AROUSAL, MOTOR CONTROL, AND PARKINSON’S DISEASE

Abstract
This review highlights the most important discovery in the reticular activating system (RAS) in the last 10 years, the manifestation of gamma (γ) band activity in cells of the RAS, especially in the pedunculopontine nucleus (PPN), which is in charge of the high frequency states of waking and rapid eye movement sleep. This discovery is critical to understanding the modulation of movement by the RAS and how it sets the background over which we generate voluntary and triggered movements. The presence of γ band activity in the RAS is proposed to participate in the process of preconscious awareness, and provide the essential stream of information for the formulation of many of our actions. Early findings using stimulation of this region to induce arousal, and the novel use of PPN deep brain stimulation for the treatment of Parkinson’s disease, although considerable work remains to be done.

Keywords
- Arousal
- Calcium channels
- Deep brain stimulation
- Mu rhythm
- Parkinson’s disease
- P13 potential
- P50 potential
- Readiness potential

Introduction
What happens when we respond to a sudden stimulus? What is required to execute voluntary movements, and what leads to involuntary movements, such as those in Parkinson’s disease (PD)? This review will address an issue not normally emphasized in the motor control and movement disorder literature, the background of information necessary for the execution of movements, both voluntary and unplanned.

In his book, I of the Vortex, Llinas proposed that what we call thinking is the evolutionary internalization of movement [1]. Llinas explored how the mind arose in evolution, and concluded that the brain’s control of organized movement gave birth to the generation of the mind. The simple but crucial point is that the evolutionary development of a nervous system is a property of actively moving creatures. Llinas suggested that prediction is the ultimate function of the brain and that the self is the centralization of prediction. The argument was advanced in linear fashion, supported by a fountain of information on the physiology of motor control [1]. One reason why this rationale is so persuasive is because the motor “card” is precisely what has been missing from play in a number of theories of consciousness and free will. The unsatisfactory nature of concepts of mindedness based on sensory perception, the uneasy discomfort generated by such hypotheses, is laid bare by this patently evident idea. While the argument is complex, the basic idea is simple, almost obvious. The theory advanced by Llinas is based on oscillations, from the oscillations of graded potentials along a nerve cell’s membrane, to the rhythmic activity of groups of nerve cells firing in phase, to the synchronization of rhythms in what are effectively analog brain systems.

If groups of neurons oscillate in phase or resonate, a global activity pattern results that provides the components for a transient construct of the external world [1]. This coherence in time is believed to form the neurological mechanism for perceptual binding. That is, to bring together the separate sensory components of a stimulus, say color, size, motion, and so on, that are processed in different regions of the cortex. This process not only serves to provide binding of sensory events, but there are coherent oscillations that subserve the precise temporal activation of muscles needed to execute a movement accurately. There is ample evidence for the coherent activation of neurons during sensory activation. For example, a visual stimulus produces coherent gamma (γ) band oscillations in the visual cortex [2, 3]. These oscillations are evident in widely separated cortical regions but they are coherent in time and frequency. That is, they are ideally manifested to perform a binding function.

Both the cortex and the thalamocortical system are endowed with intrinsic membrane properties such as sodium-dependent subthreshold oscillations or high threshold voltage-sensitive calcium channels, both of which can resonate at γ frequencies [4-6]. This means that sensory information arriving in an awake cortex and thalamus is superimposed on ongoing γ band oscillations. Basically, “specific” thalamic nuclei receive the “content” of sensory experience from primary sensory pathways and relay it to layer IV of the cortex, while “non-specific” thalamic nuclei receive the “context” of sensory experience from the...
reticular activating system (RAS), and relay it to layer I of the cortex. Temporal coherence is due to the coincidence in firing of the two pathways which, when activated together, will summate to trigger cortical output to the thalamus. The cortical output flows back down to “specific” and “non-specific” thalamic nuclei to set up a thalamocortical resonance. It is this resonance that is thought to underlie the process of perception. A common definition of consciousness is awareness, such that if there is no awareness, there is no consciousness [7]. This is fundamentally intertwined with the concept of free will. When choosing to make a movement, the implication is that there is a will that decides to engage the motor system to then induce the movement. But what happens before we make a movement?

