The concept behind this article is that respiration is not just an automatic function. Rather, it is a complex behavior that is controlled at many brain levels and structures, some autonomic and others voluntary. This article will offer a review of the organization of respiratory control and its special features as gleaned from work in animals, and from relatively recent studies in humans using updated imaging techniques. The importance of the brainstem in autonomic respiratory control, of sleep-wake states in modulating respiration, and of supramedullary regulation of breathing during volition, with emotion, and at the onset of exercise hyperpnea will be emphasized. Clinical cases will be described, including space occupying lesions in the
Respiration is also subjected to autonomic regulation, which is extensively illustrated by the case history of Mr. MDC (courtesy Dr. Rolando Del Maestro, Department of Neurosurgery, Montreal Neurological Institute, McGill University). At thirteen years of age, because of symptoms of increased intracranial pressure, this patient was diagnosed as having an invasive ependymoma, both in the fourth ventricle and in the posterior fossa. The tumor was partially removed and the region irradiated. The patient was lost to follow-up until recently, when, at the age of forty-five, due to recurring motor problems, he was discovered to have extensive gliosis and fibrosis in and around the brainstem (Figure 1).

While hospitalized, respiratory gas monitoring showed severe hypoxemia and hypercarbia, especially during sleep. These were easily alleviated with mechanical ventilation. Indeed, on one occasion, the patient refused artificial ventilation, resulting in worsening of his respiratory status to the point of coma, at which point artificial ventilation again corrected the respiratory gases and the state of consciousness. This sequence indicated that the underlying pathology of these respiratory findings was, at least in part, due to central alveolar hypoventilation, that is, to an inability of respiratory control mechanisms in the brainstem to adequately regulate breathing so as to ensure chemical homeostasis.

A case history showing the importance of the brainstem for normal breathing

The crucial role of the brainstem in normal respiratory control can be illustrated by the case history of Mr. MDC (courtesy Dr. Rolando Del Maestro, Department of Neurosurgery, Montreal Neurological Institute, McGill University). At thirteen years of age, because of symptoms of increased intracranial pressure, this patient was diagnosed as having an invasive ependymoma, both in the fourth ventricle and in the posterior fossa. The tumor was partially removed and the region irradiated. The patient was lost to follow-up until recently, when, at the age of forty-five, due to recurring motor problems, he was discovered to have extensive gliosis and fibrosis in and around the brainstem (Figure 1).

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The organization of respiratory control: Brief review

To explain this patient’s clinical respiratory findings, a review of the organization of respiratory control is in order.

First, autonomic respiration is controlled in the brainstem (Figure 2A). Current concepts indicate that the predominant, if perhaps not exclusive, origin of the drive to breathe emanates from a relatively small region in the rostral ventrolateral medulla named the preBötzinger complex. This region in the in vitro preparation of the neonatal rat comprises ~600 neurons. Of these neurons, the crucial ones are those that display an inherent inspiratory oscillatory activity and that can be identified by being endowed with both neurokinin-1 and mu-opioid receptors (the endogenous peptides of which include substance P and endomorphin-1 and endomorphin-2). Whereas, for several years after its discovery, the preBötzinger complex was considered the sole ‘kernel’ of central respiratory drive, recent studies have questioned its exclusive role, and have added neighboring neuronal networks, including neurons with pre-inspiratory activity, to the overall model of how central respiratory activity is generated. It is important to emphasize that, whereas neuronal structures and connections have been defined in some detail from animal studies, it is highly likely that these are fundamentally similar in all mammalian species, including man, although the relative size and location of various components may vary among species.

Respiratory drive is transmitted via cranial motor nuclei, such as the vagus and hypoglossus, and via the ventral respiratory neuronal group and bulbospinal pathways to upper airway, diaphragmatic, intercostal and accessary respiratory muscles (Figure 2B). The feedback loop for this autonomic respiratory drive is completed by afferent stimuli from mechanoreceptors in the lung and respiratory muscles, and from O2 and CO2 chemoreceptors. With regard to the chemoreceptors, it is currently believed that, in addition to their peripheral locations, their central distribution is extensive. This newer concept will become important as we try to explain the symptoms seen in the patient described above. Other structures in the brainstem also participate in autonomic respiratory control, including the solitary tract nucleus in the dorsal brainstem that is important for the integration of sensory autonomic information, the pontine parabrachial nuclei that participate in respiratory timing, and the sleep-wake-related gigantocellular, medullary raphé, lateral reticular, and locus coeruleus nuclei (Figure 2A).

