Yawning in neurology: a review
Bocejo em neurologia: uma revisão
Hélio A. G. Teive¹, Renato P. Munhoz², Carlos Henrique F. Camargo³, Olivier Walusinski⁴

ABSTRACT
Yawning is a stereotyped physiological behavior that can represent a sign or symptom of several conditions, such as stroke, parakinesia brachialis oscitans, parkinsonism, Parkinson’s disease and epilepsy. More rarely, it can occur in patients with intracranial hypertension, brain tumor, multiple sclerosis, migraine, Chiari malformation type I, and amyotrophic lateral sclerosis. Drug-induced yawning is an uncommon clinical condition and yawning in patients with autism or schizophrenia is very rare. The aim of this review is to describe in detail the occurrence of the phenomenon in such conditions, and its’ phenomenology and pathophysiology.

Keywords: yawning; parakinesia, stroke.

RESUMO
Bocejo é um comportamento estereotipado fisiológico, o qual, contudo, pode representar um sinal ou sintoma de várias condições neurológicas, tais como, acidente vascular encefálico, incluindo a parakinesia brachial oscitans, parkinsonismo, doença de Parkinson, e epilepsia. Mais raramente, o bocejo pode ocorrer em pacientes com hipertensão intracraniana, tumor cerebral, esclerose múltipla, migrânea, malformação de Chiari tipo I, e esclerose lateral amiotrófica. Bocejo-induzido por drogas representa outra condição clínica pouco comum. De outro modo, bocejo é muito raro em pacientes com autismo e esquizofrenia. O objetivo desta revisão foi descrever em detalhes a ocorrência deste fenômeno em tais condições, sua fenomenologia e fisiopatologia.

Palavras-chave: bocejo, paracinesia, acidente vascular cerebral.

Yawning is considered an intriguing and fascinating phenomenon with an obscure etiopathogenesis. It involves opening the mouth wide and closing the eyelids while inhaling deeply and then exhaling more briefly. A yawn typically lasts 5-10 seconds (Figure 1)¹²,⁴,⁵,⁶,⁷ and is usually accompanied by retroflexion of the head and sometimes by elevation of the arms (known as pandiculation when occurring together). Human beings yawn with a frequency of up to 28 times a day¹²,³,⁴,⁵,⁶,⁷, often after waking up and before falling asleep. Yawning is frequently contagious and is considered a sign of boredom or even disrespectful behavior in the presence of others⁸⁻¹⁰. In the 19th century, Charcot considered yawning an important clinical neurological sign, but for many years neurologists attached little importance to it¹¹. Nowadays, however, although yawning continues to be a largely under-appreciated behavior, chasmology, the study of yawning, has become the focus of considerable interest¹². The aim of this article is to review yawning in neurology.

¹Universidade Federal do Paraná, Hospital de Clínicas, Departamento de Medicina Interna, Unidade de Distúrbios do Movimento, Serviço de Neurologia, Curitiba PR, Brasil; ²University Health Network, Toronto Western Hospital, Gloria and Morton Shulman Movement Disorders, Toronto, ON, Canada; ³Universidade Estadual de Ponta Grossa, Hospital Universitário, Serviço de Neurologia, Ponta Grossa PR, Brasil; ⁴General Practice, Brou, France.

Correspondence: Hélio Afonso Ghizoni Teive; Rua General Carneiro 1103/102, Centro; 80060-150 Curitiba PR, Brasil. E-mail: hagteive@mps.com.br

Conflict of interest: There is no conflict of interest to declare.

Received 26 February 2018; Received in final form 26 March 2018; Accepted 09 April 2018.
METHODS
A literature search was performed in Pubmed, for English-language articles over the time period from 1992 to 2018, using the terms “yawning” and “CNS”, “neurology”, “neurologic disorders”. The initial references were selected using the following main exclusion criteria: 1) duplicate articles; 2) articles unrelated to the purpose of the study; 3) articles unrelated to yawning; 4) articles not including sufficient relevant data.

