Melanin-concentrating hormone: a new sleep factor?

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INTRODUCTION

Sleep occupies approximately one-third of human’s life. Sleep is not a homogeneous process. In most mammals two sleep states are recognized: slow wave sleep (SWS) and the rapid eye movements (REM) sleep. Wakefulness, SWS, and REM sleep are differentiated by the electroencephalographic (EEG) activity (synchronized during SWS, desynchronized during wakefulness, and REM sleep), cognition (consciousness during wakefulness and dreams during REM sleep), muscle activity (atonia during REM sleep), and others physiological parameters (McCarley, 2007; Torterolo and Vanini, 2010).

In the twentieth century experimental and clinical research began to unravel the mystery of sleep. Nowadays, the main neuronal networks that are involved in the generation of wakefulness, SWS, and REM sleep are known (Siegel, 2005; McCarley, 2007; McCarley, 2007; Torterolo and Vanini, 2010). The thalamo-cortical network is the pacemaker for the EEG rhythms of wakefulness and sleep, and is the scaffold for the cognitive functions. This network is regulated by the activating (waking-related) systems located in the basal forebrain, postero-lateral hypothalamus, and mesopontine reticular formation. While the preoptic area is critical for SWS generation, the mesopontine region contains the “necessary and sufficient” neuronal networks for REM sleep generation.

Neurons containing the neuropeptide melanin-concentrating hormone (MCH) are mainly located in the lateral hypothalamus and the incerto-hypothalamic area, and have widespread projections throughout the brain. While the biological functions of this neuropeptide are exerted in humans through two metabotropic receptors, the MCHR1 and MCHR2, only the MCHR1 is present in rodents. Recently, it has been shown that the MCHergic system is involved in the control of sleep. We can summarize the experimental findings as follows: (1) The areas related to the control of sleep and wakefulness have a high density of MCHergic fibers and receptors. (2) MCHergic neurons are active during sleep, especially during rapid eye movement (REM) sleep. (3) MCH knockout mice have less REM sleep, notably under conditions of negative energy balance. Animals with genetically inactivated MCHR1 also exhibit altered vigilance state architecture and sleep homeostasis. (4) Systemically administered MCHR1 antagonists reduce sleep. (5) Intraventricular microinjection of MCH increases both slow wave sleep (SWS) and REM sleep; however, the increment in REM sleep is more pronounced. (6) Microinjection of MCH into the dorsal raphe nucleus increases REM sleep time. REM sleep is inhibited by immunoneutralization of MCH within this nucleus. (7) Microinjection of MCH in the nucleus pontis oralis of the cat enhances REM sleep time and reduces REM sleep latency. All these data strongly suggest that MCH has a potent role in the promotion of sleep. Although both SWS and REM sleep are facilitated by MCH, REM sleep seems to be more sensitive to MCH modulation.

Several neurotransmitters and neuromodulators such as acetylcholine (ACh), noradrenaline (NA), dopamine, serotonin, histamine, adenosine, and hypocretins, have been involved in the control of sleep and wakefulness (Jones, 2005; McCarley, 2007; Torterolo and Vanini, 2010). Do melanin-concentrating hormone (MCH), and MCHergic system play a role in the control of sleep? Is MCH a sleep factor? So far, about 1000 papers related to MCH have been published. Notwithstanding this, only a relatively small number of publications associate MCH with sleep. In this review article we evaluate the data on MCH and sleep, and give our point of view of how the MCHergic system might regulate behavioral states.

MELANIN-CONCENTRATING HORMONE

In mammals, MCH is a cyclic neuropeptide with 19 aminoacids that has been found predominantly in neurons localized in the lateral hypothalamus and incerto-hypothalamic area (Kawauchi et al., 1983; Skofitsch et al., 1985; Bittencourt et al., 1992).

Originally, MCH was described as a circulating hormone isolated from salmon pituitaries, where it induces the aggregation of melanin granules in melanophores; this action results in a pale skin color (Kawauchi et al., 1983). This mechanism lets the animal to camouflage in response to the background color of the environment or in response to stress. This effect is opposite to
the pigment-dispersing effect of alpha-melanocyte stimulating hormone (α-MSH). Subsequently, MCH was identified in rat brain using immunohistochemical techniques (Skofitsch et al., 1985; Bittencourt et al., 1992).

MCH structure is identical in all mammals analyzed to date, including mice, rats, rabbits, and humans (Saito and Nagasaki, 2008). MCH is generated by the cleavage of a precursor of 165 amino acids, the prepro-MCH (ppMCH). ppMCH contains other peptides in addition to MCH, designated as neuropeptide EI (NEI) and neuropeptide GE (NGE; Bittencourt and Celis, 2008). In addition, the precursor may potentially give rise to an alternative splice variant termed MCH-gene-overprinted-polypeptide (MGOP), as well as a portion of the antisense-RNA-overlapping-MCH (AROM; Toumaniantz et al., 1996; Borsu et al., 2000).

