Abstract

Yawning is a physiological behavior and, as with all such behaviors, its deregulation is indicative of an underlying disorder. This chapter breaks this topic down into the types of yawning (incomplete, absent or excessive) and the direction of causality (triggering or relieving a disorder).

Yawning is a physiological behavior, an emotional stereotypy that indicates the homeostatic process of the mechanisms regulating rhythms – such as sleeping-waking, hunger-satiety or mating-relaxation – generated by the diencephalon. As with all physiological behaviors, its deregulation reveals disorders. However, a property unique to yawning, which is nevertheless physiological, is its ability to trigger certain disorders and, conversely, to cure others!

Can one talk about the ‘disease of yawning’? Here is what H. Meige and E. Feindel wrote in 1902 in their famous book *Tics and Their Treatment* [1, p. 346]:

Yawning and sneezing, which, like swallowing, are reflexive phenomena whose mechanism is voluntarily modified only with great difficulty, can hardly be perturbed except with respect to their frequency. Saenger (Observations on the idiopathic spasm of the tongue, *Monatsschrift für Psychiatrie und Neurologie*, January 1900, p. 77) reported on the case of a 29-year old woman who was not hysterical and who presented with episodes of yawning with rigidity in the arms, followed by rapid contractions of the tongue for around a minute. She recovered after several months. The case concerned an ‘idiopathic spasm’, probably some sort of tic. However, these functional anomalies of yawning or sneezing most often occur in hysterical subjects. It should also be remembered that yawning may be the aura of an epileptic seizure. Uncontrollable yawning is also seen in meningitic incidents, and in cases of tumors of the brain or cerebellum.

The analysis of clinical observations in light of the neurophysiological knowledge accumulated over a century, allows us to affirm the existence of pathological yawning, and to distinguish between its various forms: the dissatisfaction of incomplete yawning, the disappearance of yawning and the excess of repeated yawning.
Anhedonia

Frustration due to the incomplete or inharmonious development of a yawn is a frequent complaint. Yawning stretches the respiratory and facial muscles, and is sometimes associated with a generalized muscular stretching of the trunk and limbs. The yawner perceives differences in muscle tonus, a veritable and conscious extraction of the progress of the phenomenon, of its stimulus and its contextual valence, through interoceptive pathways (projections by the lateral spinothalamic tract onto the insular cortex) leading to a hedonistic perception. The dissatisfaction felt seems to be linked to an unconscious inhibition of the 'letting go' that underlies a complete yawn. Therapeutic interventions for anxiety states or for insomnia, such as relaxation or sophrology, do in fact use the control of yawning as a means of relaxation or as an anti-stress aid, a veritable proprioceptive rehabilitation of the body scheme [2, 3].

Disappearance of Yawning

Yawning is a banal action, often barely noticed or felt. It seems that its disappearance is not perceived. Complaints of missing the feeling of well-being associated with yawning, due to the absence of yawning, remain the exception. In daily life, there appear to be no particular consequences to not yawning. The reality of this fact can be questioned, since some yawns could occur without being felt or noticed. Certain extrapyramidal syndromes are accompanied by a disappearance of yawning, such as in the case of treatment with neuroleptics [4] or Parkinson's disease [5]. This disappearance indicates the activity state of the dopaminergic neurons of the paraventricular nucleus of the hypothalamus, which are necessary for yawning. Experimental pharmacology has shown the specificity of the D₃ dopamine receptors in triggering this behavior [6]. While the neuroleptics currently in use have no specific target of action, one may assume that the presence or absence of yawning in Parkinson's disease reflects the state of the neuronal population: functional or undergoing degenerative involution. The treatment of episodes of motor blocks in Parkinson's patients by apomorphine hydrochloride, a rapid-action dopaminergic stimulant, triggers yawning, as does treatment with L-DOPA, but in a less systematic fashion [7–9]. Dewey et al. [10] noted this effect in 8% of the patients they treated, and its absence in the case of a placebo injection. Yawning was clearly described by the patients not as a form discomfort, but as a signal of unblocking, which indicated the beginning of the effect of the relieving treatment [11–14].