HISTORICAL ASPECTS OF YAWNING
Phylogenetically, yawning is a very old behavior that can be detected in most vertebrates from the fetal stages to old age. Its meaning has been the subject of speculation since antiquity. In some cultures, it has been associated with Satan (Arab countries) and spirits (India), while Hippocrates associated it with apoplexy. In the Hindu world, yawning in public is a religious offense. Since the 17th century, various explanations for its meaning have been put forward. Boerhaave, the founder of clinical medicine, stated in 1680, that yawns promote “the equitable distribution of spirits in all muscles and unblock the vessels”. In the 18th century, yawning was believed to be associated with the “improvement of brain oxygenation” and, in the 19th century, it was associated with different etiologies, including hysteria, and considered a reflex. Curiously, in 1888, in his famous Leçons du Mardi à la Salpêtrière, Charcot discussed the case of a 17-year-old patient who yawned eight times a minute, or 480 times an hour, and only stopped when she went to sleep. Her condition was believed at the time to be associated with hysteria and oxygenation of the nervous centers. She had generalized epileptic seizures, complete anosmia and binaural hemianopsia. She had been amenorrheic for a year. Charcot’s young patient was most likely suffering from a prolactinoma compression of her optic chiasma and her hypothalamus.

During the 20th century, substantial progress was made in understanding yawning, and several studies on this behavior and neurophysiology, hormones, dopamine receptors and social behavior were carried out. Despite this progress, yawning remains a mysterious and fascinating physiological, or even pathological, phenomenon.

THE NEUROPHYSIOLOGY OF YAWNING – A SUMMARY
There are several hypotheses to explain yawning: physiological hypotheses (such as the respiratory and circulatory system, arousal, sleepiness, thermoregulatory, ear-pressure and state-change hypotheses) and the social/communication hypothesis. In a study published in 2011, Guggisberg et al. concluded that the hypothesis best supported by experimental evidence is the social/communication hypothesis. In contrast, Gallup argued, in 2011, that the social/communication hypothesis is unattractive and suggested that yawning is probably multifunctional across species. In their view, the most acceptable hypothesis in humans is the thermoregulatory one, according to which yawning is a brain-cooling mechanism. In 2014, Walusinski proposed a new hypothesis, according to which yawning switches the default-mode network to the attentional network by activating cerebrospinal fluid flow. The default-mode network is a set of interconnected brain areas identified in functional neuroimaging (MRI) that exhibit spontaneous physiological activity during the normal resting state. There is a high level of activity in the default-mode network when the mind is not involved in specific behavioral tasks, and a low level of activity during focused attention. According to Walusinski’s hypothesis, yawning is a homeostatic process that appears when the default-mode network is active and sleepiness increases; yawning then disengages the default-mode network to promote the attentional network, which accelerates the circulation of cerebrospinal fluid, increasing clearance of somnogens such as adenosine, prosta glandin D2, VIP, prolactin and anandamide, and reducing sleepiness. The anatomic structures involved in yawning have yet to be elucidated and, more recently, fMRI has demonstrated that different levels of the neuroaxis, including the brainstem, prefrontal cortex and subcortical regions, may be involved. In 2010, Collins and Eguibar hypothesized that there are three main neural pathways involved in the regulation of yawning (Figure 2). Two of these are formed by groups of oxytocinergic neurons projecting from the paraventricular nucleus (one of the hypothalamic nuclei) to the hippocampus, pons, medulla and spinal cord; the other is formed by ACTH (adrenocorticotrophic hormone) and melanocyte-stimulating hormone-activated neurons projecting from the paraventricular nucleus to the hippocampus via activation of cholinergic neurons. There is also direct activation of hippocampal cholinergic neurons and a serotonergic-cholinergic pathway. In summary, several neurotransmitters and neuropeptides, such as dopamine, acetylcholine, serotonin, glutamate, GABA, adrennergics, excitatory amino acids, nitric acid, ACTH-related peptides, α-melanocyte-stimulating hormones and oxytocin, seem to be involved biochemically in the mechanism of yawning. Dopamine activates oxytocin production in the paraventricular nucleus of the hypothalamus, and oxytocin in turn activates cholinergic transmission in the hippocampus and the reticular formation of the brainstem, resulting in acetylcholine induction of yawning via muscle muscarinic receptors. In general, dopamine and its agonists trigger yawning, while opioid peptides and GABA reduce its frequency. Yawning is used as an indicator of dopaminergic and oxytocinergic transmission and, in Parkinson’s disease it is an expression of therapeutic dopaminergic activity, particularly as a marker of D3 dopamine receptor activity.
It is a popular belief that yawning is contagious\cite{10,18}, and it is well known that watching another person yawn may induce the person watching to yawn too\cite{10,18}. According to Guggisberg et al.\cite{2}, who support the social/communication hypothesis, contagious yawning is due to activation of a complex network of brain areas associated with imitation, empathy and social behavior\cite{15}. In 2005, Schürman et al.\cite{19} studied this intriguing phenomenon using fMRI. They observed activation of the left periamygdalar region, suggesting a connection between yawn contagiousness and amygdala activation.