MCH RECEPTORS AND SIGNALING MECHANISMS

Several research groups almost simultaneously discovered the first MCH receptor in 1999. It was found to be an orphan G-protein coupled receptor, named SLC-1 (Bachner et al., 1999; Chambers et al., 1999; Lembo et al., 1999; Shimomura et al., 1999). Because MCH was the only ligand from rat brain extract that activated rat or human SLC-1 receptor, it was named MCHR1. In 2001, another MCH receptor was identified. This receptor was termed MCHR2 and has a 38% sequence homology with MCHR1 (Hill et al., 2001; Mori et al., 2001; Sailer et al., 2001).

The binding of MCH to MCHR1 activates diverse intracellular signaling pathways by coupling to Gi, Gq, and Go proteins, while MCHR2 is known to couple to the Gq protein (Hawes et al., 2000; Sailer et al., 2001).

ANATOMY OF THE MCHergic SYSTEM

LOCATION OF THE MCHergic NEURONS

In their classical paper Bittencourt et al. (1992) described that the ppMCH mRNA as well as MCH were localized in neurons of the lateral hypothalamic area, and the incerto-hypothalamic area of the rat. MCHergic neurons are distributed continuously in a rostrocaudal extent approximately from the level of the caudal aspect of the paraventricular nucleus to just beyond the mammillary recess in rat, cats, and humans (Bittencourt et al., 1992; Mouri et al., 1993; Torterolo et al., 2006). Photomicrographs of the MCHergic neurons in the tuberal hypothalamic region of the cat are shown in Figure 1.

A small number of MCH-containing neurons was also observed in the olfactory tubercle and pontine reticular formation in the rat, as well as in the lateraldorsal tegmental (LDT) nucleus and preoptic area of female and lactating rats, respectively (Bittencourt et al., 1992; Rondini et al., 2007, 2010).

PROJECTIONS OF THE MCHergic NEURONS

MCHergic neurons project widely throughout the central nervous system (CNS; Bittencourt et al., 1992). Because of their importance in the control of the sleep–wake cycle, we highlight the projections toward the thalamus, the tubero-mammillary nucleus of the hypothalamus, the preoptic area of the hypothalamus, the ventral tegmental area, the periaqueductal gray, the locus coeruleus (LC), the nucleus pontis oralis (NPO), the laterodorsal and pedunculopontine tegmental (LDT–PPT) nuclei, and the dorsal raphe (DR) nucleus. Furthermore, projections to other structures such as the cerebral cortex, amygdala, and motor nuclei have been described (Bittencourt et al., 1992; McGregor et al., 2005; Elias et al., 2008; Torterolo et al., 2009).

DISTRIBUTION OF THE MCH RECEPTORS

MCH receptors are distributed in diverse areas of the CNS. The MCHR1 is present in all mammals studied to date, while the MCHR2 is functional in dogs, ferret, rhesus monkey, and humans but is a pseudogene in rodents (Tan et al., 2002).

There is a widespread distribution of the MCHR1 mRNA, and like the MCHergic fibers, this receptor is present in limbic
structures and in areas related to the control of sleep and wakefulness. MCH1R have been detected also in several peripheral organs (Chung et al., 2009).

The distribution of MCHR2 nearly overlaps with that of MCH1R, but the latter shows higher relative levels of density and a wider distribution pattern (Hill et al., 2001; Mori et al., 2001). The function of MCHR2 remains unknown due to the lack of available animal models.

**MCH: VOLUME CONDUCTION THROUGH THE CEREBROSPINAL FLUID (CSF)?**

As it is the case with the central actions of oxytocin (Veening et al. 2007, 2010). the hypothalamus are also GABAergic. Besides, MCH and GABA

Sapin et al. (2010) demonstrated that 85% of the MCHergic neurons of the tuberal region of the hypothalamus of the rat. MCHergic neurons that co-localize with MCH. Elias et al. (2001) have demonstrated that the anorexigenic cocaine- and amphetamine-regulated transcript (CART), is co-expressed in 95% of the MCHergic neurons of the zona incerta and in 70% of MCH-positive neurons of the lateral hypothalamus of the rat. MCHergic neurons that co-localize with CART send ascending projections toward the septum and hippocampus, whereas the non-CART MCHergic neurons send descending projections toward the brainstem and spinal cord (Hanriot et al., 2007). In addition, all the MCHergic neurons express the novel hypothalamic molecule nesfatin-1 that also suppresses feeding (Fort et al., 2008).