Respiration is also regulated by many structures outside the brainstem. There is mounting evidence that extramedullary regions, such as several hypothalamic nuclei and the cerebellar fastigial nucleus, also modulate breathing. Respiration is also under volitional control emanating from several cortical motor regions and operating via corticospinal pathways that are analogous to those of any other motor function.
Finally, it is important to emphasize the overwhelming influence of behavioral or sleep-wake states on breathing, including the ability during wakefulness to override autonomic respiratory reflexes, the relative insensitivity of such reflexes during rapid eye movement (REM) sleep, and the total dependence of gaseous chemical homeostasis on respiratory reflexes during non-REM sleep. A conceptual schema that shows the interrelationships among the many components of respiratory control is shown in Figure 2B.

Physiologic explanation of this patient’s respiratory status

Returning to Mr. MDC, his recent computed tomography scan shows that much of his medullary circumference and portions of the solitary and hypoglossal nuclei are invaded by dense connective tissue, presumably, at least in part, a result of irradiation treatment for his tumor. As central CO₂ chemoreception occurs at and beneath the ventral medullary surface and at additional medullary regions, including the solitary tract nucleus, it is not surprising that CO₂ homeostasis in this patient was very much affected. The biggest effect was seen during sleep, when chemical respiratory homeostasis is dependent on autonomic respiratory control. Following the three successive nights without ventilatory support, the disturbed CO₂ homeostasis was manifested by an arterial Pco₂ of 120-130 Torr. It was at that Pco₂ level that the patient was comatose due to CO₂ narcosis. During daytime hours in that time period, cortical and wakefulness-related influences partially override the abnormal autonomic CO₂ reflexes, thus reducing the arterial Pco₂ to 48-55 Torr. The alveolar hypoventilation that is the fundamental pathophysiological process in this patient was also accompanied by hypoxemia in room air.

With regard to the function of the 10th and 12th cranial medullary motor nuclei, it was interesting to note that the patient’s larynx dilated and constricted normally, indicating that, at least grossly, the motoneurons innervating the recurrent laryngeal nerves were spared. By contrast, his protruded tongue deviated to the right, indicating significant invasive fibrosis in his hypoglossal nucleus on the right. These discrepant findings attest to the variable degree of gliosis and fibrosis within the medullary tissue in this patient.

Additional examples of brainstem tissue lesions affecting chemosensitivity and breathing patterns

In their seminal 1981 review of neurological diseases manifesting as respiratory control abnormalities, Plum and Leigh summarized a variety of patients’ case histories, including
their pathophysiological respiratory symptoms and signs, and described specific respiratory tests that helped define the sites of the specific lesions. Armed with less sophisticated in vivo imaging tools than we have today, many case histories included specific respiratory tests that functionally defined the locus of the pathology, and that location was ultimately put to the test by autopsy. The frequency with which such diagnoses were correctly made attests to the advantage of using respiratory studies for the definition of pathological processes, and retains the merit of this review for neurologists to this day.

In this section, however, we shall focus on more recent literature, in which the localization of pathological processes was made in vivo using current imaging technologies. The first of such examples is interesting, because it shows that even small unilateral lesions caused by circumscribed infarcts in the rostrolateral medulla and identified by magnetic resonance imaging (MRI) can cause respiratory control abnormalities presenting as hypoxemia during sleep.\(^9,10\) Whether the hypoxemia was due to central hypoventilation or to upper airway obstruction could not be conclusively determined. Such upper airway obstruction might have been secondary to an incoordination between the upper airway and main respiratory muscles, or to a loss of function in the upper airways muscles. During restful wakefulness, minute ventilation and end-tidal \(\text{Pco}_2\) (reflecting arterial \(\text{Pco}_2\)) in those patients remained normal. By contrast, a \(\text{CO}_2\) challenge elicited reduced respiratory responses when compared to the response of patients whose infarcts were elsewhere in the brainstem. This diminished responsiveness could be due to either the loss of \(\text{CO}_2\) chemoreceptors, damage to a region that processes chemosensory input, or damage to a portion of the efferent pathway normally recruited only with hypercarbic stimulation. It is noteworthy, however, that these patients displayed a perfectly normal respiratory response to exercise. The dichotomous respiratory behavior in response to \(\text{CO}_2\) versus exercise demonstrated that, at the level of those brainstem lesions, the efferent corticospinal pathways for exercise hyperpnea were in a different location from the bulbo spinal pathways responsible for autonomic \(\text{CO}_2\) responsiveness.

