic neurons, which are fast-firing (20–60 Hz) and increase their firing rate during wakefulness and REM sleep (19). Recent preliminary experiments showed that pharmacogenetic (DREADD) stimulation of BF GABAergic neurons strongly enhances wakefulness (37). Similarly, initial experiments suggest that optogenetic stimulation of BF PV neurons is wake-promoting (35), in addition to playing a role in control of cortical gamma oscillations (36).

Anatomy and Projections of BF GABAergic Neurons

Anterograde tracing studies coupled with staining for GAD showed that cortically projecting GABAergic neurons make up approximately one-third of the BF projection to the cortex. Importantly, BF GABAergic projections to the neocortex and hippocampus prominently target interneurons containing PV and somatostatin, which are thought to be involved in gamma and theta oscillations (46–48). Caudal/intermediate BF GABAergic neurons projecting to the neocortex, as well as more rostrally located BF GABAergic neurons projecting to the hippocampus, preferentially target GABAergic interneurons (46, 47, 173), in particular PV-containing fast-spiking interneurons (174) involved in theta (175) and gamma oscillations (44, 45). Somatostatin-containing cortical interneurons which target the dendrites of pyramidal neurons and appear tuned for theta-frequency firing are also a target of BF cortically projecting GABAergic neurons, as are cortical interneurons containing calbindin (173). BF PV and GABAergic neurons also project to the TRN (142, 143) and less prominently to the mediodorsal nucleus (131, 170). At present, it is unclear if only BF PV/GABAergic neurons or also other types of BF non-PV/GABAergic neurons
project to TRN. The function of the BF projection to TRN is currently unresolved but this projection may be involved in attentional suppression of TRN discharge (see previous section) and/or suppression of spindle-related bursting in TRN during wakefulness and REM sleep.

**Subtypes of Cortically Projecting BF GABAergic Neurons**

Several subtypes of cortically projecting GABAergic neurons have been identified in the BF based on the expression of different neurochemical markers (Figure 4). The most well-known are neurons containing the calcium-binding protein PV. Other, largely separate subsets express the neurokinin-3 receptor (176) and the potassium channel Kv2.2 (177, 178). In *in vitro* recordings suggest that large-sized, cortically projecting GABAergic and PV neurons can be subdivided into two groups based on the amplitude and kinetics of their H-current (24).

**PV Neurons: Anatomy and In vivo Recordings**

The vast majority of PV-containing neurons in the BF are GABAergic (16, 24). PV is a marker for cortically projecting BF GABAergic neurons (16). In the rat, immunohistochemical staining for PV labeled approximately 90% of GAD-stained neurons, retrogradely labeled from the orbitofrontal and somatosensory cortex (16). In the GAD67-GFP knock-in mouse, PV was observed in ~25% of large (>20 μm) diameter, putative cortically projecting GABAergic neurons, whereas, overall PV was present in 6.7% of all BF GABAergic neurons (24). PV-containing GABAergic neurons in the rostral part of the BF (medial septum and vertical limb of the diagonal band) project to the hippocampus, and are critically involved in control of hippocampal theta rhythm (179, 180), and associated hippocampal gamma rhythms, whereas PV-containing neurons in intermediate and caudal BF regions project to the neocortex and regulate neocortical gamma oscillations (35). Two populations of medial septal PV neurons discharge at opposing phases of the hippocampal theta rhythm. *Post hoc* identified PV neurons recorded in anesthetized rats discharged rapidly in bursts or clusters in association with cortical activation induced by tail pinch (17). In unanesthetized mice, two optogenetically identified PV neurons discharged irregularly in the gamma range (20–60 Hz) during wakefulness and REM sleep (36). Transduction of BF PV fibers with channelrhodopsin2-enhanced yellow fluorescent protein fusion proteins revealed BF PV neurons appose cortical PV interneurons, consistent with a role in control of cortical gamma oscillations (36). Direct tests of this hypothesis using optogenetic techniques demonstrated that both rhythmic and non-rhythmic stimulation of BF PV neurons preferentially enhanced cortical EEG power at gamma frequencies, whereas optogenetic inhibition of BF PV neurons reduced the ability of the cortex to respond at 40 Hz in response to a train of auditory stimuli delivered at 40 Hz (36). Together, these data strongly implicate BF PV neurons in the behavioral state-related increases in cortical gamma oscillations, which are observed during wakefulness and REM sleep (181, 182).

