NORMAL RESPIRATION

Respiration requires an integrated system of control (CNS), movement of a bellows (ribs and associated muscle) to draw air into the lungs, exchange of gas within the lung, and a feedback system (chemoreceptors, stretch receptors) to close the loop. This lecture will present some basic elements of the bellows part of this system, providing a basis for evaluation of respiratory patterns in the dog.

LUNGS COLLAPSE, RIBS SPRING OUT

Lungs are a viscoelastic tissue suspended within the pleural cavity. If the chest is opened to the atmosphere the lungs collapse until small airway occlusion prevents further loss of gas. The collapse won’t stop until the lung volume is quite a bit smaller than the normal functional residual capacity (FRC) in the intact animal. The lung volume in a normal animal is maintained by support from the thoracic wall. Because there is an airtight seal within the pleural space and the lungs are coupled to the chest wall with a viscous fluid, the tendency of the lungs to collapse is countered by the tendency of the ribs to spring out. Between breaths, these opposing forces yield a pleural pressure of -5 cm H\(_2\)O at the 8th-9th interspace in an average size dog. Breathing takes place around this equilibrium, with most of the energy of breathing expended to increase the thoracic volume above the equilibrium point during inspiration.

BUCKET HANDLES

The ribs in the dog (and cat) are arranged in such a manner that most movement, and the inspiratory effect of that movement, is related to cranial displacement of the ribs during inspiration (Fig 1). Because of the ‘bucket handle’ relationship between the ribs and the spine and the caudal sweep of the ribs when the lungs are at functional residual capacity (FRC), cranial displacement of the ribs results in an increase in thoracic diameter and intrathoracic volume, a reduction in pleural pressure, and inspiration.(1) Therefore, any muscles that move the ribs forward and outward will contribute to inspiration, and any that move the ribs caudally and/or inward will contribute to exhalation. Until recently, the usual explanation for intercostal muscle function was based on the theory of Hamberger (1697-1755). He proposed that the external intercostals, because they were anchored high on the cranial rib (closer to its axis of rotation) and lower on the caudal rib (further away) at each intercostal space, must have a net effect of raising the caudal rib, producing inspiration. Similarly, the internal intercostals, with their origins high on the caudal rib and insertions lower on the cranial rib at each space, must have a net effect of pulling the cranial rib more caudally, producing exhalation. This served, without experimental verification, as the leading explanation for intercostal function until the 1980’s. Since then major advances in understanding have been obtained by the efforts of a small number of physiologist using the dog as a model. From this work comes an appreciation that the act of breathing is very complex, and the effects of muscle activation depend on many factors including location (dorsoventral and craniocaudal), muscle mass, patterns of innervation, and locomotion.(2)

RESPIRATORY MUSCLE AND ‘ABDOMINAL BREATHING’

The primary muscles of inspiration include the diaphragm, levator costae, internal intercostals (ventro-cranial), and external intercostals (dorsocranial). The primary muscles of exhalation include the triangularis sterni and dorsocaudal internal intercostals. Accessory muscles of respiration are those that do not normally participate in breathing unless called upon during periods of high demand. Accessory muscles of inspiration include the scalenes and the sternomastoids. Accessory muscles of exhalation include the four abdominal muscle groups: rectus abdominis, external oblique, internal oblique, and transversus abdominis. Accessory muscles of respiration have other primary roles such as movement of the head and neck (sternomastoids), stabilization of the spine (scalene), flexors and rotators of the trunk (rectus and obliques), and their function is synchronized with breathing when conditions demand. Contraction of the abdominal muscles increases intra-abdominal pressure and forces the diaphragm to a more cranial position within the rib cage. They can ONLY assist with exhalation, and the term “abdominal breathing” is therefore properly used to indicate vigorous exhalation.
As depicted in figures 2 and 3, the external and internal intercostals in the cranial intercostal spaces have an inspiratory bias, and the same muscles in the caudal intercostal spaces have an expiratory bias.