The reduced \(\text{CO}_2\) responses with small unilateral lesions are important, in that they dispel the notion that a redundancy in \(\text{CO}_2\) chemoreceptive sites\(^7\) and a diffuse dispersion of bulbo spinal pathways\(^11\) guarantee a complete compensation for circumscribed damage. Furthermore, the studies of Morrell et al\(^9,10\) indicate that respiratory control abnormalities are not always easily apparent. When they do exist, such perturbations are likely, however, to manifest during sleep and to be expressed at least by an abnormally elevated carbon dioxide level and/or by reduced oxygenation. Therefore, at a minimum, regular monitoring of oxygenation during sleep serves as a precaution against such likelihood. Moreover, the diagnosis of breathing control abnormalities must at times be sought out deliberately with the help of specific respiratory tests such as \(\text{CO}_2\) challenge.

Respiratory control abnormalities, however subtle, have various clinical implications: (1) Cyanotic episodes, or sleep apnea/hypoventilation that are not explained by lung disease may reveal the existence of brainstem pathology; (2) Superimposed respiratory illness such as pneumonia can precipitate full-blown respiratory failure in vulnerable patients; (3) Sleep-related hypoventilation can be associated with exquisite sensitivity to the depressant effects of narcotic or sedative agents, leading to unexpected respiratory failure and occasional demise,\(^8\) and (4) The recurrent hypoxemia that is inherent to sleep-related hypoventilation carries a risk of long-lasting effects on many organ systems, for example, on the cardiovascular system, including systemic and pulmonary hypertension and cor pulmonale, and on the central nervous system, including an alteration of neuronal function and decreased neuronal survival.\(^12,13\)

Some of these implications are exemplified in a report on children with brain tumors who were referred to a sleep clinic for an evaluation of symptoms such as daytime hypersonnolence, respiratory insufficiency, hypoxemia and/or apnea.\(^14\) Of the 14 patients included in this report, one patient serves as an example for respiratory control abnormalities signaling brain malignancy: This patient had been initially referred for an evaluation of snoring and observed apnea, was found to have apnea and hypoventilation during sleep, but a CT scan of the brain at that time appeared normal. Two years later, when she returned with respiratory insufficiency and congestive heart failure, MRI study revealed a cervical-medullary astrocytoma that had probably also caused her initial respiratory symptoms. Another child who had a teratoid-rhabdoid tumor in the posterior fossa exemplifies the occurrence of overt respiratory abnormalities with acute lung pathology, in that this child developed both daytime and nighttime hypoxemia during a subsequent bout of pneumonia. A third child serves as an example for the increased sensitivity to sedation consequent to medullary pathology: Four years after resection and radio/chemotherapy for a medulloblastoma in the posterior fossa involving the brainstem, sedation for a colonoscopy that is usually uneventful precipitated an acute respiratory failure in that patient.

Medullary metastases can also result in a variety of respiratory effects depending on their location and/or extent of invasiveness. For example,\(^15\) a 52-year-old woman, who had been irradiated palliatively for large intracerebral adenocarcinomatous metastases (probably of pulmonary origin) displayed increasing hypersonnolence and was found dead one morning. Detailed autopsy revealed a hitherto undiscovered solitary metastasis in the brainstem. That small tumor completely replaced the right solitary tract nucleus and produced edema spreading to the contiguous reticular formation. Whereas the larger brain metastases in this patient may have also invaded supramedullary respiratory regions or pathways, the authors ascribed the sudden death to the disruption of activity in the solitary tract nucleus, because of its crucial role in the integration of afferent and efferent respiratory information, and because of its extensive connectivity both with its counterpart and with other respiratory medullary structures. The hypersonnolence in this patient has been attributed to the involvement of the reticular formation, and the time of death has been ascribed to the vulnerability of respiratory control mechanisms at the transition between sleep and wakefulness. Of interest, a much more extensive metastasis in the medulla in another patient\(^16\) produced completely different functional disturbances: This 67-year-old man suffered from hypcarbia and hypoxemia that were not attributable to an abnormality in respiratory rhythm, but rather to a perturbation in the motor execution of breathing activity.
Indeed, when challenged with CO₂ or hypoxia, this patient’s breathing frequency increased appropriately, but the tidal volume of each breath remained reduced. As well, when asked to take large breaths, his voluntary breathing efforts were also reduced. In this case, the authors concluded that the respiratory rhythm generator, the location of which is unknown in man, was somehow spared, thus permitting respiratory oscillation and increased respiratory rate, but that the major effluent respiratory outflow to the respiratory muscles was the culprit, thus preventing an increase in breath amplitude.