Recently, utilizing MCH immunohistochemistry and in situ hybridization to detect GAD67 mRNA, Sapin et al. (2010) demonstrated that 85% of the MCHergic neurons of the tuberal region of the hypothalamus are also GABAergic. Besides, MCH and GABA co-localize in the LDT and medial preoptic area (Rondini et al., 2007, 2010).

**ELECTROPHYSIOLOGICAL CHARACTERISTIC OF THE MCHergic NEURONS**

*In vitro* recordings have shown that the MCHergic neurons present a resting potential ranging from −62 to −50 mV, slightly more hyperpolarized than nearby hypocretinergic neurons (Guyon et al., 2009). MCHergic neurons also showed a relatively low level of spontaneous activity compared with hypocretinergic and GABAergic neurons of the region.

Glutamate and GABA regulate MCHergic neuronal activity. Accordingly, both AMPA and NMDA depolarize these neurons (van den Pol et al., 2004). The activation of group I of metabotropic glutamate receptors also excite MCHergic neurons (Huang and van den Pol, 2007). The GABA agonist muscimol hyperpolarizes MCHergic neurons while there are no data regarding the effect of GABA
g agents Guyon et al. (2009).

**SYNAPTIC EFFECTS OF MCH**

Studies in lateral hypothalamic neurons *in vitro* have shown a predominantly inhibitory effect of MCH both at pre- and postsynaptic levels (Gao and van den Pol, 2001). Hence, MCH inhibited the synaptic activity of both glutamatergic and GABAergic inputs to lateral hypothalamic neurons. In addition, MCH reduced the amplitude of glutamate-evoked currents and the amplitude of miniature excitatory currents, indicating an inhibitory modulation of postsynaptic glutamate receptors. Furthermore, MCH attenuated L-, N-, and P/Q-type calcium voltage-activated currents (Gao and van den Pol, 2002).

MCHergic neurons express MCHR1 autoreceptors; by this mean the neuropeptide inhibits MCHergic neurons via a depres-

sion of voltage-dependent calcium channels without affecting the resting potential (Huang et al., 2007). Therefore, MCHergic neurons would exert a negative feedback on their own activity.

**ROLE OF MCH IN THE CONTROL OF ENERGY HOMEOSTASIS AND MOOD**

The high conservation of MCH structure along phylogeny sug-

MCHergic neurons are mainly located, is an integrative region par excellence. This region has been implicated in a wide variety of functions, such as energy homeostasis, wakefulness and sleep, regulation of motor activity, motivation, and reward (Bernardis and Bellinger, 1996). MCHergic neurons seem to be implicated in some of these functions.

Due to the strong relationship among sleep, energy homeosta-

There are strong experimental evidences that MCH increases food intake and reduces energy expenditure such as:

1. Acute infusion of MCH into the lateral ventricles induces feeding in rodents. Fasting induces an up-regulation of MCH mRNA expression (Qu et al., 1996).
2. MCH mRNA levels are also elevated in the obese leptin-deficient ob/ob mice, suggesting that leptin negatively regulates MCH expression (Qu et al., 1996). In fact, by means of double knockout mice for MCH and leptin, it was demonstrated that
MCH is a critical mediator of the leptin-deficient phenotype (Segal-Lieberman et al., 2003).

3. The stimulating effect of food intake has been confirmed in transgenic mice, in which an over-expression of MCH causes obesity (Ludwig et al., 2001), while prepro-MCH knockout mice (MCH-KO) are hypophagic and lean (Shimada et al., 1998). Indeed, MCH is the only known hypothalamic peptide whose genetic deletion results in leanness. In these MCH-KO mice, an increase in temperature, heart rate, and metabolic rate occurs along with an increase in energy expenditure (Astrand et al., 2004). A toxin-mediated genetic cell ablation strategy using a truncated ataxin-3 has been applied to induce apoptosis of MCHergic neurons; these mice developed a late onset syndrome characterized also by leanness, hypophagia, and increased energy expenditure (Alon and Friedman, 2006).

4. Mice knockout for the MCHR1 are lean, although in contrast to MCH-KO mice, they are hyperactive, hyperphagic, and have altered metabolism (Marsh et al., 2002). In addition, blockade of MCHR1 with specific antagonists has anorectic effects (Borowsky et al., 2002).

5. The MCHergic neurons are under the direct control of NPY-containing neurons of the arcuate nucleus; these neurons are part of the neuronal network that governs feeding and energy expenditure (van den Pol et al., 2004). Furthermore, peripheral signals of energy abundance such as glucose, enhance MCHergic activity in vitro (Burdakov et al., 2005).