Figure 2. Inspiratory actions of the internal intercostals (left) and external intercostals (right). The shaded areas represent locations where the mechanical advantage and net respiratory effects favor inspiration at FRC. The portion of the internal intercostals between the bony rib elements of the first few spaces remain electrically silent during breathing, and the external intercostals in this region, when active, are active only during inspiration.

Figure 3. Expiratory actions of the internal intercostals (left) and external intercostal (right). The shaded areas represent locations where the mechanical advantage and net respiratory effects favor exhalation at FRC. This region of the internal intercostals, when active, is active only during expiration. This region of external intercostals is electrically silent during tidal breathing.

The muscle mass of the external intercostals is greatest in the dorsocranial region where they possess the greatest mechanical advantage for inspiration. The parasternal internal intercostals are a special group of internal intercostal muscles that play a greater role than the externals in inspiration during rest. These muscles also have a large mass and they possess a mechanical advantage for inspiration in every intercostal space. Their orientation is such that contraction produces outward movement of the distal rib relative to the sternum. They have a comparatively large effect on inspiration because they move the chondral aspect of the ribs outward, away from the midline. In the dog any given movement in the outward direction is roughly 4 times more effective at increasing lung volume than comparable movement in the cranial direction.

The levator costae and parasternal and external intercostals are active during inspiration. Levator costae originate on the transverse process of the thoracic vertebrae and extend caudally to insert on the dorsal-cranial aspect of the rib so when they contract they rotate the ribs cranially. This muscle works in concert with the parasternal intercostals to provide most of the rostral rib displacement during inspiration. The levator costae and parasternal intercostals appear to function exclusively for breathing. In contrast, the function of the interosseous portions of the intercostal muscles may be given over to stabilization of the torso during activity. For example, in trotting dogs the activity of the intercostal muscles becomes synchronized with locomotion and drifts relative to the phase of respiration. Therefore, in active dogs the interosseous portions of the intercostal muscles are more important for stabilizing the trunk for locomotion than for any direct role in breathing. However, their action stabilizes the rib cage, which is essential for optimal function of the other respiratory muscles.

**EXHALATION: NEVER PASSIVE IN DOGS**

The triangularis sterni and internal intercostals are active during expiration. In particular, the triangularis sterni muscle is active even during quiet breathing, and is independent of both body position and intercostal activity. Figure 4 illustrates the orientation of this muscle from a view inside the thorax. The muscle extends from the caudal half of the deep aspect of the sternum to the chondral (parasternal) portion of ribs 2-7. When it contracts, the triangularis sterni displaces the ribs caudally and the sternum cranially, positioning the ribs for a greater mechanical advantage and lengthening the parasternal intercostal muscles to enhance their function during the next breath.
DIAPHRAGM

The diaphragm is now considered to be a combination of two functionally discrete muscles, the costal and crural diaphragm. The muscles separate the abdominal cavity from the thoracic, and diaphragm function is essential for effective ventilation. Contraction of the diaphragm in inspiration causes it to flatten and move the liver and abdominal viscera caudally into the abdominal cavity. However, because it is anchored to the chondral arch and caudal ribs, contraction of the diaphragm pulls this region of the rib cage rostrally. When working in concert with the parasternal intercostals and levator muscles the diaphragm assists in the cranioventral rib displacement of inspiration to increase thoracic girth and volume. The effect on airway pressure of this combined effort is significantly greater than the sum of the individual contributions from diaphragm and rib muscles.(6)

DON'T CONFUSE FAST BREATHING WITH PANTING

Panting is a thermoregulatory maneuver and is the most important method of evaporative cooling in dogs exposed to heat or exercise. It has minimal impact on gas exchange. When the skin (environmental trigger) or core temperature (exercise) temperature rise, respiration shifts to a pattern optimized to remove heat from the body. The diaphragm and other muscles of respiration generate a rhythmic motion that cycles at a frequency of 3.5-5 Hz. The precise frequency is closely related to the resonant frequency for the individual, thus body conformation and the unique characteristics of that individual’s respiratory system determine the rate. The oropharynx and larynx move in synchrony to 'valve' the system and provide an efficient unidirectional flow of air over the evaporative surface of the mouth.(7) When the need is minimal, dogs will pant through the nose. As need for heat loss increases, dogs oscillate between two patterns: a) inhalation through the nose, exhalation through the nose and mouth and b) inhalation through the nose and mouth, exhalation through the nose and mouth. Lingual blood flow increases 6 fold, and the tongue, wet from increased salivation, hangs out of the mouth as it lengthens to increase its surface area.(8)