In contrast, Platek et al.\cite{20}, using fMRI in 2005 to study contagious yawning, showed activation of the left periamygdalar region, suggesting a connection between yawn contagiousness and amygdala activation. In contrast, Platek et al.\cite{20}, using fMRI in 2005 to study contagious yawning, showed activation of the left periamygdalar region, suggesting a connection between yawn contagiousness and amygdala activation. In 2005, Schürman et al.\cite{19} studied this intriguing phenomenon using fMRI. They observed activation of the left periamygdalar region, suggesting a connection between yawn contagiousness and amygdala activation. Since the latter study, various papers on the relationship between contagious behavior, such as yawning and itching/scratching, and the activation of mirror neurons and the neurological mechanism of social behavior have been published\cite{19,20,21}. In 2014, Provine\cite{22} stated that contagious behaviors such as yawning have mirror-like properties.
Yawning can be present in the clinical picture of several neurological diseases, including stroke, multiple sclerosis, neuromyelitis optica spectrum disorders, Parkinsonism, Parkinson’s disease, migraine, vasovagal syncope, brain tumor, intracranial hypertension, Chiari malformation type I and epilepsy, and can be induced by drugs1,11.

**Yawning in sleep disorders**

In general, excessive yawning occurs more frequently in patients with sleep disorders (e.g., insomnia and sleep obstructive apnea) and is most common in children and young adults11.

**Excessive yawning in clinical conditions**

Other clinical conditions associated with yawning are functional digestive disorders (e.g., dyspepsia and irritable bowel syndrome), motion sickness and hypoglycemia in diabetic patients under insulin therapy11. In addition, vasovagal syncopal or presyncopal episodes frequently present with yawning as well as malaise, dizziness, visual obscuration, nausea, pallor and loss of consciousness11. Excessive yawning has also been reported in patients with depression and can be caused by the many medications used in neurology, psychiatry and internal medicine11. This issue is discussed in the following section.

**Drug-induced yawning**

Drug-induced yawning represents a rare and frequently not very serious adverse side-effect1,11,24. Antidepressants, dopaminergic agents, opioids and benzodiazepines are the main pharmacologic classes associated with yawning11,24. The most widely-used antidepressants are serotonergic agents (selective serotonin reuptake inhibitors), including fluoxetine, paroxetine, escitalopram, venlafaxine and duloxetine11,24. Dopaminergic drugs include levodopa, dopaminergic agonists (pramipexole, ropinirole, rotigotine and apomorphine) and monoamine oxidase inhibitors11,24. Blin et al.25 evaluated eight healthy volunteers, in 1990, after they used apomorphine and after a placebo, allowing 48 hours for the medication to be washed out. They found that this dopamine agonist induced blinking and yawning in all their participants. Another interesting finding is the occurrence of yawning in the withdrawal syndrome after prolonged use of opioids or coffee11,24. In 2007, Sommet et al.26 studied drug-induced yawning reported in the French Pharmacovigilance Database between 1985 and 2004. They found 28 reports involving 38 drugs, especially serotonergic agents, dopaminergic agents, opioids, and benzodiazepines26.

**Yawning and stroke**

Yawning has been observed secondary to acute hemorrhagic or ischemic anterior or posterior circulation stroke27,28,29,30. In general, it indicates damage to the brainstem reticular formation and cortical and subcortical structures27,29. In 2006, Cattaneo et al.29 published a case report of two patients with pathological yawning as a presenting symptom of brainstem ischemia involving the upper pons and the pontomesencephalic junction. Yawning has been described in patients with locked-in syndrome due to verteobobasilar stroke caused by a thrombosed megadolicobasilar artery30. Another potential cause of yawning in patients with stroke is intracranial hypertension or even herniation27. More recently, Krestel et al.23 studied 10 patients with acute anterior circulation stroke and yawning using neuroimaging (brain MRI with diffusion-weighted images). They found a correlation between ischemic lesions in the insula and caudate nucleus and a period of abnormal yawning and suggested that the insula may be the main region of the brain responsible for serotonin-mediated yawning23. A rare and very interesting clinical condition involving yawning known as parakinesia brachialis oscitans is related to stroke and will be described below.