The presence of MCHergic fibers and receptors in the limbic system tends to suggest that the MCHergic system plays a role in the control of mood and emotion. Borowsky et al. (2002) have shown that systemic administration of an MCHR1 antagonist produces antidepressant effects (Borowsky et al., 2002). We have also shown that the microinjection of a low dose of MCH (50 ng) into the DR of the rat evokes a depressive-like behavior as shown by a significant increase in the immobility time as well as a decrease in climbing behavior in the forced swimming test (Lagos et al., 2011b). The depressive-like effect induced by MCH was prevented by systemic pre-treatment with fluoxetine, a selective serotonin reuptake inhibitor. Consistent with these results, the immunoneutralization of MCH by microinjection of antibodies anti-MCH into the DR has antidepressant effects; anti-MCH antibodies decreased the immobility time in the forced swimming test without modifying the motor activity measured in the open field test.

The above-mentioned data suggest that MCH modulates the activity of serotonergic DR neurons, and could be involved in the control of mood. Furthermore, antagonism of the MCHR1 produces also a decrease in anxiety in animals models (Borowsky et al., 2002; Chaki et al., 2005).

Additional functions of MCH via central or peripheral MCHR1 such as the control of reproduction or the immune response have been also described (Lakaye et al., 2009; Wu et al., 2009).

ROLE OF MCH IN THE CONTROL OF SLEEP AND WAKEFULNESS

From 2003 on, several investigators have examined the role of MCH in sleep, and the evidence indicates that it promotes sleep (Torterolo et al., 2003a; Verret et al., 2003; Willie et al., 2003).

ANATOMICAL CONSIDERATIONS

We mentioned above that MCHergic fibers and receptors are present in regions related to the control of sleep and wakefulness. In addition, MCHergic neurons have been identified in key areas responsible for the generation of wakefulness and REM sleep such as the LDT (only in female rats), and the pontine reticular formation (Bittencourt et al., 1992; Rondini et al., 2007). Last but not least, MCH is present in the tanyocytes of the DR of the cat, where serotoninergic REM “OFF” neurons are present (Torterolo et al., 2008).

RELATIONSHIP WITH HYPOCRETINERGIC NEURONS

Due to the importance of the hypocretinnergic system in sleep physiology and pathology, the interaction between the MCHergic, and the hypocretinergic system is worth to be considered. The hypocretinergic neurons degenerate in narcolepsy, a paradigmatic sleep disorder (Nishino et al., 2000; Peyron et al., 2000; Thannickal et al., 2000). The activity of the hypocretinergic neurons increases during active wakefulness, and they are believed to be central in the maintenance of this state (Torterolo et al., 2001, 2003b; Lee et al., 2005; Milejkovskiy et al., 2005).

A strong relationship exists between hypocretinergic and MCHergic neurons in the hypothalamus. The MCHergic neurons are intermingled with hypocretin (also called orexin)-containing neurons in the lateral hypothalamus, mainly at the tuberal and tuberohypothalamic levels (Torterolo et al., 2006). MCHergic fibers are in close relationship with hypocretinergic neurons and vice versa, which tends to indicate the existence of reciprocal synaptic contacts between both types of cells (Guan et al., 2002; Torterolo et al., 2006). This fact as well as the presence of hypocretinergic receptors on MCHergic neurons suggests an important functional interaction between both systems (Backberg et al., 2002). In fact, hypocretin increases MCH mRNA expression in hypothalamic neurons, directly excites MCHergic neurons and increases glutamate release onto them (Bayer et al., 2002; van den Pol et al., 2004). On the other hand, MCH modulates hypocretin/orexin-mediated effects on behavioral state and synaptic transmission in the lateral hypothalamus (Rao et al., 2008). Rao et al. (2008) demonstrated that the efficacy of glutamatergic synapses on hypocretinergic neurons is enhanced in MCHR1 knockout mice, and hypocretin-1-induced firing is facilitated. On the contrary, in wild-type mice, MCH significantly attenuated the hypocretin-1-induced enhancement of spike frequency in hypocretinergic neurons, but not its basal activity. Furthermore, in these neurons, MCH attenuated hypocretin-1-induced enhancement of the frequency of miniature excitatory postsynaptic currents (EPSCs). These effects imply that MCH exerts a unique inhibitory influence on hypocretinergic signaling as a way to fine-tune the output of these neurons.

Interestingly, hypocretinergic and MCHergic neurons respond in a different way to most homeostatic signals such as glucose (Burdakov et al., 2005), or to waking-related neurotransmitters such as NA (Bayer et al., 2005). In this regard, it is of note that while hypocretinergic neurons express the α1 adrenergic receptors,
MCHergic neurons express the α2 adrenergic receptors, which are related to activation or inhibition of its target, respectively (Modirrousta et al., 2005).