**Compression-related causes of brainstem respiratory control abnormalities**

Cervicomedullary compression in patients with achondroplasia can also cause severe respiratory symptoms, such as apnea and cyanotic episodes. The pathophysiology underlying those respiratory abnormalities can be divided into two categories: The first is directly related to skeletal abnormalities of upper airway structures, expected to produce upper airway obstruction, or of the chest, likely to produce restrictive lung pathology. The second pathophysiological category is a compression or indentation of the cervicomedullary brain tissue by osseous and fibrous elements. Such compression is sometimes combined with an obstruction of cerebrospinal fluid flow and with an elevated intracranial pressure.

When apnea and cyanosis were clinically observed in patients with achondroplasia, a polysomnographic sleep study revealed the full extent of the respiratory control abnormalities that included apnea and hypoventilation during sleep. Even in the absence of overt respiratory symptoms, however, sleep studies done in these patients for other reasons often revealed the existence of respiratory control abnormalities during sleep. Indeed, such respiratory abnormalities in achondroplasia are functionally more damaging than the skeletal motor disturbances associated with this syndrome, as these breathing perturbations may lead to acute respiratory failure and sudden death when they go undiagnosed. By contrast, however, when such perturbations are diagnosed in a timely fashion, and when they can clearly be attributed to the presence of a cervicomedullary compression, a surgical release of the compression can result in a complete relief of the respiratory symptoms. A concomitant upper airway obstruction, however, must be corrected separately.

Children with severe myelomeningocele can display similar sleep-disordered breathing manifestations to those in achondroplasia, although the underlying pathophysiology in myelomeningocele is much more complex. In a cross-section study of a clinic population of children with myelomeningocele, those children who had a motor or sensory myelomeningocele level above L3, combined with a severe, Grade 3 Chiari II malformation (comprising brainstem abnormalities including a cervicomedullary kink and compression) were most inclined to exhibit respiratory perturbations during sleep. Those disturbances included hypoventilation, primarily in REM sleep, increased apnea and hypopnea, and central as well as obstructive apnea. The obstructive apnea appeared to be largely due to an incoordination of upper airway muscles, also of central origin, rather than to the more common childhood-related causes such as enlarged adenoids and/or tonsils. Inasmuch as the central respiratory abnormalities caused by this congenital malformation last a lifetime, and because these abnormalities can be exacerbated by reduced lung function due to skeletal deformities such as scoliosis, the respiratory dysfunction in these patients must be diagnosed, treated and followed closely by medical teams that include neurologists.

**Congenital central hypoventilation syndrome**

Brainstem-related respiratory control abnormalities can also exist in the absence of any obvious structural pathology. In this regard, it is important to be cognizant of the existence of the relatively rare, but life-long affliction of congenital central hypoventilation syndrome. The predominant patho-
physiology of this syndrome is an inability to maintain chemical respiratory homeostasis, primarily during sleep in less severe cases, and during both sleep and wakefulness in more severe cases. Whereas the precise location or mechanism underlying this syndrome is yet unknown, the rhythmic but shallow breaths registered in these patients during sleep indicate an existence of a respiratory oscillator, but an abnormality in either the perception, integration, or motor neuron response to CO$_2$ and O$_2$ chemostimulation. The general location of this abnormality is thought to reside in the brainstem. This premise is strengthened by mounting evidence for other autonomic dysfunctions associated with this syndrome the location of which is also in the brainstem. The embryological origin of these abnormalities is thought to stem from the neural crest, and several genes have been associated with this syndrome, although none exclusively.

That there can be a genetic transmission of this syndrome is clearly evident from a study of a young woman who was afflicted with congenital central hypoventilation, as proven by a lack of responsiveness to CO$_2$ rebreathing (Figure 3A). She gave birth to an infant who, soon after birth, displayed wide swings in transcutaneous Pco$_2$ levels depending on behavioral state: Pco$_2$ was very high during sleep, and decreased to normal levels upon wakefulness (Figure 3B), as is typical of this syndrome. Despite its rareness (1 per 50,000 live births), one should be aware of this clinical entity so as to provide the life-long support that these patients require, including mechanical ventilation or phrenic nerve pacing.

In summary, the brainstem is crucial to the automated control of breathing and to chemical homeostasis. Experiments of nature, that is, tumors, fibrosis, infarcts, compression, distortion and structural abnormalities, as well as congenital conditions of as yet unclear etiology, can alter this homeostasis. Such disturbances are most clearly apparent during sleep, as this is the state in which the organism relies on automated respiratory control for the maintenance of chemical homeostasis.