**Parakinesia brachialis oscitans**

Occasionally, in patients with acute hemiplegia, the onset of yawning is associated with an involuntary raising of the paralyzed arm11,22. This phenomenon was named *parakinesia brachialis oscitans* in 2005 by Walusinski et al.31, who studied four cases and compared them with other cases published in the literature over the previous 150 years. Parakinesia brachialis oscitans was probably first described by Erasmus Darwin, the grandfather of Charles Darwin, in 1794, in his book “Zoonomia of the laws of organic life”31. In their seminal paper, Walusinski et al. proposed a possible mechanism for this abnormal involuntary movement. They suggested that resection of the cortico-neocerebellar tract of the extrapyramidal system disinhibits the spino-arheocerebellar tract, enabling motor stimulation of the arm by the lateral reticular nucleus, which has a link with respiratory and locomotor rhythms.31 Several case reports have since been published describing this phenomenon, including others by Walusinski et al. and a few other reports32,33,34,35,36,37. In 2013, Zorzetto et al.38 also published a case report of parakinesia brachialis oscitans in the setting of thrombolytic therapy (Figure 3). In 2015, Kang and Dhand36 published a case report of a 63-year-old man who presented with an acute ischemic infarct in the left middle cerebral artery distribution, aphasia and paralysis of the right arm, which consistently rose to his chest when he yawned. After yawning, paralysis always resumed. In a recent publication, Alves et al.39 describe a patient who has voluntary control over that movement.

**Yawning, parkinsonism and Parkinson’s disease**

In general, yawning is used as an indicator of activity of the dopamine-oxytocin pathway12. In patients with Parkinson’s disease (PD), it is associated with the presence of dopaminergic activity1,13 and has been considered an aura for an L-dopa-induced “on” state17. It is very well known that
PD patients who receive apomorphine injections (a direct D1/D2 dopamine receptor agonist) frequently present with yawning before positive motor effects, sometimes in association with penile erections\(^40\). Yawning was described by von Economo in patients with acute encephalitis lethargica or postencephalitic parkinsonism\(^8\). Curiously, Sandyk et al.\(^41\) reported a series of five patients with progressive supranuclear palsy (Richardson’s syndrome) who presented with balance deficits, falls, oculomotor abnormalities and bursts of repeated yawning without any correlation with their dopaminergic treatment. In 2013, Giganti et al.\(^42\) studied 18 untreated early-PD patients and compared them with 18 age-matched healthy controls. They showed that the circadian distribution of yawning was not altered in untreated patients with early-stage PD and concluded that yawning is a behavioral marker of sleepiness in \textit{de novo} PD patients\(^42\).

**Yawning and epilepsy**

Yawning can occur in different epileptic syndromes\(^11,43,44,45,46,47,48\). It has been described in the peri-ictal period in children with infantile spasms and in patients with temporal lobe epilepsy (peri-ictal yawning preceding complex partial seizures or yawning in the post-ictal period)\(^43,44,45,46,47,48\). In 2011, Specchio et al.\(^46\) reported ictal yawning in an adult patient with drug-resistant focal epilepsy. Kuba et al.\(^45\) investigated the incidence and lateralizing value of peri-ictal yawning in patients with temporal lobe epilepsy who had undergone surgery for epilepsy. They observed peri-ictal yawning in 4.1% of 97 patients and, in 1.8% of these patients’ 380 seizures, yawning occurred in the post-ictal period\(^46\). They also observed that peri-ictal yawning occurred only in patients with right-sided, nondominant temporal lobe epilepsy, suggesting that it may have a lateralizing value.\(^45\) Wasade et al.\(^47\) published an interesting case report, in 2016, of a young male who presented with controllable yawning expressed as focal seizures of frontal lobe epilepsy. In 2012, Nicotra et al.\(^48\) discussed the case of an elderly patient who presented with pathological yawning as an ictal seizure manifestation.

**Yawning and intracranial hypertension**

Some patients with intracranial hypertension, brain swelling or herniation due to stroke, head trauma or brain tumors may present with yawning together with headache and seizures\(^11\). In 2014, Saura et al.\(^49\) described intractable yawning associated with mature teratoma of the supramedial cerebellum. Yawning may also occur as a presenting symptom of Chiari malformation type 1\(^50\).