We will pursue a new line of evidence that suggests that the preparation for movement begins at much lower levels than previously thought. The preamble to volition and response time is subcortical in origin, and arises from the moment we awaken until the moment we fall asleep [8].

The readiness potential, the mu (μ) rhythm, and the P50 potential

What do these three well-known waveforms have in common? The readiness potential (RP), the μ rhythm, and the P50 midlatency auditory evoked potential all are recorded in the same region of the cortex. The RP was originally described by Kornhuber as a negative direct current (DC) shift that began 600-800 ms before a voluntary, uncued movement [9]. The RP was present at maximal amplitude at the vertex in the region of the supplementary motor cortex and precentral cortex [9, 10]. Pioneering studies by Libet first showed that when people consciously set a goal to engage in a behavior, their conscious will to act begins “unconsciously” [11]. The studies of Libet employed the RP, which is known to have an early component that precedes the movement by as much as 1 s, and a late component that precedes the movement by 400 ms. Libet’s subjects were asked to move voluntarily, and were instructed to subjectively time the moment at which they felt the “will” to move as well as the onset of the actual movement. Figure 1 is a diagram of the vertex-recorded RP with the timing of the subjectively perceived “will” and perception of actual movement. The early and late phases of the RP preceded the “consciously” determined will to move by hundreds of milliseconds. These authors concluded that cerebral initiation of spontaneous, freely voluntary acts can begin “unconsciously”, before there is any subjective “conscious” awareness that a decision to act was initiated cerebral [11].

Even simple movements appear to be generated “subconsciously”, and the “conscious” sense of volition comes later [12]. Hallett described the details of studies showing that voluntary movements can be triggered with stimuli that are not “consciously” perceived, that movement may well occur prior to the apparent planning of the movement, and that not only the sense of the movement having occurred, but also the sense of willing the movement, happen before the actual movement [12]. However, Libet suggested that voluntary acts begin “unconsciously”, before there is subjective “conscious” awareness that a decision to act was initiated by the brain. This conclusion has been extrapolated to suggest that there is no free will. In response, Libet suggested that, although the movement was indeed initiated “unconsciously”, it was subject to veto once it reached consciousness [13]. This has been regarded as unsatisfactory and not answering the question of whether there is free will or not. The question is complex because so many factors influence the sense of volition such as the perception of time, the conditions under which the movement is executed, the perception of volition, etc. [12].

We proposed an alternative view simply by concluding that it is the interpretation of the results that assumes that the process preceding the movement is “unconscious” [8]. There is no evidence that this is the case. The preparation for movement generates brain processes that are clearly related to the intent, but these occur during waking, and therefore should be labeled “preconscious”, not “subconscious”. The replacement of the word “preconscious” for the word “subconscious” significantly alters the conclusion of these studies. That is, the conclusion should have been: “voluntary acts begin preconsciously, before there is subjective conscious awareness that a decision to act was initiated by the brain”. We will see below how the RAS is an ideal candidate for the process of preconscious awareness.

The μ rhythm was first described by Gastaut [14] as an 8-12 Hz wave present over the vertex and bilaterally across the precentral motor cortex, basically at the EEG C3, Cz (vertex), and C4 electrode placements. The μ rhythm is present when the body is at rest, but the rhythm is “suppressed” or “blocked” when the person performs a motor action, or, after practice, when the person views another or visualizes a motor action. That is, it is more likely that, just as with the alpha (α) rhythm, the μ rhythm is merely replaced by faster activity when the function of that region is called for; that the μ rhythm over the vertex and precentral cortex is shifted to higher frequencies, not “suppressed”, from idle speed to higher frequencies, when motor events are called for.

![Figure 1. The human readiness potential (RP). Representation of a vertex-recorded RP and the timing reported by subjects performing an uncued voluntary movement. The estimation of the sense of will (signified by “W”) or intent to move occurred well after the beginning of the RP, and the sensation of movement (signified by “M”) occurred even later, and well after the beginning of the RP as in the Libet study [11].](image)
The μ rhythm is also suppressed during tactile stimulation [15], as is the α rhythm [16]. Both the occipital α rhythm and the precentral μ rhythm have been referred to as "idle rhythms" or "resting state" activity [17]. The μ rhythm has been associated with somatosensory information, while a faster beta (β) rhythm in the precentral region has been associated with actual motor processing [18]. That is, these "idle" or "resting" rhythms are of slower frequencies and are "blocked" or "suppressed" (actually replaced) by higher frequencies when sensory or motor events are called for.