**Sleep, breathing, and brain hypoxia**

The automated control of breathing by the brainstem is modulated by a great number of influences. One of the most important is the behavioral state. The importance of sleep-wake states in the control of breathing is already evident from the previous clinical examples which demonstrate that hypoventilation of any etiology manifests primarily during sleep, that wakefulness is a powerful stimulus to breathing that can overcome inherent brainstem pathology and normalize ventilation, and that the transition between sleep and wakefulness creates a vulnerability in the maintenance of breathing in compromised patients.

Sleep-wake states, the mechanisms of which are of ongoing interest and active research, modify breathing under normal conditions as well. Breathing control is truly automated during quiet, or non-rapid-eye-movement (non REM) sleep, a state associated with an inhibition of brain regions that produce arousal. During wakefulness, the afferent reticular activating
The principle behind the study is that a repetition of such episodes may produce functional consequences that outlast the removal of the obstruction. In children, such functional consequences are suspected, because post adenotonsillectomy improvements in school performance, for example, can be partial, modest, or non-existent, and because cardiovascular complications are sometimes only partially resolved.\(^{10,32}\) Neuroanatomical consequences are also suspected, as surmised from MRI studies in brains of adult patients after a long history of obstructive sleep apnea.\(^{12,13}\) These MRI studies have documented a decrease in the volume of grey matter in various brain regions. This volume loss has been interpreted as a possible indication of neuronal death consequent to the chronic intermittent hypoxia, hypercapnia and/or altered cerebral perfusion that accompany obstructive sleep apnea.\(^{12,13}\)

In addition to aggravating the hypoventilation associated with brainstem pathology discussed above, sleep states are linked to several specific respiratory-related syndromes. The most common of these is obstructive sleep apnea syndrome. Whereas in the adult population, obstructive sleep apnea can be found in both non REM and REM sleep, in the pediatric population, such apnea is closely linked to REM sleep. This link can be seen in the section of a polysomnographic study from a child referred to the sleep laboratory for an evaluation of severe snoring (Figure 4). In this section, REM sleep can be identified by a constellation of low voltage, high frequency electroencephalogram, presence of horizontal eye movements, and very low electromyographic chin activity (as contrasted with the high voltage, low frequency electroencephalogram, absence of eye movements, and moderate electromyographic chin activity associated with non REM sleep). At the same time, paradoxical breathing movements are present, but there is no air flow. Hence, oxygen saturation decreases and bradycardia follows suit. This is a typical episode of obstructive sleep apnea during REM sleep. This episode is followed by abrupt arousal and wakefulness, recognized from the constellation of low voltage high frequency electroencephalogram, increased electromyographic chin activity, and body movement, and by a resumption of vigorous consensual thoracic and abdominal breathing movements, airflow and normalized oxygenation and heart rate. It is evident that the arousal and wakefulness provide a powerful stimulus to breathing, thus interrupting the apneic episode. Such cycling can occur hundreds of times per night. In severely afflicted patients, this recurring breathing pattern can be diagnosed from overnight oximetry studies that show clusters of oxygen desaturation episodes, thus obviating the need for full polysomnography.\(^{25,26}\) Obstructive sleep apnea results in recurrent intermittent hypoxia, an increased threshold for arousal, sleep fragmentation and deprivation, increased daytime somnolence and confusion, decreased concentration and learning, and increased cardiovascular complications including hypertension and cardiac failure in both children and adults.\(^{27,28}\)

The pathophysiology of upper airway obstruction ranges from the relatively common enlargement of adenoids and/or tonsils or from more complicated craniofacial abnormalities in children to other structural upper airway narrowing and disturbances in upper airway neuromotor tone in adults.\(^{29}\) Whereas adenotonsillectomy in children, and even continuous positive airway pressure in adults, can reverse or improve much of the acute pathophysiology,\(^{30,31}\) there are indications that the recurrent intermittent hypoxia associated with prolonged, severe obstructive sleep apnea may have long term functional consequences that outlast the removal of the obstruction. In children, such functional consequences are suspected, because post adenotonsillectomy improvements in school performance, for example, can be partial, modest, or non-existent, and because cardiovascular complications are sometimes only partially resolved.\(^{10,32}\) Neuroanatomical consequences are also suspected, as surmised from MRI studies in brains of adult patients after a long history of obstructive sleep apnea.\(^{12,13}\) These MRI studies have documented a decrease in the volume of grey matter in various brain regions. This volume loss has been interpreted as a possible indication of neuronal death consequent to the chronic intermittent hypoxia, hypercapnia and/or altered cerebral perfusion that accompany obstructive sleep apnea.\(^{12,13}\)