Excessive Yawning

The most common cause of frequent and repeated yawning is sleep debt, particularly in children and young adults. Campaigns for the prevention of falling asleep while
driving emphasize this warning sign of the risk of falling asleep involuntarily. Research on accident prevention is leading to the development of programs for the automatic detection of yawnning and blinking by the driver, in order to trigger an alarm that forces the driver to stop [15]. Drowsiness can be assessed by establishing an Epworth score, in order to uncover a syndrome of sleep apnea or another cause of hypersomnia. In children, drowsiness is often manifested by excessive agitation, punctuated by yawnning, with an attention-fixing deficit. Before invoking the diagnosis of a possibly hyperactive child, it is advisable to eliminate any sleep apnea syndrome caused by an obstructive hypertrophy of the tonsils or adenoids. Evans [16] describes, in the same spirit, two cases of inhalation of foreign bodies into the bronchi, which manifested as suffocation alternating with yawnning in succession.

Dyspepsia or the sensation of a full stomach and slow digestion, and an irritable colon are often associated with salvoes of yawns. Considering the importance of the autonomic nervous system, and the parasympathetic system in particular, in the regulation of digestive functions, it is not surprising that yawnning appears to be associated with problems that are essentially functional. The term ‘gut brain’ or digestive neurology has been used in this context, in order to avoid restricting the pathophysiology of this system to motor deficits, and to also take into account the sensitivity of the digestive system (viscerosensation, a component of interoception) [17, 18]. The discovery of hypocretin, a neuromediator that triggers sensations of hunger and arousal, could also provide an explanation [19]. It is possible that leptin, the messenger of satiety, and ghrelin, another messenger signaling hunger, play a role that has not yet been elucidated [20]. These functional digestive disorders should be seen in light of a vasovagal episode. The circumstances under which it is triggered are multiple: violent emotion, intense pain, the sight of blood, staying in a confined overheated space, etc. The most common form of the malaise, a loss of consciousness – which occurs in subjects of all ages, regardless of their state of health – is preceded by a rich parade of premonitory disorders that attest to a hyperstimulation of the parasympathetic system: pallor, nausea, salvoes of yawnning, visual disturbances, etc. The appearance of yawnning is a sign that should attract the attention of a health-care worker during invasive exploration, and allow him/her to anticipate a loss of consciousness and subsequent fall. Dorsal decubitus, or an injection of atropine, eliminates the malaise and the yawnning [21]. Motion sickness or kinetosis is a related functional disorder that is often accompanied by repetitive yawnning before the onset of vomiting [22]. The beginning of hypoglycemia in a diabetic receiving insulin therapy is accompanied by a feeling of hunger, profuse sweating and repeated yawnning, similar to the feeling of hunger in non-diabetics.

With a progression that is often insidious, an increase in the frequency of yawnning becomes embarrassing both because of the brief pause in activity that it causes, and because of the negative social connotations perceived by both the yawner and the company they are in. Often occurring in salvoes of 10–20 successive yawns, the daily number often exceeds 100. Thus, the famous patient presented at the Leçons
du Mardi de la Salpêtrière on October 23, 1888 by Prof. J.-M. Charcot [23] yawned 8 times a minute, or 480 times an hour! The most frequent cause, in our times, is of iatrogenic origin. Numerous medications used in neurology and in psychiatry lead to an increase in the frequency of yawning. We will review them here in decreasing order of frequency.