We should emphasize that the μ rhythm, like the RP, is present at the vertex (EEG Cz electrode location and also laterally at C3 or C4 depending on which hand or arm movement is being performed contralaterally). The vertex is also where the P50 potential is maximal. The human P50 potential is a midlatency auditory evoked response evoked by a click stimulus and recorded at the vertex [19]. The P50 potential peaks at a latency of 40-70 ms. The P50 potential was also referred to as the P1 potential because it is the first positive wave following the early brainstem auditory evoked response (BAER) that occurs at a 5-10 ms latency, and the primary auditory response, called Pa, at the superior temporal gyrus, that occurs at a 25 ms latency. The Pa response is directly related to the auditory lemniscal, "specific" thalamic nuclei, and the auditory cortex, while the P50 potential is related to the non-lemniscal, reticular, intralaminar or "non-specific" thalamic nuclei ascending pathways [19].

The human P50 potential has three main characteristics: 1) it is sleep state-dependent, that is, it is present during waking and rapid eye movement (REM) sleep, but is absent during deep slow-wave sleep (SWS) [20,21], so that it is present when the cortex is activated and the EEG shows high frequency activity and, of course, when the pedunculopontine nucleus (PPN) is active, 2) it is blocked by muscarinic cholinergic antagonists, such as scopolamine, so that it may be mediated, at least in part, by cholinergic neurons such as those in the PPN [22], and, 3) it undergoes rapid habituation at stimulation rates greater than 2 Hz, indicative of a "reticular" pathway. For example, the primary auditory cortex Pa potential can follow stimulus frequencies close to 20 Hz. However, the P50 potential cannot follow such high frequencies of stimulation, implying that it is not generated by a primary afferent pathway, but by multisynaptic, low security synaptic elements of the RAS [20-22].

The P50 potential decreases and disappears with progressively deeper stages of sleep and then reappears during REM sleep at full amplitude [20, 23]. None of the earlier latency auditory evoked potentials (BAER or Pa primary auditory cortex potentials) possess this sleep-dependent characteristic. This suggests that the P50 potential is functionally related to states of arousal. In addition, the P50 potential is most likely generated, at least in part, by a cholinergic mesopontine cell group such as the PPN, since its cells are preferentially active during waking and REM sleep, but inactive during SWS [19, 24-26]. The P50 potential has been localized to the frontal lobes in the region of the vertex, a source we confirmed using magnetoencephalography [27]. Studies in cats [20, 21] and rats [28-30] showed that the feline "wave A" and the rodent P13 potential are the animal equivalents of the human P50 potential. These waveforms can be blocked by agents injected locally into the PPN to decrease its output, so that there is little doubt that these potentials are state-dependent, reticular in origin, and modulated in the same manner as the PPN, and therefore the equivalent of the human P50 potential. The P50 potential is, therefore, a measure of PPN output, an arousal-related waveform recorded at the level of the vertex in the human. Moreover, changes in its manifestation will inform about the function of the PPN in disease.

We hypothesize that the RP (in particular the early component), μ rhythm and P50 potential are all at least in part generated by PPN activity. This activity sets the background of activity for assessing sensory events, and precedes the execution of voluntary movements. The PPN, located at the transition between the midbrain and pons, is part of RAS and is in charge of the EEG high frequency states of waking and REM sleep (as opposed to low-frequency activity during SWS). The PPN, however, has become a target for deep brain stimulation (DBS) for the treatment of PD in human subjects. We suggested this possibility based on animal studies using stimulation in the region of the PPN, which at the time was thought to be part of the mesencephalic locomotor region (MLR) [24, 25, 31]. As we will see below, it took some 20 years for these findings to be translated to the bedside.