The other close correlation between sleep and respiratory-related syndromes relates to the Sudden Infant Death Syndrome. Although the precise pathophysiology of this syndrome is yet to be determined, a leading hypothesis ascribes it to developmental abnormalities in the autonomic control of breathing patterns, chemoreflexes, cardiac reflexes, thermoregulation and/or asphyxic arousal that manifest in early infancy and during sleep, and that may be related to abnormalities in genes involved in the development of the autonomic nervous system.\(^{33}\) The subtle pathophysiological abnormalities are presumed to be augmented when the infants sleep in the prone position. In that position the infants may be rebreathing their own air, thus incurring progressive hypoxemia and hypercapnia. Whereas in normal infants, such circumstances would prompt arousal, a change in head position, increased ventilation, and correction of the respiratory chemical imbalance, these infants appear to have an elevated threshold of arousal and sluggish reflex responses. Such delay in arousal and responsiveness prevents these infants from benefitting from the powerful effect of wakefulness on breathing.\(^{19,33}\) A repetition of such episodes may produce recurrent hypoxemia which, in turn, may result in apoptosis in various brainstem and supramedullary regions\(^{34}\) that may further increase the vulnerability of these infants and set the stage for their demise.

**Supramedullary control of breathing: Volition**

In addition to the general excitatory effect on respiration provided by arousal and wakefulness, volition is a powerful stimulus of breathing that can modulate and override autonomic respiratory reflexes. It has been important to demonstrate where the drive for volitional respiratory control emanates from for both scientific reasons and for a better understanding of clinical situations in which this drive to breathe is disrupted, as will be described later.

The localization of supramedullary regions generating the volitional control of breathing has been based on a study employing positron emission tomography (PET) imaging in human volunteers.\(^{35}\) The principle behind the study is that a functionally-induced, increased synaptic and neuronal activity uses up oxygen and generates CO\(_2\), and that the resultant local reduction in oxygen and elevation in CO\(_2\) concentrations increase blood flow in the activated brain region. This regional blood flow increase can be imaged from H\(_2\)\(^{15}\)O that is infused intravenously at the time that this process takes place, and that is
preferentially distributed to the region of high blood flow.

Using this approach, the volunteers were asked to perform volitional inspiratory or expiratory maneuvers. The PET images thus obtained were collected three-dimensionally and subtracted from those obtained during passive breathing using positive pressure ventilation. Volitional inspiration was found to involve the primary and suplemental motor cortical regions bilaterally and the premotor cortex and ventrolateral thalamus unilaterally, confirmed subsequently. Of note, volitional expiration activated the same regions, although more extensively and with greater bilateral representation, and, in addition, activated the cerebellum. These findings document the participation of the cortex in volitional breathing, localize the motor regions involved, and emphasize the more extensive cortical representation during volitional expiration than during inspiration. Considering the fact that during normal breathing expiration is a passive process, the extensive participation of the cortex in volitional expiration emphasizes the essential role of active expiratory control in the execution of complex human activities such as speaking, singing, playing wind instruments, and expulsive maneuvers.

Exercise hyperpnea

During moderate exercise, the increased respiration is tightly matched to metabolic requirements and is termed exercise hyperpnea. The mechanisms involved in this hyperpnea include peripheral mechanisms through afferent signals from chemosensory organs, from the exercising skeletal muscle itself, and from the vascular dilation in the exercising muscle. Central mechanisms have also been invoked, especially at the onset of exercise, during which there is a central command to initiate an increased breathing activity in anticipation of the exercise. Using experimental animals, that central command had been thought to emanate from hypothalamic regions. More recent studies in humans have attempted to reveal whether the cortex might participate in the initiation of the central command for exercise hyperpnea.

To identify and define the role of the cortex in exercise hyperpnea, the study by Thornton et al involved hypnotizing volunteer adult subjects and, under hypnosis, asking them to imagine various modalities of exercise. During these imaginary exercise maneuvers, the investigators recorded the volunteers’ respiration and performed PET scanning so as to learn what portions of the brains were activated by each maneuver. The state of hypnosis was advantageous, because it isolated the subjects from environmental stimuli, diminished any perception of afferent stimulation, allowed a more directed and reproducible imagination of exercise without peripheral feedback, and ensured immobility, thus permitting more accurate scanning. Imagining riding a bicycle up a steep incline yielded a PET scan that included both the cognition of exercise and the imagined hard work of the exercise maneuver, whereas imagining riding a bicycle downhill rendered a scan that included the same cognition but without the imagined hard work. The volunteers were also asked to volitionally match the breathing patterns produced by the two exercise modalities, thus furnishing a PET scan of the volitional portion of those activities. Because increased ventilation without a concomitant increase in metabolism induces hypocarbia, CO₂ levels were monitored and maintained by the addition of CO₂ to the inspired air so as to avoid any confounding effect of hypocarbia on the observations. When the three-dimensional PET scans were subtracted from one another, the supplementary motor areas, the right premotor area, and the dorsolateral prefrontal cortex were identified as the cortical regions that participated in the exercise hyperpnea that was unencumbered by feedback mechanisms. These cortical regions identified by imaging in man have been proposed to initiate the behavioral response to exercise, thus dispelling the notion that the central command for exercise hyperpnea emanates solely from the hypothalamus.