**WAKING AND SLEEP-RELATED MECHANISMS NEUROTRANSMITTERS REGULATE MCHergic ACTIVITY**

Serotonergic neurons of the dorsal and median raphe nucleus, as well as noradrenergic neurons of the LC are active during wakefulness (Monti and Jantos, 2008; Monti, 2010b; Torterolo and Vanini, 2010). Midbrain dopaminergic neurons are also involved in the promotion of wakefulness (Monti and Jantos, 2008). On the other hand, cholinergic neurons are active both during wakefulness and REM sleep (Torterolo and Vanini, 2010). All these neurotransmitters regulate the activity of the MCHergic neurons.

Noradrenaline hyperpolarizes the MCHergic neurons by direct synaptic actions mediated by the α2 adrenergic receptor (van den Pol et al., 2004). Furthermore NA has presynaptic effects; it decreases the frequency of EPSCs and increases the frequency of miniature inhibitory postsynaptic currents (IPSCs; van den Pol et al., 2004).

Serotonin hyperpolarizes MCHergic neurons in the presence of TTX indicating a direct postsynaptic effect (van den Pol et al., 2004).

Dopamine decreases the excitability of MCHergic neurons by decreasing the membrane resistance without modifying the resting potential (Guyon et al., 2009).

Acetylcholine hyperpolarizes MCHergic neurons through muscarinic receptors and increases GABA release onto these neurons through nicotinic mechanisms (van den Pol et al., 2004).

Cannabinoids are sleep-promoting neuromodulators (Murillo-Rodriguez, 2008), and they increase the firing rate of the MCHergic neurons. Acting through the CB1 receptor, these substances induce depolarization of MCHergic neurons and decrease the activity of nearby GABAergic neurons (Huang et al., 2007).

In summary, extrahypothalamic waking-related neurotransmitters (NA, serotonin, ACh) inhibit MCHergic neurons, thus suggesting that in order to generate or maintain wakefulness, these neurons should be inhibited. Notwithstanding this, it is intriguing that ACh promotes REM sleep and simultaneously inhibits MCHergic neurons (see below).

**MCHergic NEURONAL ACTIVITY DURING WAKEFULNESS AND SLEEP**

Harthoorn et al. (2005) have demonstrated an up-regulation of the content of MCH in hypothalamic neurons during the resting phase of the light–dark cycle of the rat, when this animal sleeps more.

Modirrousta et al. (2005), studied the c-fos expression (as an index of neuronal activity) of the MCHergic neurons after a sleep rebound that followed a short protocol (3 h) of total sleep deprivation in the rat. Although c-fos expression was observed in a very small proportion of MCHergic neurons (≈3%), the authors demonstrated that MCHergic neurons were activated during the sleep rebound (that consisted mainly in SWS) in comparison to wakefulness. In contrast, almost no MCHergic neurons expressed c-fos during sleep (mainly SWS) induced by an adenosine receptor agonist in rats (Satoh et al., 2006). In the cat, we did not observe Fos immunoreactivity in MCHergic neurons either during a SWS rebound that followed a short protocol of total sleep deprivation, or during REM sleep induced by microinjections of carbachol into the NPO (Torterolo et al., 2006). However, it should be considered that the occurrence of a false negative is one of the inconveniences of this technique.

Verret et al. (2003) also studied the c-fos expression in the MCHergic neurons of the rat during a REM sleep rebound that followed a prolonged protocol (72 h) of REM sleep deprivation. The authors observed that in comparison to the waking period, during the REM sleep rebound a large number of Fos+ cells were immunoreactive for MCH and almost 60% of the MCH-containing neurons were Fos positive. The MCHergic neurons that expressed c-fos either did or did not co-localize CART (Hanriot et al., 2007).

In accordance with the studies mentioned above, MCHergic neurons did not express c-fos in wakefulness induced by application of GABA_A antagonists in the lateral hypothalamus (Goutagny et al., 2005).

Hassani et al. (2009) made use of in vivo extracellular recordings of identified MCHergic neurons of head restrained rats to describe their activity during the sleep–wakefulness cycle. These neurons were silent during wakefulness and began to discharge during the SWS transition; their firing rate slightly increased during SWS and reached a maximum during REM sleep. During this period of time they fired in doublets or groups of spikes, and often in relation to twitches of the whiskers and muzzle. Furthermore, the MCHergic neurons discharged in a reciprocal manner to the hypocretinergic neurons. Although the sleep “ON” pattern of discharge of the neurons that were analyzed was clear, it remains to know if this is the general pattern of activity of all the hypothalamic and extrahypothalamic MCHergic neurons. Besides, due to the strong relationship of the MCHergic neurons with the control of energy homeostasis, it would be important to know if in conditions of positive energy balance there are MCHergic neurons active during wakefulness, as suggested by the in vitro studies of Burdakov et al. (2005).