The mesencephalic locomotor region and the pedunculopontine nucleus

The MLR was originally described as a region that required very specific parameters in order to induce locomotion on a treadmill in the decerebrate cat [32]. That is, stimulation at low levels (typically <100 mA) needed to be localized to the lateral cuneiform nucleus (LCN), of increasing current amplitude, of long duration (0.5-1.0 ms) pulses, and delivered at 40-60 Hz. We became interested in the PPN since it is a descending target of basal ganglia structures [33, 34], and carried out experiments to determine if the PPN and the MLR represented overlapping structures. We were able to use electrical stimulation as well as injection of neuroactive agents into the region of the PPN to induce locomotion [35], and to record from locomotion-related PPN neurons, with some cells being active in relation to limb alternation, while others were related to the duration of the locomotor episode [36]. We then addressed why, in order to induce locomotion, LCN, but not medial (MCN), cuneiform nucleus stimulation was required, ramping up of the current was required, stimulation at 40-60 Hz, but not higher or lower was required, and why long duration pulses were required. On a practical basis, two other peculiarities applied to these parameters. When current was ramped up and locomotion was initiated, further increase in the current would lead to cessation of stepping, therefore, current had to be backed down in order to maintain locomotion. The other issue was that stepping was never immediate and always ensued but only after 1-2 s of stimulation. These requirements have finally been satisfactorily resolved.

We localized electrical stimulation sites within the region of reduced nicotinamide adenine dinucleotide phosphate diaphorase...
(NADPH-diaphorase) labeled cells (cholinergic neurons in the PPN are selectively labeled by this histochemical method) to induce locomotion on a treadmill in the precollicular-postmamillary transected cat. We also used neuroactive agents such as N-methyl-D-aspartate (NMDA) injected into the region of cholinergic cell labeling in transected cats. This ensured that activation of cell bodies by these agents, and not merely fibers of passage as was possible using electrical stimulation, was responsible for the stepping. Alternation between antagonists in the same limb, between agonists in opposite limbs, and a proximodistal delay of agonists in the same limb signaled that an appropriate locomotor step cycle was being induced following PPN stimulation chemically or electrically. Moreover, we found similar responses in the cat and the rat [24, 31-33, 35-43]. It is critical to realize that the reason such localization was possible was the use of very low current levels, below 100 μA, usually ~50 μA. By applying higher current levels locomotion can be induced following stimulation of more distant locations such as the MCN.

Since we were recording or injecting agents into a region that we had first physiologically identified to induce locomotion, we used the term “MLR” in most of the studies to describe our findings. However, we became convinced that the lowest threshold sites for inducing locomotion were located within the PPN [36]. We questioned the interpretation of the MLR as a locomotion-specific area, showing instead that the region stimulated was a “rhythmogenic” region that was part of the RAS [24-26, 36, 39-42]. Therefore, the term “MLR” needs to be retired.

The PPN has long been identified as a cholinergic cell group [44], but recent studies established that this nucleus includes non-overlapping populations of cholinergic, glutamatergic and GABAergic neurons [45]. Sagittal sections are optimal for visualizing the PPN, since it has a wedge shape extending from the dorsolateral to the ventrolateral mesopontine region (Fig. 2A). The pars compacta of the PPN is located in the LCN, posteriorly and dorsal to the lateral portion of the superior cerebellar peduncle (SCP), in the optimal site for inducing locomotion on a treadmill at low thresholds [37]. As the nucleus ranges anteroventrally, it is embedded in the superior cerebellar peduncle and ends in the posterior substantia nigra. The lowest threshold sites for inducing locomotion were located within the PPN pars compacta in the LCN [37, 39]. Stimulation at more ventral sites did not induce stepping, but rather changes in muscle tone [46], an effect that may also have been due to suddenly switching on the stimulus (see below). Stimulation of more medial regions such as the MCN, laterodorsal tegmental nucleus (LDT), the medial partner of the PPN, or anteroventrally in the region of the substantia nigra, did not induce locomotion [24, 26, 40-42]. This explains why stimulation of only LCN, but not MCN, in which the PPN is embedded, produces reliable stepping on a treadmill.