**PV Neurons: In vitro Recordings and Intrinsic Properties**

*In vitro* recordings from identified PV neurons using genetically modified mice expressing fluorescent markers, post hoc staining (183), or based on intrinsic membrane properties have revealed many similarities between MS/DB GABAergic/PV neurons projecting to the hippocampus and those projecting to the neocortex. Both groups are very fast-firing (24, 183), likely due to their expression of the delayed rectifier Kv3.1 (Kcnq1) potassium channel (184), and whose extremely fast kinetics enables fast repolarization of the action potential (185). Both groups of BF GABAergic/PV neurons exhibit a "depolarizing sag" during hyperpolarizing current pulses due to a hyperpolarization-activated cation current (H-current) (24, 183), a property which distinguishes them from cortical PV interneurons and TRN GABAergic/PV neurons. This current counteracts prolonged hyperpolarization and is also often present in neurons which show rhythmic firing, providing a depolarizing influence in the interburst interval following afterhyperpolarizations. Thus, this current may be important for the cluster and burst-like firing of these neurons recorded *in vivo* (17). In fact,
Neurokinin3 Receptor Immunoreactive Neurons

The Neurokinin3 Receptor (NK3R) is the most selective receptor for neurokinin B, produced from the precursor preprotachykinin (PPTB). PPTB is present in a small subset (5%) of projection neurons in the neostriatum which projects to the substantia innominata (188). Neurokinin B and NK3R are essential for normal reproduction (189). Thus, one plausible functional role for this population of BF GABAergic neurons is control of sexual arousal. Immunohistochemical staining in rats and in GAD67-GFP knock-in mice revealed that NK3R is present on a distinct subset of cortically projecting BF GABAergic neurons (176). About 92% of NK3 receptor positive neurons showed signals for GAD67 mRNA (176). Only 10–15% of NK3R expressing neurons were PV-positive and only 1.7% of PV-positive neurons were NK3R positive (176). About 1.7% of NK3R neurons were positive for calretinin and none contained calbindin, NPY, or somatostatin.

Kv2.2 Channel Immunoreactive Neurons

The potassium channel Kv2.2 is abundantly expressed in one group of BF GABAergic neurons (177). Less than 4% of Kv2.2 immunoreactive neurons were positively for PV, establishing this as a separate group of GABAergic neurons (178). Sleep deprivation experiments showed that this group of neurons expressed more Fos, suggesting that this is a wake-active group of GABAergic neurons. Furthermore, knockout of the Kv2.2 channel led to more activity and promotion of wakefulness over sleep (178).

Neurotransmitter Regulation of Wake-Promoting BF GABAergic Neurons

Our recent in vitro recordings in GAD67-GFP knock-in mice and PV-Tomato mice revealed that putative cortically projecting BF GABAergic and PV neurons are strongly excited by cholinergic inputs (31). Similarly, MS/DB GABAergic/PV neurons projecting to the hippocampus are excited by cholinergic agonists (190). Bath application of cholinergic agonists revealed that septohippocampal neurons are excited via M1 receptors (191) and indirectly by activation of nicotinic receptors on glutamatergic neurons (192). Two subpopulations of putative neocortically projecting BF GABAergic neurons (large Ih, small Ih) are excited via M1 muscarinic receptors and M1 muscarinic receptors, respectively (31). Optogenetic stimulation of cholinergic fiber terminals revealed an additional excitatory effect mediated by nicotinic receptors (31). Interestingly, blockade of cholinergic receptors in the BF blocked the ability of optogenetic stimulation of cholinergic neurons to increase wakefulness, indicating that cholinergic modulation of behavioral state may depend on local interactions with BF GABAergic and/or glutamatergic neurons (193). Septohippocampal GABAergic/PV neurons are also excited by noradrenaline (194), histamine (195), serotonin (196), and orexin/hypocretins (197, 198). Preliminary experiments suggest that this is also true for putative neocortically projecting GABAergic neurons (32). Anatomical tracing studies revealed that PV containing BF neurons receive direct input from cortex (144) and dopaminergic inputs from the substantia nigra-VTA (199).

Summary

Several subsets of wake/REM active GABAergic neurons are present in the BF (Figure 4). They are strongly excited by wake-promoting neuromodulatory systems. Their projections target interneurons in the hippocampus and neocortex, as well as TRN GABAergic/PV neurons allowing control of cortical rhythms, attention, and wakefulness. The precise functional role of these different subtypes awaits further study but BF PV projection neurons appear to have a particular role in control of cortical gamma oscillations.