Antidepressants, in particular serotonergic antidepressants, are the ones most often found to be involved [24, 25]. The effect involves the entire pharmacological class, and all molecules studied have been implicated (fluoxetine, paroxetine, escitalopram, duloxetine, venlafaxine, etc.) [26–28]. This symptom is very often wrongly interpreted both by patients and their treatment providers. Attributed to asthenia, the persistence of a depressive state or a lack of effectiveness of the treatment, this frequent repetitive yawning can wrongly lead to an increase in the recommended dosage, which in turn accentuates the problem, whereas stopping the treatment would allow the symptoms to disappear within a few days. There is never any associated somnolence. This effect is sometimes accompanied by erection of the clitoris and involuntary orgasms [29]. Serotonergic antidepressants are nevertheless considered to be better tolerated than tricyclic antidepressants or monoamine oxidase inhibitors. It is not easy to interpret the somewhat inconsistent mechanism controlling this side effect. In fact, in addition to their activity as serotonergic agonists (potentially involving 5-HT₄ receptors), these molecules also have adrenergic, muscarinic cholinergic and histaminergic effects [30, 31]. Unfortunately, as this is only rarely reported to pharmacovigilance agencies, there are no statistics to assess the frequency of this iatrogenic effect, nor any studies showing that it is revealed by association with another psychotropic drug [32]. Curiously, while tricyclic antidepressants have atropinic side effects, thus being inhibitors of yawning, and are reputed to lead to impotence, there have been reports regarding excessively frequent salvos of yawning that accompany involuntary orgasms with clomipramine [33, 34].

Concerning dopamine agonists, we have already discussed apomorphine hydrochloride injected during episodes of motor block in Parkinson’s patients in order to restore motor function. It is not possible to invoke a real iatrogenic effect here. Apomorphine hydrochloride, at lower doses administered orally, is used to treat male impotence. Data sheets accompanying the product indicate yawning as a rarely reported side effect seen during the initial clinical studies. Following commercialization, no follow-up data are accessible. Data for other dopaminergic agonists (bromocriptine, lisuride, pergolide, ropinirole, pramipexole, selegiline and piribedil) are no doubt lacking as a result of faulty pharmacovigilance, since animal models show evidence of the same iatrogenic risks [35, 36].

Detoxification after prolonged use in heavy users of coffee or of opiates is accompanied by a withdrawal syndrome that includes the occurrence of repetitive yawning over several days [37–43]. The clinical profile of a neonate with a mother who consumes opiates, legal (methadone, morphine) or not, until the end of her pregnancy,
can be striking, with respiratory irregularities and pauses interspersed with salvos of yawning [44, 45].

Migraine is one of the most common disorders in humans, affecting 10–20% of the population at least once per year. It appears to be secondary to a combination of environmental and genetic factors. Clinical and pharmacological findings, as well as recent developments in genetics, confirm the hypothesis that a dysfunction in dopaminergic transmission plays a role in the pathophysiology of a migraine attack. Prodromes (changes in mood, yawning, somnolence, food aversion, etc.) may be related to dopaminergic overstimulation. The dopaminergic system also plays a role in the headache phase, on the one hand by participating in nociceptive pathways, and on the other by intervening in the regulation of cerebral arterial circulation. Apomorphine induces more yawning in migraine sufferers than in non-migraine sufferers [8, 46–48]. The shimmering blind-spot is a classic sign of a visual aura. However, a large number of sufferers have noted that repeated yawning in salvos also acts as an aura before an attack [49–51]. More rarely, the attack ends with repeated yawning, accompanied by drowsiness and a postdrome profile [52, 53].