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**Figure 2.** The pedunculopontine nucleus (PPN). A. Sagittal plane. Histochemical NADPH-diaphorase (reduced nicotinamide adenine dinucleotide phosphate diaphorase) labeling of only the cholinergic neurons in the PPN revealed the wedge-shaped structure of this nucleus. Posteroventrally is located the pars compacta of the PPN, which is located within the lateral cuneiform nucleus (LCN) ventral to the inferior colliculus (IC). As the nucleus descends, its cells are intermixed with the fibers of the superior cerebellar peduncle (SCP). The body or pars dissipata of the PPN is located more ventrally and extends to the posterior edge of the substantia nigra (SN). Anterior is to the right. B. Semi-horizontal. This section is angled along the long axis of the PPN from the posteroventral to the medioventral direction (see Figure 2A for sagittal orientation). Histochemical NADPH-diaphorase labeling of cholinergic cells shows that PPN neurons of the pars compacta (pc) are located dorsally, are found within the cuneiform nucleus (CF), and laterally to the locus coeruleus (LC). Medial to the LC are cells of the laterodorsal tegmental nucleus (LDT) embedded within the central gray. Laterally, the pars dissipata (pd) of the PPN descends in an arc towards the posterior edge of the SN.
Figure 2B is a semi-horizontal section taken along the long axis of the PPN. The section is from the parabrachial area dorsally and posteriorly, angling downwards and anteriorly towards the posterior end of the substantia nigra. Basically, it is a view looking down from the top of the midbrain at the cholinergic neurons of the PPN and LDT. Medially and dorsally within the central gray is the aggregation of cholinergic cells of the LDT. More laterally to the LDT is the space occupied by the locus coeruleus (LC), whose cells are not labeled in this section. Dorsal to the LC is the CF, which contains the remainder of the pars compacta of the PPN in its lateral side. More ventrally and lateral to the LC is located the ventral edge of the pars compacta of the PPN, and the curvature of the PPN as it descends towards the posterior pole of the substantia nigra. Note the more compact nature of the cells in the pars compacta posteriorly (top of the view) compared to the pars dissipata anteriorly (bottom of the view). This section makes clear the reason why the optimal target for inducing locomotion at low current levels is so small, containing the aggregation of cells of the pars compacta of the PPN, which is embedded in the LCN, but not in the MCN. In keeping with these findings, a recent study on a rodent model of PD showed that stimulation of anterior PPN induced freezing and worsened gait, but gait was improved by posterior PPN stimulation [47].

The bottom line is that we believe the MLR is not a locomotion-specific region but rather an effect elicited by PPN stimulation. Further evidence that the PPN is responsible for these effects lies in the physiology of PPN neurons.

Pedunculopontine nucleus physiology

Why does stimulation of the PPN need to be applied at 40-60 Hz to induce stepping? We found that, when we applied depolarizing steps, all PPN cells increased firing frequency and then plateaued at 40-60 Hz as current levels were increased [48]. That is, these neurons could not be induced to fire any faster than 40-60 Hz, regardless of the level of current delivered intracellularly. This property was further investigated to determine the mechanism responsible for such a unique characteristic by patch clamp recording from neurons in the presence of synaptic blockers (to prevent afferent signals) and tetrodotoxin (to prevent action potential generation). In our investigation of the intrinsic properties of these cells, we applied current steps to attempt to drive high threshold channels, but the steps were unable to depolarize the membrane sufficiently to maintain depolarization to reveal membrane oscillations, probably due to the activation of potassium channels by the sudden depolarization [49]. However, when we applied ramps (instead of steps) to gradually depolarize the membrane to activate high threshold channels, we were able to elicit membrane oscillations in the β and γ band range. We found that all PPN neurons, regardless of transmitter or electrophysiological type, manifested intrinsic membrane oscillatory activity at β/γ frequencies (20-60 Hz) [49]. We used specific calcium channel blockers to identify both P/Q- and N-type high threshold, voltage-dependent channels as responsible for PPN neuron β/γ band oscillations. It should be noted that cells outside of the PPN do not show similar properties, making the cellular boundaries of the nucleus specific to this kind of activity. The presence of a frequency plateau in PPN cells explains the requirement to stimulate this region at 40-60 Hz to induce locomotion. Early locomotion studies revealed that suddenly switching on the current at previously determined threshold levels elicited only decrements in muscle tone instead of stepping [24, 46, 50]. Pulses delivered to the PPN had to be increased gradually until locomotion was induced after 1-2 s, in many cases then backing off the current, and only then was continuous stepping evident [24, 26, 50]. We referred to this characteristic as “recruiting” locomotion. Single cells in the PPN also need to be "recruited" to fire at γ band frequencies. The requirement to use ramps to elicit oscillations helps explain why current has to be gradually increased to depolarize PPN neurons slowly, i.e. to avoid activating potassium channels, and thus induce stepping. Sudden application of high amplitude currents would activate potassium channels and fail to sufficiently depolarize neurons. This would instead lead to inactivation of PPN neurons, explaining the decrement in muscle tone evident with sudden onset stimulation. This accounts for the need for ramping up current, instead of suddenly switching it on, in order to “recruit” stepping. In order to maintain such depolarization, the current needs to be reduced slightly, otherwise the depolarization will exceed the window for high threshold calcium channels. This explains the characteristic need to back off the current once locomotion is induced.