Air hunger

One of the most unpleasant respiratory sensations that occur either during strenuous exercise or with pulmonary disease is dyspnea, or shortness of breath. One of its components is breathlessness, or air hunger, defined as an uncomfortable urge to breathe. This sensation has many descriptors and its intensity can be ranked and quantified. Air hunger can also be elicited in normal subjects by letting them inhale gases enriched with CO₂, by making them breathe through a face mask, or by limiting the size of mechanically provided breaths. Several studies have attempted to identify those brain regions that play a role in this sensation.

Brannan et al mapped such regions by PET scanning in healthy adults exposed to a high (8%) CO₂ mixture, and found that many limbic and paralimbic regions “lit up”, consistent with the classical notion that these afferent and “old” brain areas are associated with strong emotions, including fear, panic, and anticipation of pain. Many other brain regions were activated as well, however, including cortical areas and the cerebellum. Whereas the PET scans obtained by these investigators are informative, they probably represent a combination of a general increase of cerebral blood flow due to the direct vascular effect of the high CO₂ with the regional increases in cerebral blood flow due to the neuronal activation by the CO₂ stimulus. The latter, however, likely encompasses both the supramedullary regions recruited in the overall respiratory response to hypercarbia and those regions involved in the unpleasant sensations that accompany hypercarbia.

Several studies have attempted to distinguish among these components and isolate those brain regions activated by air hunger alone. In the first of this series, PET scans were obtained in healthy adult subjects that were artificially ventilated throughout the experiment, thus removing the contribution of volitional breathing from the equation. The experiment comprised changing the gaseous composition versus the volume of the mechanical breaths: When the Pco₂ was held at above normal levels, large tidal volumes did not elicit air hunger, but low tidal volumes did. When Pco₂ was held at below normal levels, neither large nor small tidal volumes elicited any sensation of air hunger. Thus it was a tidal volume that was inappropriately small for the prevailing chemical stimulus that generated the intense sensation of air hunger. Such a combination specifically activated the insular cortex, a region known to integrate visceral information with sensation, behavior and learning relevant to homeostasis. In this case, the extent of neuronal insular activation depended on the balance between the afferent stimulus to breathe (CO₂) and the afferent visceral.
information regarding the inflation status of the lung (mechanoreceptors) that, if adequate, would diminish the desire to breathe.

A subsequent study by the same group\textsuperscript{40} refined the experimental approach, and also used a different imaging technique, blood oxygenation level dependent functional magnetic resonance imaging (BOLD fMRI), to map the activated brain regions. Briefly, this imaging method relies on the relatively lesser degree of hemoglobin deoxygenation in blood distributed to activated brain regions, due to increased regional blood flow and volume in those regions. The altered ratio between oxygenated and deoxygenated hemoglobin modifies the magnetic output, registered as a change in BOLD fMRI.\textsuperscript{45} The advantages of this method versus PET scanning are (1) lack of ionizing radiation; (2) a greater signal to noise ratio, and (3) a greater spatial resolution. Using this method, and employing a similar experimental protocol in healthy adults to that used in the previous study,\textsuperscript{42} the investigators found that, in addition to the insula, many other cortical and subcortical regions were activated by the air hunger, including the anterior cingulate, operculum, cerebellum, amygdala, thalamus, basal ganglia and frontoparietal regions associated with attention. This finding indicates that the insula functions in concert with a larger neuronal network in integrating air hunger. More recently, the same group\textsuperscript{43} dispelled the previously held notion that only hypercarbia, but not hypoxia, provokes air hunger, by showing that when ventilatory drive is made to be equal, the two chemical stimuli result in the same degree of air hunger. It is likely that the afferent projection of both chemical stimuli to the insular cortex emanates from the solitary tract nucleus, as this nucleus integrates these chemical stimuli and has direct projections to the insula.\textsuperscript{44}

**Experiments of nature: Ondine’s curse and the ‘locked-in syndrome’**

The preceding sections addressed the roles of the brainstem and of supramedullary regions in normal physiological mechanisms controlling breathing. Ondine’s curse and the ‘locked-in syndrome’ are medical conditions that illustrate what happens when some of these mechanisms are severely disrupted.