Yawning can be studied from several angles during the course of a stroke. During the occurrence of an attack (ischemic or hemorrhagic) deficits in vigilance occur, accompanied by salvos of yawning, whether or not the victim is conscious [54–56]. This could be due to intracranial hypertension resulting from the stroke. In the case of a deep coma (Glasgow score = 3), the occurrence of repetitive yawning is a sign of herniation, a grave prognostic sign. Apart from this extremely serious evolution, yawning during the course of a stroke indicates damage to cortical and subcortical circuitry, and to a mechanism of secondary vigilance stimulation controlled by the reticular formation of the brainstem, a mechanism probably common to the yawning that occurs during a partial seizure in temporal lobe epilepsy. During an ischemic attack affecting the territory of the lenticulostriate arteries, damage to the internal capsule and/or the lentiform nucleus leads to complete hemiplegia due to the lesioning of the pyramidal tracts, while the extrapyramidal pathways are spared. In this case, during yawning, the paralyzed arm can be seen to move, bringing the hand up to the mouth. The arm drops immediately following the end of the yawn. We have named this syndrome, which is not a synkinesis, ‘parakinesia brachialis oscitans’. With regard to its phylogeny, it has been shown that in quadrupeds like the dog and the horse, there is a synchronization of the ventilatory rhythm with that of gait: one ventilatory cycle per gait cycle, with concomitant acceleration while running. In humans, bipedalism has led to the loss of this automatic synchronization, retaining only the swing of the arms while walking, but no strict synchrony with ventilation. After a stroke has interrupted cortical control, the subjacent neurological structures retrieve their ancestral functions, which are normally inhibited by the overlying cerebral structures as a result of evolution. During the movement of the diaphragm while yawning, the paralyzed arm receives motor stimulation from the lateral reticular nucleus of the medulla, which couples ventilation and locomotion in animals, an extrapyramidal signal that is not
inhibited by the ischemic lesion. In two stroke profiles, the persistence of yawning and emotional facial expressions signals the dissociation between automatic and voluntary pathways [57]. In the case of ‘locked-in’ syndrome caused by an occlusion in the basilar artery territory, there is quadriplegia, in association with bilateral facial paralysis. However, physiological yawning continues to occur [58–60]. Similarly, in bilateral anterior opercular syndrome or Foix-Chavany-Marie syndrome, the muscles of the face, tongue and laryngopharynx are paralyzed during all voluntary acts, including voluntary smiles or grimaces, as well as during language articulation, whereas the expression of emotions, eye blinking, laughing, coughing, swallowing and yawning remain possible. No voluntary imitation of these movements can be carried out. The cause is ischemic, or rarely post-traumatic bilateral damage to the frontoparietal opercular areas [61, 62].

Intracranial hypertension, whether related to a stroke, tumor [63–65] or head trauma [66], can be revealed by headaches and by disturbed vigilance associated with salvos of yawns and with convulsions. Certain coma scores used in the USA take into account the presence of yawning in these situations [67, 68]. Vegetative states of postischemic or other origins also display ‘automatic-voluntary dissociation’ with the persistence of frequent yawning [69].

J.-M. Charcot presented a patient in 1888, referred to previously, and reported by Gilles de la Tourette [70] in 1890 as suffering from hysteria. Nevertheless, this young woman of 23 years was amenorrheic, epileptic, suffered from a bitemporal loss of the visual field and yawned 480 times an hour. In reality, she was probably developing a prolactinoma of the pituitary gland. I have also personally observed a 39-year-old acromegalic male with prognathism and moderately protruding brows, suffering from persistent asthenia with salvos of almost 200 yawns per day [unpublished data]. Similarly, Wong [65] reported a case of a mucocele of the sphenoid sinus compressing the pituitary stalk, which was revealed by the occurrence of yawning repeated every 15 s. There are a number of clinical arguments to show that an unexplained excess of yawning could be the result of a hypothalamic-pituitary disorder, the mechanism of action of which could be the oversecretion or inappropriate release of oxytocin or other neuromediators due to compression [71, 72].

There has been little interest in the behavioral disturbances that precede or succeed an epileptic seizure by a few minutes to several hours. Nonetheless, these anomalies could provide orienting data that would help to localize the anatomical origin of focal seizures. Before as well as after a temporal, or sometimes a frontal, epileptic seizure, different automatisms such as rubbing the nose with the fingers, yawning or sighing can be observed. J.H. Jackson [73] wrote in around 1876:

These symptoms do not occur during but after the paroxysm of the seizure; these are movements that are too well coordinated to result from an epileptic discharge; there exists, I think, a double condition: (1) negatively, a loss of control; (2) positively, an augmentation of the activity of the inferior functional center. In any case, the association or the sequence of gestures is very significant.
In accordance with behavioral data, there is a phylogenetic origin to these stereotypical behaviors. They are noticeable during the fetal stage, and continue into the postnatal period and throughout life. Scratching the face, rubbing the nose, yawning and sighing have been described as automatic behaviors that occur before or after absence seizures or focal seizures. They can also be seen in healthy subjects as they awake from sleep. They can be distinguished as being physiological (upon waking) or pathological (for example, in temporal lobe epilepsy) depending on whether their velocity is harmonious or not, and whether their repetition is brief or prolonged. These behavioral automatisms are related to the activation of the brainstem or the spinal cord, the seat of their motor and integrative centers. The cortex, where the seizure is localized, is not involved. Thus, we see the reappearance of ancestral behavioral automatisms that are necessary for survival (such as walking, swimming, mating and other rhythmic activities, to which yawning also belongs), by a liberating ictal cortical disconnection [74–78].