Panting requires a dramatic increase in minute ventilation, but blood gases do not change appreciably (although in severe heat stress panting will cause a significant respiratory alkalosis for a short period). This is possible because most of the increase in ventilation that occurs during panting is dead space ventilation, and the inspiratory volume is only slightly larger than the anatomical dead space of the airways. Ventilation is maintained owing to the principles of jet ventilation: individual molecules of gas enter the trachea at very high speeds. Mixing of gas in distal airways is enhanced by asynchronous contractions of the crural and costal segments of the diaphragm, and there is net displacement of carbon dioxide out and oxygen in.(9)

HOW SICK DOGS BREATHE

It is suspected that mammals generally adopt a respiratory pattern that meets their metabolic needs with the least metabolic energy cost. Because of the phenotypic diversity in healthy dogs and cats, how this is achieved varies dramatically between patients - what works best for a Borzoi may not work at all for a Pug. In addition to these species and breed differences, respiratory diseases cause characteristic changes in respiratory frequency, tidal volume, airflow velocity, and patterns of muscle activation that can provide clues for the clinician to help localize the cause of abnormal respiratory function. This lecture will present the basic elements of how respiratory patterns change with disease states and emphasize important
inspiration and exhalation. The intercostal muscles ‘suck in’ from markedly negative pleural pressures created by vigorous inspiratory phase becomes noticeably prolonged. As the problem becomes more severe, more energy is expended on both a prolonged inspiration and exhalation that is characteristic of fixed obstructions. Respiratory frequency falls as the change appreciably with the phase of respiration. As the airway lumen narrows with advancing disease, the animal will adopt approaches dyspnea it must focus more and more conscious effort on the act of breathing. Animals that are interactive and engaged with their environment usually still have significant physiological reserves; those that appear withdrawn and focused on the act of breathing may be close to the edge of respiratory arrest.

Where a given patient lies on this spectrum must be inferred from their behavior. In general, as an animal approaches dyspnea it must focus more and more conscious effort on the act of breathing. Animals that are interactive and engaged with their environment usually still have significant physiological reserves; those that appear withdrawn and focused on the act of breathing may be close to the edge of respiratory arrest. A notable exception to this generalization is the puppy. Juvenile dogs will often remain active, alert, and hungry even with advanced respiratory failure and may go straight to the brink of death before showing compelling behavioral signs of distress. Dyspnea is a true medical emergency; our goal is to relieve it within minutes of presentation. In contrast, the terms associated with increased resistance to airflow but may also be caused by anything leading to clinically significant hypoxemia or hypercapnia (especially if acute). Like pain, its presence is inferred from facial expressions and behavioral signs of distress. Dyspnea is a true medical emergency; our goal is to relieve it within minutes of presentation. In contrast, the terms referred to the physical manifestations of increased work of breathing. This may be objectively measured, but most of the time is inferred from subjective appraisal of physical signs. Whereas an animal with dyspnea is having a crisis, an animal with labored breathing may be very well adapted to its condition and be free from distress.

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**DYSPNEA VS INCREASED RESPIRATORY EFFORT**

The term dyspnea refers to the experience of distress secondary to respiratory disease. In man it is most often associated with increased resistance to airflow but may also be caused by anything leading to clinically significant hypoxemia or hypercapnia (especially if acute). Like pain, its presence is inferred from facial expressions and behavioral signs of distress. Dyspnea is a true medical emergency; our goal is to relieve it within minutes of presentation. In contrast, the terms referred to the physical manifestations of increased work of breathing. This may be objectively measured, but most of the time is inferred from subjective appraisal of physical signs. Whereas an animal with dyspnea is having a crisis, an animal with labored breathing may be very well adapted to its condition and be free from distress.