**STUDIES ON GENETICALLY MODIFIED MICE**

Studies in knockout mice for both MCHR1 and MCH, show that these animals have an altered sleep architecture (Adamantidis et al., 2008; Willie et al., 2008).

MCH-KO mice have a reduction in SWS and an increase of wakefulness during baseline conditions (Willie et al., 2008). In response to fasting these animals exhibit a marked hyperactivity, accelerated weight loss, and an exaggerated decrease in REM sleep compared to the wild-type littermates. The decrease in REM sleep results from a reduced number of REM sleep bouts. Following 6 h of sleep deprivation, these animals have a normal sleep rebound. This data support a role for MCH in vigilance state regulation, particularly in response to a challenge in energy homeostasis.

Adamantidis et al. (2008) studied the spontaneous sleep and its homeostatic regulation in mice with deletion of the MCHR1 gene. In contrast to previous data showing that MCH promotes sleep, the authors described that the lack of functional MCHR1 results in a hypersomnia-like phenotype. In comparison with wild littermates, these mice exhibit higher amounts of REM sleep during the active (dark) phase. Furthermore MCHR-1 knockout mice...
showed a more pronounced rebound of SWS in response to sleep deprivation. In addition, the authors found that modafinil, a stimulating drug, induced less wakefulness in MCHR1 knockout mice than in wild-type littermates. Then, these MCHR1 knockout animals exhibit altered vigilance and sleep homeostasis, but the sign of the alteration seems the opposite to previous results. The authors discussed that these unexpected results could be related to the development of compensatory mechanisms; the resulting adaptive changes would oppose those caused by the blockade of the endogenous peptide.

**EFFECT OF MCH ADMINISTRATION ON SLEEP VARIABLES**

**Intraventricular microinjections of MCH**

Intraventricular microinjections of MCH into the active (dark) phase of the rat, produced a marked increase in REM sleep (up to 200% above control values) and a moderate increase in SWS (up to 70% of control values; Verret et al., 2003). The increment in REM sleep time was due to an increase in the frequency of REM sleep episodes, whereas their duration remained unchanged.

**Intracerebral microinjections of MCH**

Using cholera toxin subunit-b as a retrograde tracer in cats, we have demonstrated that ≈15% of the lateral hypothalamic neurons that project to the NPO, contained MCH (Torterolo et al., 2009). The NPO of the cat (also called medial pontine reticular formation or perilocus coeruleus α) or its corresponding nucleus in the rat, which is called the sub-laterodorsal nucleus, is an integral part of different models of REM sleep, and is considered to exert executive control over the initiation and maintenance of this behavioral state (Reinoso-Suarez et al., 2001; Xi et al., 2001; Chase and Morales, 2005; Siegel, 2005; Fuller et al., 2007; Luppi et al., 2007; McCarley, 2007). A single injection of a cholinergic agonist, such as carbachol, generates REM sleep with a very short latency (down to 30 s) and a duration that can exceed 2 h (Baghdoyan et al., 1987). Microinjections of MCH within this region produced a significant decrease in REM sleep latency with an increase in the time the animal spent in this behavioral state. The increment in REM sleep time was due to a increase of REM sleep episodes duration and frequency (Torterolo et al., 2009).

As mentioned earlier, the DR has a dense network of MCH-immunoreactive fibers and tanyctyes (Torterolo et al., 2008). In this respect, we have studied the effect of MCH administration into the DR on sleep during the light period of the light–dark cycle of the rat. MCH induced a marked and dose-dependent increase of the time the animal spent in REM sleep, and also a moderate increase of SWS; on the other hand, wakefulness was reduced (Lagos et al., 2009). The increase of REM sleep time was due to an increment in the frequency of REM sleep bouts, while the duration of the episodes was not modified. **Figure 2** shows representative hypnograms of an animal microinjected with MCH or vehicle (control) into the DR.

The blockade of physiologically released-MCH in the DR by immunoneutralization (microinjections of antibodies anti-MCH) increased wakefulness and suppressed REM sleep, while SWS tended to remain unchanged (Lagos et al., 2011a). The decrement in REM sleep time was due to a decrease in the frequency of REM sleep episodes whereas the duration of the episodes was not modified (Figure 3).

**Systemic administration of MCHR1 antagonists**

Systemic administration of MCHR1 antagonists in rats decreased SWS and REM sleep time. This treatment also increased
wakfulness as well as SWS and REM sleep latencies (Ahnaou et al., 2008). The reduction of REM sleep was related to a smaller number of REM sleep bouts, while the duration of the REM sleep episodes was unchanged.

MCH: PROMOTION OF SLEEP AND SUPPRESSION OF WAKEFULNESS

Most of the preceding data strongly suggest that the MCHergic system promotes sleep, but especially REM sleep. Indeed, reversible inactivation by microinjection of a GABA<sub>A</sub> agonist into the hypothalamic area where the MCHergic neurons are located, strongly suppresses REM sleep (Lin et al., 1989).