Goldie and Green [79] presented three observations taken from old reports of Gowers [80], Penfield and Jasper [81] and Symonds [82], of children suffering from ‘petit mal’ seizures, the beginnings of which are signaled by repeated yawning. In addition to the association between temporal lobe epilepsy and yawning, Penfield and Jasper [81] described a rare type of epilepsy, diencephalic epilepsy, the existence of which has been questioned by some epileptologists. It brings together the brutal stimulation of sympathetic and parasympathetic activity: a feeling of disconnection from the environment without loss of consciousness, a ‘vasomotor storm’ with flush, profuse sweating, rapid rise and fall of the body temperature, pulse and arterial pressure, lacrimation, salivation, pupillary inequality and irregularities of the ventilator rhythm. Salvos of yawning and irrepressible hiccupping accompany this spectacular and very unpleasant profile. The cases described reveal the presence of thalamic tumors [83–87]. D’Mello et al. [88] also reported salvos of yawns as an iatrogenic effect of vibratory massage or of the withdrawal of neuroleptics, persisting several months after convulsions are provoked. He proposed a lesioning of the hypothalamus by way of explanation.

Occurring at the threshold between absence seizures and epilepsy of a psychogenic origin, prolonged hyperventilation (voluntary or unconsciously induced) can alter the level of consciousness, mimicking an epileptic fit, with the appearance of automatic movements such as smiling, swallowing or yawning [89, 90]. Positive or negative emotions are accompanied physiologically by modifications in cardiorespiratory activity, mediated by the autonomic nervous system. Thus, an increase in the frequency of ventilation is one of the signs of a panic attack. The hyperventilation syndrome comprises a neurotic profile on its own, leading to respiratory alkalosis, accompanied in more than 30% of cases by repeated widely spaced yawns that could be seen as a homeostatic counterstimulation of the parasympathetic system [91–95]. The use of relaxation techniques and yoga designed to trigger relaxing yawns allows these problems to be treated [96].
In 2006, in a study published exclusively on the internet, SR Gallezzo of Holyoke Community College (Mass., USA) [97] demonstrated the existence of a link between the frequency of yawning reported by depressive patients and the severity of the depression, using a small sample of 31 patients. Depression was evaluated with the help of the Goldberg scale, and yawning on the basis of a questionnaire proposed by Greco and Baenninger [98]. Statistical analysis of the results demonstrated a significant link: the more the patient yawned, the severer the depressive state. This was the first time that such a study was performed. To the surprise of the author, there was a negative correlation between age and yawning. It should be noted that the number of yawns produced diminishes physiologically with age, and that the predictive value of the relation between yawning and depression is probably lost after 65 years. It should also be remembered that antidepressants, in particularly serotonergic antidepressants, promote repeated yawning.

It is at present quite rare to encounter a schizophrenic who does not take neuroleptics, which, by their mode of action, inhibit yawning. In the past, the reappearance of yawning in a schizophrenic was interpreted as a resumption of contact with the environment and socialization [99]. Salvos of yawns, such as those seen in at least three of the five cases reported by Gilles de la Tourette in 1890, could have a psychogenic cause, as a form of non-verbal language. These have been described as an urgent and irrepressible feeling, following a sensation of epigastric heaviness with ascendant retrosternal constriction, relieved by undergoing a ventilatory period of yawning that yields a brief feeling of pleasure such as those described by patients affected by nervous tics [100, 101].