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**FIXED AIRWAY OBSTRUCTION**

**Neoplasia, constrictions, compression from abscesses or trauma:** A ‘fixed’ obstruction refers to one that does not change appreciably with the phase of respiration. As the airway lumen narrows with advancing disease, the animal will adopt a prolonged inspiration and exhalation that is characteristic of fixed obstructions. Respiratory frequency falls as the inspiratory phase becomes noticeably prolonged. As the problem becomes more severe, more energy is expended on both inspiration and exhalation. The intercostal muscles ‘suck in’ from markedly negative pleural pressures created by vigorous diaphragmatic contraction, and the animal may actively exhale, first by contracting the intercostal muscles alone, later by a combination of intercostals and abdominal muscles. The term ‘abdominal breathing’, referring to active contraction of the abdominal wall, can only assist with exhalation and is a response to air trapping that slows exhalation to unacceptable rates.

Neoplasia and constrictions from scarring are typically insidious in onset, initially causing clinical signs only with exercise and usually allowing plenty of time for adaptive strengthening of the respiratory muscles. If acute, as for example from cervical swelling from trauma, the animal may have the same physical features as aspiration of a foreign body, including dyspnea and cyanosis.

**Foreign body:** This disorder is characterized by an acute onset of respiratory difficulty, often accompanied by coughing/retching and behavioral signs of dyspnea. Marked cyanosis may be present as hypoxemia occurs before there is any adaptive shift in hemoglobin oxygen affinity and cardiovascular responses are vigorous, with good perfusion of skin and mucus membranes. In the author’s experience, inspiration is more severely affected than expiration but this would be expected to vary with location of the obstruction. Most obstructions are at the level of the larynx or tracheal bifurcation.

**Brachycephalic syndrome:** This syndrome is classically characterized as restriction of the upper airway secondary to excessive or extreme phenotypic expression of the abnormalities selectively bred for in some chondrodysplastic breeds. Although some anatomic features of the syndrome produce a dynamic obstruction, most individuals have at least some component of fixed obstruction. Any combination of stenotic nares, stenotic nasal cavity, elongated soft palate, myopathy of
the pharyngeal opening musculature, laryngeal dysplasia, everted laryngeal sacculles, and trachea hypoplasia are possible.(10-12) Collapse of the cervical and/or intrathoracic portion of the trachea is sometimes seen. Some dogs have concurrent abnormalities of respiratory drive(13), and some acquire rib fractures and/or hiatal hernia.(14) The breathing pattern is characterized by prolonged inspiration, and (usually) comparatively easy exhalation.(15) Affected dogs routinely develop stertorous sounds with the slightest provocation. In spite of evidently high resistance to air flow, affected animals are generally not focused on breathing and relatively free from distress, even if their arterial pO2 is very low. This finding fits with the observation in many species that chronic hypercapnia and hypoxemia invoke adaptive responses that resets chemoreceptor response thresholds to relatively high values of pCO2 and low values of pO2.

DYNAMIC AIRWAY OBSTRUCTION

Laryngeal paralysis: Although some dogs with idiopathic laryngeal paralysis have features of a fixed airway obstruction, this is probably the best example of a dynamic extrathoracic obstruction.(16) In this disorder, deterioration of the recurrent laryngeal nerves results in a loss of innervation of the abductor muscles of the larynx. It may occur as an isolated neuromuscular abnormality or be part of a polyneuropathy that may include swallowing disorders and/or megaesophagus. Most often, it is insidious in onset and many owners may recognize that clinical signs have been present for months only in retrospect after a crisis. During inspiration, the larynx is prone to collapsing into the lumen secondary to the subatmospheric pressure that creates a Venturi effect as described in Bernoulli’s principle. Vigorous inspiration becomes prolonged and either stertorous (if the laryngeal structures vibrate) or attended by a high pitched squeak (if obstruction is nearly complete). Many dogs seem to be able to prevent this for long periods by adopting an inspiratory pattern that is slow in frequency with a relatively prolonged inspiration and large tidal volume. They can thus avoid symptomatic obstruction unless forced to increase ventilation by the demands of activity or environment (by inducing panting from heat exposure). When signs occur in these dogs they often come on dramatically as the dog worsens the obstruction with rapidly escalating inspiratory efforts that in turn worsen the obstruction. Many present with cyanosis (from acute worsening of the obstruction) and hyperthermia (from the heavy muscular effort).