Why the need to use long duration pulses to elicit stepping? We hypothesize that the need to use long duration pulses is related to the high threshold calcium channels present in PPN neurons. We used calcium imaging to visualize ramp-activated calcium channels in PPN cells [51]. These voltage-dependent high threshold P/Q- and N-type calcium channels are located in the dendrites of PPN neurons. This explains the need to apply high levels of depolarization of the cell body in order to ultimately depolarize the higher resistance dendrites sufficiently to activate the high threshold calcium channels and promote gamma band oscillations. Long duration pulses, along with ramping up current levels, would be effective methods for activating dendritic calcium channels that would in turn allow PPN neurons to fire maximally at γ band frequencies.

Figure 3A shows the responses elicited in a representative PPN neuron to intracellular application of current steps. PPN cells fired maximally at 40-60 Hz when intracellular steps were applied [48]. We then identified the intrinsic membrane properties that caused PPN neurons to fire at these frequencies and found that they were due to the presence of high threshold voltage-dependent calcium channels [49]. Figure 3B, C shows that these channels required that ramps be used to depolarize the cell sufficiently to reach the high thresholds, since sudden current steps activated potassium channels that prevented the maintenance of such depolarization. These properties explain the need to stimulate the PPN at specific frequencies, ramping up current up to certain levels, and then backing off to maintain depolarization.
Oscillations were mediated by N- and P/Q-type voltage-dependent calcium channels [49]. For activation of high threshold calcium channels (around -20 mV in the soma). Further studies showed that these oscillations could be gradually increased to induce membrane oscillations that could be maintained within the window of high threshold, voltage-dependent calcium channels (-20 mV). C. Recordings in the same neuron but using square current steps (dark gray record represents the response to higher amplitude square current while light gray record represents the response to lower amplitude square current steps). Note that the membrane potential failed to be maintained and repolarized below the window for high threshold, voltage-dependent calcium channels (-20 mV). C. Recordings in the same neuron but using ramps of increasing amplitude (dark gray record represents the response to lower amplitude current ramp while the light gray record represents the response to higher amplitude current ramp). Note that the membrane potential could be gradually increased to induce membrane oscillations that could be maintained within the window for activation of high threshold calcium channels (around -20 mV in the soma). Further studies showed that these oscillations were mediated by N- and P/Q-type voltage-dependent calcium channels [49].

Years before the MLR was described, in the initial identification of the RAS, investigators stimulated the region of the PPN to transform the EEG from slow wave activity and sleep to fast activity and arousal [52]. They used chloralose-anesthetized or midbrain-transected (decerebrate) cats and stimulated the region of the mesencephalic reticular formation. They also performed lesions immediately anterior to the PPN that eliminated the effects of such stimulation. These studies typically used stimulation frequencies of 300 Hz, but established that 50 Hz stimulation was close to the lowest effective frequency. Interestingly, the effects of stimulation had a latency, typically of 1-2 s, that is, the effect of stimulation was not instantaneous, but ultimately effective in “recruiting” high frequency EEG activity. The frequency of stimulation and latency used in these RAS studies are remarkably similar to those used in the MLR studies.

But why does the RAS modulate arousal as well as postural and locomotor pathways? The RAS is a phylogenetically conserved system that modulates arousal and flight-or-flight responses. During waking, man’s ability to detect predator or prey is essential to survival. Under these circumstances, it is not surprising that the RAS can also modulate muscle tone and locomotion. This system is thus intrinsically linked to the control of the motor system in order to optimize attack or escape. During REM sleep, the RAS generates the atonia and keeps us from acting out our dreams. In fact, only our diaphragm and eye muscles appear to be acting out dream content. Therefore, during both waking and REM sleep, two states modulated by the PPN, the RAS can influence muscle tone and locomotion via the same reticulospinal systems, as well as arousal through ascending pathways to the intralaminar thalamus.