Sandy [102] reported a series of 5 patients in the initial stages of Steele-Richardson-Olszewski syndrome, or progressive supranuclear palsy, with balance deficits, oculomotor problems and salvos of repeated yawning that were reduced, paradoxically, by the administration of dopaminergic agonists. Louwerse [103] reported, in a series of 200 patients with the bulbar form of amyotrophic lateral sclerosis or Charcot’s disease, the existence of excessive yawning in salvos in around 10% of them, which is also the case in the study carried out by Williams [104]. Wicks [105] saw this problem in 47% of patients (n = 539). Present during the initial phase of the disease and concurrent with the appearance of swallowing deficits, the yawning disappears with the aggravation of paralysis. A brainstem attack in multiple sclerosis can yield the same profile [106].

Between 1917 and 1925, an epidemic of encephalitis described as lethargic by C. von Economo (1876–1931) [107] struck the whole of Europe. After an initial meningitic phase, a lethargic phase lasting several weeks set in. These acute symptoms, often fatal, were followed in those who survived by a ‘chaos of abnormal movements’ (Pierre Marie), with the mixing of depressive or delirious manifestations, oculogyric crises, Parkinsonian syndromes, etc. In 1921, Sicard and Paraff [108] reported observations in which the initial lethargic phase was followed by spasmodic attacks of hiccupsing, laughing or crying, and salvos of repetitive yawning several times a
day. The homogeneity of the process of morbidity and the relative consistency of the lesion sites were in contrast to the polymorphism of the clinical manifestations. Contemporary discoveries confirm the soundness of the hypotheses put forward by C. von Economo. Autoimmune injury to the basal ganglia of the hypothalamus leads to a rarefaction of orexin/hypocretin-containing neurons, among others, and decreases, either transiently or permanently, the secretion of certain neuromediators responsible for arousal.

**Yawning Triggers a Disorder**

The act of yawning is not merely a simple opening of the mouth but a complex coordinated movement, bringing together a flexion followed by an extension of the neck, a wide dilatation of the laryngopharynx with maximal descent, and an opening of the eustachian tubes [109]. Each of these elements can lead to pain or to dysfunctions.

Yawning is the most frequent cause of mandibular subluxation, ahead of laughing, vomiting and dental care. The subluxation is anterior; in other words, the mandibular condyle juts out in front of the temporal condyle and into the temporal fossa, from which it cannot be extricated due to the tonicity of the masticator muscles. The subluxation is most often bilateral. It is generally possible to reduce it promptly without anesthesia [110, 111]. Relapses occur frequently. Although this type of luxation may occur during normal physiological yawning [112–115], several publications have described mandibular luxation as a complication of uncontrollable yawning while on serotonergic antidepressants [116, 117] or during the induction of anesthesia [118–120].

In addition, yawning can trigger painful cramps in the geniohyoid muscle [121].

When a patient presents with otalgia (showing no noticeable lesion of the ear) or for atypical facial pain, the presence of pain aggravated by yawning is a good indication of a disorder of the temporomandibular joints, such as arthritis, ankylosis, mandibular dysfunctions or myalgias of the masticator muscles [122–125].

Yawning can be the release mechanism or the trigger for an attack of glossopharyngeal neuralgia [126, 127]. As in the case of swallowing, it can initiate choreoathetotic movements of the lips and the laryngopharynx, following a brainstem stroke [128]. In 2001, D.E. Jacome described two patients with paroxysmal headaches, in whom the only triggering factor was yawning [129, 130]. In 2004, he described two other cases in which yawning led to extracephalic pain that appeared neuralgic [131]. In the first case, neck pain radiated towards a shoulder, and in the second, intense pain invaded the anterior portion of the neck, where a thyroid tumor of the clear-cell carcinoma type was developing. In both cases, a spontaneous wide opening of the mouth did not trigger pain; this was only triggered by yawning. Another clinical type of neuralgia that can be triggered by yawning is short-lasting neuralgiform unilateral pain affecting the orbital-periorbital area and associated with autonomic phenomena
consisting mainly of conjunctival injection, tearing, and rhinorrhea (Charlin-Sluder syndrome) [132].

The Marin Amat syndrome (or inverted Marcus Gunn syndrome) consists of a synkinetic closing of the eyelids of one eye while opening the mouth, notably during yawning [133].