Ondine’s curse is an extreme manifestation of sleep apnea, in which respiratory activity is entirely dependent on conscious volitional breathing, and ceases when the patient’s attention is diverted, upon the onset of sleep, or when anesthesia is administered. This syndrome is named after the German mythical sea nymph, Ondine, who thrust a curse on her unfaithful mortal knight, Hans, that he would lose all autonomic functions, and would have to remember to breathe voluntarily to survive. He died when he finally fell asleep. As reviewed recently by Bolton et al\textsuperscript{46}, this rare condition can be caused by an extensive destruction of the medullary ventrolateral respiratory regions and their reticulospinal efferent and afferent connections due to pathologies ranging from vascular events, large and extensive space occupying lesions, poliomyelitis, tract resection for intractable pain, or degenerative diseases. These patients require artificial ventilation or phrenic pacing to survive.

By contrast, in an illustrative case study of the ‘locked-in syndrome’,\textsuperscript{47} a 44-year-old man who suffered a bilateral infarction of the ventral pons displayed spontaneous breathing that was very regular, and that adequately maintained a normal Pco\textsubscript{2}. However, the patient could not exert any volitional breathing maneuver. By contrast, emotional outbursts provoked significant irregularities of his breathing rhythm. This patient is a testament to the fact that volitional breathing operates via corticobulbar pathways which were nonfunctional in his case. Further, these corticobulbar pathways are distinct from limbic respiratory pathways that conduct supramedullary respiratory responses to emotion, and that did remain functional. Under normal conditions, the two pathways eventually merge, but in his case, only the limbic pathways were intact. It is also noteworthy that this patient survived due to his pontomedullary respiratory oscillator which operated in isolation from the corticospinal pathways. It is because of this autonomic oscillator that this patient’s ventilatory response to inhaled CO\textsubscript{2} was normal, and that he could sense breathlessness during the hypercarbic stimulus. Another instructive lesson was derived from the observation that when the patient’s Pco\textsubscript{2} level was artificially reduced by mechanical hyperventilation while he was awake, he became apneic. In healthy subjects, such apnea never happens during wakefulness, and can only occur when the subject is in non REM sleep, as the wakeful stimulus that provides cortical excitation overrides such apnea and sustains breathing via the corticospinal pathways. The disrupted corticospinal pathways in this patient obviated such cortical protection against hypocarbia. The lessons that can be learned from this syndrome is that, whereas the pontomedullary oscillator is essential for basal respiratory homeostasis, the brain as a whole is required for normal breathing, and that the wakefulness stimulus and intact corticospinal pathways are crucial for maintaining stable, versatile and appropriately adaptive breathing patterns.

**Breathing and Emotion**

A vast lore exists in the medical literature about the effects of emotion on breathing.\textsuperscript{48} The best known example is the emotive hyperventilation syndrome that can result in extreme hypocarbia, respiratory alkalosis, and muscle spasms due to an
acute reduction in ionized calcium. The relationship between emotion and breathing is an everyday experience, however, as evidenced from Figure 5. This figure depicts respiratory and pulse signals of a person listening to two successive styles of music, dissonant atonal electronic music by Stockhausen versus melodic soothing music by Chopin. The difference in respiratory and pulse patterns between the two is striking: It is highly excited and irregular in the first, and very stable and regular in the second, showing the different emotionally-driven respiratory and cardiac reactions to the two types of music. The pathways governing the emotion to respiration operate via limbic structures and are anatomically distinct from the corticospinal pathways, as noted in the ‘locked in syndrome’ case study discussed above.

SUMMARY AND CONCLUSIONS

Breathing is not only an automated function, but is a complex behavior that involves many brain regions. These regions are cortical and subcortical regions that are not usually considered with regard to respiratory control. These regions are activated by respiratory-related sensations and initiate complex respiratory activities. Clever experimental designs and specialized respiratory functional tests are required for demonstrating respiratory manifestations of brain pathology. Since respiratory abnormalities often become apparent during sleep, monitoring of breathing during sleep should be an inherent practice for any neurologically related pathology. Such monitoring will not only help in identifying abnormal patterns and chemical imbalances when the pathology is known, but will sometimes reveal an unknown pathology, and will also prevent respiratory crises from occurring during sleep or at the transition between sleep and wakeful states.

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