**Hypocretinergic neurons**

MCH inhibits waking-related hypocretinergic neurons (Rao et al., 2008), and by this means the MCHergic system may facilitate SWS and REM sleep. In fact, sleep attacks, sleep onset REM, and the intrusion of REM sleep-like atonia during wakefulness (cataplexy) characterize narcolepsy, where there is a suppression of the hypocretinergic activity due to a degeneration of the hypocretinergic neurons (Guilleminault and Fromherz, 2005; Mignot, 2005).

**Serotonergic neurons**

Serotonergic neurons of the DR are active during wakefulness, reduce their discharge during SWS, and are essentially silent during REM sleep (REM-off neurons; Jacobs and Azmitia, 1992). Furthermore, serotonin release decreases during REM sleep in diverse brain areas (Portas et al., 2000). Application of 5-HT<sub>1A</sub> receptor agonists into the DR facilitates REM somn occurrence, an effect mediated by the activation of inhibitory somatodendritic 5-HT<sub>1A</sub> autoreceptors (Portas et al., 1996; Monti et al., 2002). The inhibition of serotonergic neurons of the DR following the microinjection of the GABA<sub>A</sub> agonist muscimol also increases REM sleep time (Nitz and Siegel, 1997). This outcome is similar to the one obtained following intra-DR microinjections of MCH (Lagos et al., 2009). In preliminary studies, microinjection of MCH into the DR of the cat produced similar effects (Devera et al., 2007).

Based upon the wake-related discharge pattern of serotonergic neurons it has been suggested that these neurons promote also wakefulness; in fact, there is strong evidence that the serotonergic neurons of the DR have a key role in the activation of the EEG (Dringenberg and Vanderwolf, 1998).

Therefore, it is likely that the microinjection of MCH into the DR inhibits the serotonergic neurons and by doing so, reduces wakefulness and increases REM sleep.

**GABAergic neurons of the NPO**

Due to the postulated inhibitory actions of MCH, we believe that in order to facilitate the generation of REM sleep, MCH must inhibit the GABAergic “wake-ON” neurons in the NPO, thus allowing the activation of “REM sleep-ON” neurons (Xi et al., 1999; Luppi et al., 2007; Brown et al., 2008).

**Other waking-related neurons**

MCHergic neurons project also to other neural structures involved in the control of sleep and wakefulness including the basal forebrain, the tubero-mammillar nucleus of the hypothalamus, the LDT–PPT, the LC, and the ventrolateral periaqueductal gray (vPAG). Interestingly, the bilateral inhibition or lesions of the vPAG and nearby areas generates an important increase in REM sleep amount (Sastre et al., 1996; Vanini et al., 2007; Kaur et al., 2009); therefore it is expected that MCH would promote REM sleep when applied onto this region (Peyron et al., 2009). We believe that MCH promotes sleep (both SWS and REM) acting not only on the vPAG, but acting in tandem on several waking and sleep-related regions.

MCHergic neurons innervate also the specific and non-specific nuclei of the thalamus, including the reticular nucleus (Bittencourt et al., 1992). By actions onto these nuclei as well as onto the neocortex, the MCHergic neurons could influence the EEG rhythms and behavioral states (Steriade, 2005).

**PHYSIOLOGICAL CONSIDERATIONS**

Energy conservation and restoration is one of the probable functions of sleep (Benington and Heller, 1995). Thus MCH would have an integrative role in energy conservation by promoting food intake, decreasing metabolism, and inducing sleep.

MCH decreases temperature, heart rate, and metabolic rate by central actions that enhance the parasympathetic/sympathetic tone ratio and decrease the release of thyroid hormones. Somatomotor activity is also reduced by MCH (Saito and Nagasaki, 2008). Given the profile of the activity of the MCHergic neurons it is possible that these actions are maximally expressed during sleep.

In order to conserve energy, MCHergic neurons could be involved also in the promotion of sleep in accordance with metabolic demands. It is likely that in conditions of energy abundance such as post-feeding time, MCH would first decrease the level of wakefulness, and then induce sleep. Furthermore, in special cases when energy conservation is imperative, MCHergic system may play a critical role. One case is hibernation, where the whole body metabolism falls to 1–2% of the basal value, and the body temperature could reach the freezing point of water (Heller and Ruby, 2004). Interestingly, the entrance to this state is through SWS.