Tracheal collapse, bronchial collapse: These terms describe the obstruction of a large airway due to chondromalacia, compression, or in the case of the trachea, a redundant dorsal ligament. Clinical signs will depend upon which portion of the airway system is affected. Extra-thoracic tracheal collapse will tend to occur during inspiration when intraluminal pressure is subatmospheric and the dorsal ligament is sucked into the tracheal lumen. Intrathoracic collapse is more likely to occur during exhalation when pleural pressures approach (or exceed) intraluminal airway pressure. Either event may stimulate coughing via stimulation of submucosal nociceptors, but intrathoracic collapse usually causes more severe problems than cervical tracheal collapse. Animals with cervical tracheal collapse are often asymptomatic except with activity; once they begin collapsing the airway they may quickly transition to a pattern characterized by vigorous attempts at inspiration with loud stertor or sounds of “choking off.” Animals with intrathoracic airway collapse may have no signs other than persistent cough. If severe, they may begin trapping air at relatively high lung volumes and be forced to recruit accessory abdominal muscles of expiration in an effort to forcibly ‘burp’ air past obstructed airways.

Feline asthma: If bronchospasm is severe, this syndrome has all the features of a fixed airway obstruction. In most patients, however, it is likely that restriction to airflow is worse on exhalation as small airway diameter falls with declining lung volume. Wheezes are commonly auscultated, but auscultatory evidence of large airway collapse is typically lacking.

RESTRICTIVE LUNG DISEASE

Restrictive diseases are characterized by pulmonary changes that reduce lung compliance and make the lungs more difficult to inflate. At the same time, the underlying disorder typically compromises the oxygenating capacity of the lung. Arterial hypoxemia and lung inflammation or edema are direct respiratory stimulants and provoke an increase in desired minute alveolar ventilation. Thus, when severe, the animal is faced with the need to increase ventilation with lungs that are increasingly stiff and difficult to inflate. The archetypical pattern that results from this combination is breathing at a relatively fast rate and small tidal volume.

Clinically, the animal develops an increased frequency with rapid shallow inspirations. As inspiration becomes more difficult (or fatigue develops with acute illness), one of the first behavioral changes is a reluctance to lie in lateral recumbency. As difficulties progress, the dog may have trouble remaining in sternal recumbency for prolonged periods and position changes become more frequent. Finally it may be difficult to lie down for any significant period and the dog will prefer to sit or stand, even in the face of exhaustion. During this progression, the dog begins to extend the head and neck and recruit accessory muscles of inspiration, including the scalenes and the sternomastoids(17;18). The forelimbs are abducted as breathing difficulty becomes greater. The head may bob up and down with respiratory effort. If the elastic recoil of the lung is sufficient, exhalation may be completely passive. If there is excessive air trapping from airway collapse, or if passive recoil of the lung does not provide sufficient emptying the dog may exhale actively (contraction of intercostal muscles) and may recruit accessory muscles of expiration (abdominal muscles).

In contrast to what is seen with airway obstruction or pleural space disease, hypoxemia from lung disease typically causes more pallor than cyanosis, yielding a pale-to-grey color of the mucus membranes. If the underlying cause is rapidly progressing the animal may develop fatigue of the respiratory muscles quickly and go on to develop hypoventilation (respiratory failure).

Pulmonary edema/pneumonia: Any increase in the water content of the lung tissue will reduce its compliance by
increasing the opening pressure required to inflate pulmonary exchange units. As edema develops, regions of the lung will lose ventilation and venous admixture increases. Respiratory frequency increases, in some cases associated with stimulation of pulmonary C fibers (juxtacapillary receptors) from the increased fluid pressure of the interstitium, and in some from hypoxic stimulation of chemoreceptors. Because the lungs are relatively stiff, the prototypical pattern will be relatively shallow breaths at a relatively high frequency. As these disorders tend to develop rapidly, respiratory fatigue may develop quickly and hasten death in the absence of ventilatory support.