Wake-Active GABAergic Neurons are Fast-Firing, Whereas Sleep-Active GABAergic Neurons are Slow Firing

In vitro, most putative ascending subcortical GABAergic neurons are fast-firing neurons, both in terms of their spontaneous firing rate and their maximum firing frequency. Those which contain PV are even faster-firing and have extremely brief action potentials and afterhyperpolarizations (24). Similarly, in vivo, these subpopulations of GABAergic neurons show high rates of firing during waking and REM sleep. This fast-firing is at least partly due to the presence of Kv3.1 and Kv3.3 potassium channels. These channels are present in cortical interneurons, TRN neurons, and BF GABAergic neurons projecting to the hippocampus and neocortex. Knockout of Kv3.1 channels alone leads to a relatively mild arousal phenotype involving fourfold increased gamma (20–60 Hz) activity during wakefulness and reduced delta oscillations during all states (200). Given that GABAergic/PV neurons are thought to be involved in promoting gamma activity, these results suggest that developmental compensation occurs. Knockout of the Kv3.3 channel led to no discernible phenotype. However, double knockout mice lacking the genes encoding both of these proteins have severe deficits in sleep/wake behavior (201–203).

In contrast to the fast-firing of wake/REM sleep active GABAergic neurons, the majority of sleep-active, presumptive GABAergic neurons in the ventrolateral preoptic area, median preoptic area, and lateral hypothalamic area (MCH neurons) tend to be silent at rest and fire slowly (<15 Hz), even during sleep, the state when they discharge fastest (20, 38, 40, 203–205). These differences in firing rates suggest alternative complements of voltage-gated ion channels and neurotransmitter receptors in wake and sleep-active GABA neurons. Thus, differential pharmacological modulation may be possible.

Wake-Active GABAergic Neurons May Be a Prominent Target of Hypnotic Agents

As described in the previous section, most wake-promoting GABAergic neurons are neurons which exhibit a high spontaneous and maximal firing rate, consistent with a role in controlling the fast theta, beta, and gamma-band EEG oscillations typical of wakefulness. GABAergic hypnotic agents such as diazepam, zolpidem (Ambien™, α1 subunit selective agent), and eszopiclone...
Furthermore, genetic removal of GABA<sub>A</sub> receptors from histamine neurons on aminergic and cholinergic neuromodulatory systems. However, lesions or inactivation of these neuromodulatory systems have little effect on total amounts of sleep and waking [reviewed in Ref. (9)], suggesting that this explanation may not be correct.

Conversely, pharmocogenetic tools may be particularly useful for dissecting out their role and neuropharmacology is likely to be very important in understanding the properties of ascending GABAergic neurons. Thus, further study of these neurons to determine their functional role and neuropharmacology is likely to be very important in order to develop novel therapeutic compounds to modulate cortical activation, memory, reward, and sleep. Furthermore, understanding the properties of ascending GABAergic neurons may allow novel treatments for diseases involving disorders of cortical activation and wakefulness.

How do ascending subcortical GABAergic systems turn a negative (inhibitory postsynaptic effects) into a positive (cortical activation/arousal)? Ascending cortically projecting GABAergic neurons in the BE, VTA, and possibly also the ZI, target inhibitory neocortical interneurons, allowing disinhibitory effects and entrainment of fast cortical oscillations. Thalamic-targeting GABAergic neurons exert tonic inhibitory control, which can be suppressed in behaviorally important situations, potentiating thalamocortical transmission. Ascending GABAergic neurons exhibit narrow action potentials, brief hyperpolarizations, and are fast-firing. They often exhibit burst or cluster firing, and like cortical interneurons many are electrically coupled, properties which will enhance their action on their post-synaptic targets and enable them to synchronize their activity into fast oscillations. Most ascending, wake-active GABAergic neurons also express potassium channels and GABA<sub>A</sub> receptor subunits, which are the targets of anticonvulsant and hypnotic/anesthetic agents. Thus, further study of these neurons to determine their functional role and neuropharmacology is likely to be very important in order to develop novel therapeutic compounds to modulate cortical activation, memory, reward, and sleep. Furthermore, understanding the properties of ascending GABAergic neurons may allow novel treatments for diseases involving disorders of cortical activation and wakefulness.

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