Fractures of a long styloid apophysis are rare. Throat or neck pain, reduced neck mobility, hoarseness and a lateral mass in the neck are all signs that point to this condition. They occur secondary to yawning or a direct shock, such as those encountered during a traffic accident [134, 135]. These should prompt a search for an Eagle’s syndrome – in which lateral neck pain, triggered by yawning or mastication, is associated with the pressure exerted by the styloid process within the tonsillar fossa [136, 137]. X-rays of the neck reveal the calcification of the stylohyoid ligaments (a discovery that sometimes occurs by chance), which becomes symptomatic whether or not there is a fracture [138, 139].

Idiopathic carotidynia is a controversial diagnosis, based on clinical arguments. The international classification of headache disorders established by the International Headache Society proposes the presence of at least one of the following criteria in order to retain this diagnosis: localized hypersensitivity, swelling and an increase in pulsations. The pain is typically unbearable, pulsatile, permanent and localized at the carotid bifurcation, but it may radiate towards the mandible, cheek, eye or ear. The symptoms are often exacerbated by swallowing, chewing, yawning and contralateral movements of the neck. The cardinal clinical sign is an exaggerated sensitivity of the carotid bulb to palpation, which often reveals prominent swelling with an accentuation of visible and palpable pulsations. Other serious causes (dissection of the carotid) should be excluded. Its evolution appears to be benign in most cases [140, 141].

Yawning Relieves a Disorder

The eustachian tubes play a role in the aeration of the middle ear, or pressure equalization [142]. Physiologically, the eustachian tubes open during swallowing due to the contraction of the peristaphyline muscles. Defective opening occurs as a result of tube dysfunctions. These could be a consequence of inflammatory phenomena affecting the ear and throat, or functional phenomena, such as during variations in pressure during air travel. The powerful contraction of the peristaphyline muscles during yawning explains its airing effect on the tympanic cavity, which serves to restore hearing in case of serous otitis or during rapid descent while in flight [143, 144].

Dysbaric facial paralysis is a rare event, but one which usually has a favorable outcome. It is linked to a change in pressure in the medium in which the victim evolves, and is responsible for ischemic neuropraxia of the facial nerve, e.g. during diving or flying. If hyperbaric oxygen therapy is not available immediately, either the Toynbee
maneuver, provoked yawning or swallowing can accelerate recovery and limit the risk of sequelae [145].

The extrinsic or intrinsic musculature of the larynx is particularly sensitive to emotional stress. Its hypercontraction is a common denominator in all forms of functional or psychogenic dysphonia and aphonria. Yawning-whistling is a technique that has come into increasing use over the last few years, in particular as a therapy for vocal fatigue. The technique is particularly effective in combating the excessive elevation of the larynx and constriction of the glottis that characterize vocal fatigue. This condition corresponds to an excessive muscular effort by the respiratory, vocal and resonance systems, which tends to raise the larynx. This technique aims to open the glottis and to reposition the larynx at its lowest point in order to reduce muscular effort to a minimum, which is what yawning accomplishes physiologically.

Pieter Brueghel the Elder painted, in around 1560, a work posthumously named *The Yawner*, currently on display at the Royal Museums of Fine Arts in Brussels. Marsden [146] contests this interpretation and believes that it depicts a blepharospasm or oromandibular dystonia, the contraction of which can be relieved by yawning, as noted by a good number of patients.

**Concluding Remarks**

At the end of this vast panorama, the consultation of a patient complaining of excessive yawning can be schematized. The first step consists of looking for an iatrogenic effect, the most frequent cause. Then, a search for abnormal drowsiness or sleep debt should allow any syndrome of sleep apnea to be uncovered. Functional causes leading to an anxiety disorder, possibly associated with a hyperventilation syndrome, should be treated by relaxation and yoga. This should be correlated with the occurrence of dyspepsia. Clinical examination is needed to detect pituitary/hypophyseal endocrine anomalies, intracranial hypertensive syndromes, partial temporal seizure, stroke or Charcot’s disease. Finally, repeated yawning could form part of a tic disease.

**References**