Outputs from the PPN activate reticulospinal systems that lead to profound hyperpolarization of motoneurons, which is the mechanism responsible for the atonia of REM sleep [53]. However, cholinergic projections from the PPN to the medullary reticular formation are known to induce decreased muscle tone at some sites, while producing stepping movements at other sites. This suggests the presence of a heterogeneous, distributed system of reticulospinal motor control. The required parameters of stimulation for eliciting these differing effects are important such that instantaneous, high frequency trains (similar to ponto-geniculo-occipital, PGO, burst neurons that may drive the atonia of REM sleep) trigger pathways that lead to decreased muscle tone, while lower frequency tonic stimulation leads gradually to the “recruitment” of locomotor movements [24, 26, 50]. Given the extensive evidence, it is to be expected that the PPN should modulate both posture and locomotion through descending projections, in addition to modulating arousal through ascending projections. Stimulation of the PPN would be expected to affect both postural and locomotor status, as well as arousal and wake-sleep functions.

**Parkinson’s disease**

PD patients, in addition to tremor, bradykinesia and akinesia, show sleep disturbances that include increased REM sleep drive, decreased SWS, frequent awakenings leading to daytime sleepiness, all resulting in insomnia [55]. These observations suggest that the RAS, especially the PPN that is in charge of waking and REM sleep, is overactive in PD. We carried out a study of the P50 potential in PD patients,
Preconscious awareness

The “preconscious” mind includes those things of which we are aware, but we are not paying attention (to them). If we choose to pay attention, we bring them to the “conscious” mind [72]. We proposed that it is the activation of the RAS during waking that induces coherent activity (through electrically coupled cells) and high frequency oscillations (through P/Q-type calcium channel and subthreshold oscillation activity), and lead to the maintenance (through activation of G-proteins) of the background of γ activity necessary to support a state capable of reliably assessing the world around us on a continuous basis [73, 74]. Our results suggest that a similar mechanism to that in the cortex for achieving temporal coherence at high frequencies is present in the PPN, and in its subcortical targets. We suggested that γ band activity and electrical coupling generated in the PPN may help stabilize coherence related to arousal, providing a stable activation state during waking that participates in preconscious awareness [73, 74].

Functionally, the process of preconscious awareness would need to be fairly continuous during waking in order to provide the sensory foundation for planned behavior. Importantly, this “stream of preconsciousness” would need to begin upon waking. It has been shown that increases in blood flow in the thalamus and brainstem begin within 5 min of waking, but as much as 15 min elapse before significant changes in frontal cortex blood flow are observed [75]. This is more surprising when considering that waking follows the last REM sleep episode of the night, during which frontal cortex blood flow is low [76]. The sudden onset of waking after the last REM sleep episode of the night does not instantly increase frontal lobe blood flow. Instead, upon waking, significant increases in the brainstem and thalamus precede restoration of blood flow to the frontal lobes.

The importance of subcortical structures in the determination of states of awareness is being growingly emphasized. Damasio proposed that the brainstem is critical to the formulation of the self, which is critical to the formulation of feelings [77]. Penfield extensively described the responses to cerebral cortex stimulation in a large number of epilepsy patients. Penfield arrived at the conclusion that “There is no place in the cerebral cortex where electrical stimulation will cause a patient to believe or to decide” [78]. In addition, he emphasized that, while cortical seizures localized to specific cortical regions elicit sensory or motor effects but maintain consciousness, petit mal seizures in “mesothalamic” (midbrain and thalamus) regions always eliminate consciousness.
Based on these results, Penfield proposed the presence of a "centrencephalic integrating system" that fulfills the role of sensorimotor integration necessary for consciousness.

**Conclusion**

The presence of γ band activity in the RAS, the most important discovery in the RAS in the last 10 years, participates in the process of preconscious awareness, and provides the essential stream of information for the formulation of many of our actions. The clinical implications of such a system are far-reaching and modulate such functions as fight-or-flight, arousal, wake-sleep control, posture and locomotion, and even volition and free will.

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