**Migraine and yawning**

Migraine represents a very common neurological problem, and the pathophysiology of migraine attacks has been shown to be related to dopaminergic transmission\(^51,52\). In the premonitory phase, yawning can be present in association with malaise, somnolence and mood changes among other manifestations\(^51,52\). Some migraineurs may experience excessive yawning during the visual aura before the attack\(^51,52\). However, excessive yawning in migraine can be improved with dopamine receptor antagonists, which are effective therapeutic agents for this condition\(^51,52\).

**Yawning, multiple sclerosis and neuromyelitis optica spectrum disorders**

Some patients with multiple sclerosis (MS) present with thermoregulatory dysfunction\(^53\). Therefore, sleep problems and yawning could be symptoms of MS\(^51,52\). Gallup et al.\(^54\) also showed that yawning can provide symptom relief in MS patients. In 1996, Postert et al.\(^54\) published a case report on pathological yawning as a symptom of MS and, in 2014,
Yawning and amyotrophic lateral sclerosis

Yawning can be observed in patients with amyotrophic lateral sclerosis (ALS)\(^5\). Williams\(^6\) published a case report, in 2000, of a 64-year-old woman who presented with progressive, very frequent yawning characterized by bouts of 20 to 30 successive yawns in association with painful cramping and a sensation of choking. During the follow-up, the patient developed bulbar palsy, and an electromyogram confirmed the diagnosis of amyotrophic lateral sclerosis\(^6\). In 2007, Wicks\(^7\) published a paper in which he noted that excessive yawning is a common sign in the bulbar-onset form of amyotrophic lateral sclerosis.

Yawning, autism and schizophrenia

Contagious yawning can be explained by the relationship between yawning and social empathy\(^2\)\(^,\)\(^3\)\(^,\)\(^4\)\(^,\)\(^5\)\(^\)\(^,\)\(^6\)\(^,\)\(^7\). Interestingly, in some psychiatric disorders, such as autism and schizophrenia, yawning is very rare\(^8\)\(^,\)\(^9\)\(^,\)\(^10\)\(^,\)\(^11\)\(^,\)\(^12\). In 2007, Senju et al.\(^13\) evaluated 24 children with autism spectrum disorder and 25 age-matched, normally-developing children, in terms of their frequency of yawning, using video clips of either yawning or control mouth movements. They concluded that contagious yawning is impaired in autism spectrum disorder, supporting the idea that contagious yawning is based on the capacity for empathy\(^14\). Haker and Rössler\(^15\), in 2009, compared changes in yawning patterns in a group of 43 schizophrenic patients and an age- and sex-matched group of healthy controls using a video with sequences of yawning, laughter and neutral faces. They observed that schizophrenic patients showed a significantly lower rate of yawning, suggesting that susceptibility to contagious yawning is reduced in patients with impaired social empathy\(^16\).

Yawning – problems and solutions

Yawning can sometimes cause complications, such as mandibular subluxation, painful cramps in the geniohyoid muscle and fracture of the styloid apophysis\(^17\). More rarely, it can trigger an attack of glossopharyngeal neuralgia, idiopathic carotidynia or even Marin Amat syndrome, a form of acquired facial synkinesis manifesting as involuntary eyelid closure on jaw opening that often occurs following the aberrant regeneration of the facial nerve after a peripheral facial palsy\(^18\)\(^,\)\(^19\). In contrast, yawning can improve Eustachian tube dysfunction and dysbaric facial paralysis and even acts as a therapy for vocal fatigue\(^20\). Gallup and Gallup\(^21\) studied the frequency of yawning as an initial signal of fever relief.

CONCLUSION

Yawning is a stereotyped physiological behavior that can be a sign or symptom of several neurological conditions, such as stroke (including stroke with parakinesia brachialis oscillans), parkinsonism and epilepsy. More rarely, yawning can occur in patients with intracranial hypertension, multiple sclerosis, migraine and amyotrophic lateral sclerosis and can be induced by drugs. It is rare in patients with autism spectrum disorders or schizophrenia\(^12\)\(^,\)\(^13\)\(^,\)\(^14\)\(^,\)\(^15\)\(^,\)\(^16\)\(^,\)\(^17\)\(^,\)\(^18\)\(^,\)\(^19\)\(^,\)\(^20\).

References