Another paradigmatic physiological state is lactation, where we hypothesize that the activity of the MCHergic system would be very high. In fact, during this condition MCH is expressed in neurons of the medial preoptic area, a critical region for reproductive control and maternal behavior (Rondini et al., 2010). During the lactating period in the rat, the animals appear somnolent and the EEG is synchronized before milk ejection (Lincoln et al., 1980). Lactation is also associated with an increase in SWS in women (Blyton et al., 2002). Furthermore, it has been reported an increase in the amount of REM sleep after delivery (Petre-Quadens and De Lee, 1974). These authors also showed a gradual decrease in REM sleep in bottle-feeding mothers compared to breastfeeding mothers where REM sleep values remain high. Therefore, it would be important to investigate if the cause of this hypersomnia during lactation is promoted by MCH.

On the other hand, depression of the MCHergic neuronal activity may mediate the effect of long-term sleep deprivation. This condition leads to a marked increase in energy expenditure and body weight loss (Rechtschaffen and Bergmann, 2002). In fact, this syndrome partly recalls the effect of the deletion of the MCH gene.
MCH in Sleep: Working Hypothesis

Recently, Peyron et al. (2009) suggested that the MCHergic neurons are inhibited during wakefulness by extrahypothalamic waking-related neurons, and during SWS are inhibited by GABAergic neurons of the sleep-promoting preoptic region. On the other hand, glutamatergic REM sleep “ON” neurons of the sublaterodorsal nucleus would drive the MCHergic neurons. Thus, according to this proposal the MCHergic neurons are inhibited or driven by the classical activating or sleep-promoting system.

A complementary view is to consider that MCHergic neurons play a more active role in the induction of sleep; in other words, these neurons would drive the sleep-promoting regions instead of being driven by them. In addition, MCHergic neurons would inhibit the activating systems and promote sleep. A simple model is presented in Figure 4.

The MCHergic System in Pathology

In spite of the large number of preclinical studies, there are almost no data regarding the pathophysiology of the MCHergic system.

Preclinical studies have demonstrated that MCH is related to energy homeostasis and it has been suggested that a dysfunction of this system could lead to obesity. In fact, there is one study that reports a trend toward association of several MCHR1 single-nucleotide polymorphisms with an obese phenotype in independent study groups of obese German children and adolescents (Wermter et al., 2005). Furthermore, MCHR1 antagonists could be a therapeutic option for treating obesity (Rivera et al., 2008).

Kokkotou et al. (2008) demonstrated that mice genetically deficient in MCH have substantially reduced local inflammatory responses in a model of experimental colitis. Likewise, mice receiving treatments with an anti-MCH antibody developed attenuated colonic inflammation and survived longer. Furthermore, there is an increased colonic expression of MCH and its receptor in patients with inflammatory bowel disease, suggesting a role of MCH in inflammatory processes in the intestine (Kokkotou et al., 2008).

We have mentioned earlier that there is preclinical evidence to support an increase in MCHergic activity in depression and that the antagonism of this system has antidepressant effects (Borowsky et al., 2002; Lagos et al., 2011). In addition to pro-depressive effects, MCH promotes the occurrence of REM sleep in laboratory animals. Interestingly, in patients with a diagnosis of major depression (where there would be an increase in MCHergic tone), REM sleep occurrence is increased (Adrien, 2002). Accordingly, it could be of interest to determine the role of MCH in patients with depressive disorder.

Postpartum emotional distress or “blues” occurs in about 75–80% of new mothers about 3–5 days after birth (Lee, 1998). This “blues” is typically confined to the 1st week postpartum. More serious forms of maternal depression and postpartum psychosis occur in less than 10% of new mothers. Regarding the probable increment in activity of the MCHergic neurons during postpartum and lactation, it would be important to investigate if MCH is related to these emotional distresses.

As above-mentioned, there is a close anatomical and functional relationship between hypocretinergic and MCHergic neurons. Therefore, it is imperative to know how MCHergic neurons function in narcolepsy, where there is a degeneration of the hypocretinergic neurons; in fact, the malfunction of MCHergic neurons could play a role in the pathophysiology of this disease.

Parkinson’s disease (PD) but not Huntington’s disease, is accompanied by decrease in the number of MCHergic neurons (Thanickal et al., 2007; Aziz et al., 2008). The tentative relationship between a loss of MCHergic activity and some symptoms of PD has not been studied yet; however, a decrease in the activity of MCHergic cells could partly explain the high incidence of the REM sleep behavior disorder in this pathology (Thorpy and Adler, 2005).

Conclusions and Future Directions

MCHergic neurons constitute a powerful regulatory system with wide and divergent projection. MCH is involved in the central regulation of food intake, energy balance, and mood. Recent evidence strongly suggests that MCH has a regulatory role in the control of sleep, mainly REM sleep. New experimental approaches are needed to unravel the role of MCH in sleep. Clinical research is also needed in order to determine the involvement of this peptide in a number of neuropsychiatric conditions.

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