Idiopathic pulmonary fibrosis: This progressive disorder of cats and dogs results in loss of function (hypoxemia) and compliance, and is may be accompanied by cough. Compared to pulmonary edema, this disorder progresses relatively slowly and there is ample time for training of the respiratory muscles and development of fatigue resistance. Affected animals are often not in respiratory distress until functional loss is severe.

**PLEURAL SPACE DISEASE**

The thoracic girth is normally maintained by the opposing forces of lung (which wants to collapse) and rib cage (which wants to spring outward). Separation of the two with a layer of air or fluid in the pleural space produces a reduction in lung volume and an increase in thoracic girth. The effects of pneumothorax on breathing has been studied more than the effects of pleural effusion, but it appears that dynamic lung compliance, inspiratory duration and tidal volume decrease significantly with either disorder.\(^\text{(19,20)}\) As lung volume falls exchange units are lost as the reduction in volume becomes sufficient to favor small airway collapse, leading to hypoxemia due to venous admixture and if severe, hypoventilation. At the same time this is occurring the total intrathoracic volume is growing, flattening the diaphragm (placing it at a mechanical disadvantage) and shifting external intercostal muscle function towards exhalation.

As the disorder progresses, inspiratory efforts increase and accessory muscles of inspiration may be recruited. The diaphragm will contract maximally, maintaining good excursion in spite of its flattened profile in exhalation. Pneumothorax directly stimulates phrenic nerve activity in a manner independent of the effects on gas exchange.\(^\text{(19)}\)

**DIAPHRAGMATIC WEAKNESS OR PARALYSIS**

Diaphragm paralysis may be caused by injury to the phrenic nerve(s), degeneration of the phrenic nerves, or severe motor unit diseases such as botulism. If the condition is isolated to phrenic nerve involvement, dogs compensate by increasing the activity of the intercostal, levator costae and parasternal muscles. Although this may be a conscious effort in response to chemical stimulation, contraction of the normal diaphragm stimulates receptors whose output inhibits efferent activity to the intercostal and levator costae muscles. Clinically, the dog has an increased frequency, marked inspiratory excursion of the anterior ½ of the thorax, and sometimes paradoxical movement of the abdomen. The latter is characterized by a failure of the abdomen to maintain its girth during inspiration or a frank reduction of girth if the diaphragm and abdominal viscera are sucked into the thorax by the vigorous inspiratory efforts happening up front.

**INTERCOSTAL PARALYSIS**

Weakness or paralysis of the intercostal muscles may be seen as part of a generalized myopathy, motor unit disease, and spinal cord injury due to trauma or neoplasia. In this patient, the thoracic wall may fail to maintain its normal position during tidal breathing and move paradoxically during diaphragm contraction. Because these dogs are typically paralyzed, if their lungs are otherwise normal they usually have adequate respiratory function with diaphragm movement alone.

**DISORDERS OF THE BONY STRUCTURES FORMING THE THORACIC BELLOWS**

Several disorders affect the bellows mechanism of respiration, including rib fractures and breakdown of the chondral arch. Rib fractures are painful and may limit tidal volume, prompting a compensatory increase in respiratory frequency. In addition, fractured ribs disrupt the action of the intercostal muscles and may produce paradoxical motions of regions of chest wall that further compromise ventilation.

Some animals with stiff lungs or chronic airway disease will eventually suffer collapse of the costochondral arch, possibly secondary to excessive forces generated by a hypertrophied diaphragm. When this happens, the diaphragm is shortened, losing its dome shape and most of its mechanical advantage. When viewed with fluoroscopy, contraction of the diaphragm results in pulling the caudal aspect of the sternum closer to the spine, instead of forcing the caudal ribs to move in a cranial direction and increasing thoracic volume. Because these animals typically have chronic lung or airway disease and already suffer significant functional loss, their condition is often made acutely worse when the contribution of the diaphragm to inspiration is compromised.

**Reference List**