If you are coming here because your dog has just been diagnosed with Megaesophagus, you are probably asking yourself, *why is this happening to my dog?*

**Canine Megaesophagus**

Let’s review what Canine Megaesophagus is all about first. Megaesophagus is a disorder of the esophagus. The esophagus is the tube which connects the mouth to the stomach. With megaesophagus, the esophagus is enlarged (dilated) and there is a decrease in peristalsis. Peristalsis is the wave like contractions that moves the food from the esophagus to the stomach. ME dogs have a dilated esophagus due to lack of muscle tone. This causes food to sit in the esophagus, unable to travel to the stomach and sometimes forming pockets. When this happens a dog “regurgitates” the food that is stuck in the esophagus. The dog will open his mouth and food will simply roll out. It is generally a quiet and effortless motion opposed to vomiting. Vomiting is an active motion associated with retching and heaving digested food.

Megaesophagus may be **congenital** (born with it) or **acquired**. Acquired ME can be further divided into two subclasses, **idiopathic** (cause unknown) or it can be **secondary** to another disease. Unless your dog is born with ME, it is very important to look at what could have caused the ME. ME is secondary to many diseases, diseases that can be treated. Many times the symptom of Megaesophagus can resolve once the proper drugs are in place. Let’s take a look...
Congenital ME

Puppies with congenital ME usually show signs when they are being weaned from their mother’s milk. While milk can slide down easily, solid food tends to bulk up and the puppy will regurgitate their food. Sometimes it is noticeable right from the start. Breeders often refer to these puppies as “milk nose”. They are often the runt of the litter. It is largely the result of “esophageal hypomotility” (decrease or absence of movement or peristalsis). Sometimes esophageal hypomotility is due to delayed maturity of the esophageal function that may or may not improve with age. This is why it is so important to diagnose and start treatment early in order to prevent pockets from forming and further dilation of the esophagus. Thoracic radiography (xray) is the common diagnostic tool to determine megaesophagus. A barium contrast esophagram (barium swallow) can confirm the presence of PRAA (Persistent Right Aortic Arch).
Persistent Right Aortic Arch (vascular ring anomaly) is a birth defect that happens when blood vessels, known as aortic arches in the heart, fail to deteriorate as the embryo grows. The right aortic arch passes near the esophagus. If this blood vessel does not dissipate in the newborn, the esophagus is trapped between it and the heart. Food is restricted from passing by this stricture (narrowing), causing regurgitation. Surgery is recommended to correct this condition. It should be performed as early as possible to prevent further damage to the esophagus. Many puppies born with PRAA do not have true megaesophagus. Once the stricture is removed these puppies can eat normally in the majority of cases. Please click on our PRAA tab for further information about the surgery.

Congenital Myasthenia Gravis
Congenital MG is an inherited autosomal recessive condition (meaning two copies of an abnormal gene must be present in order for the disease or trait to develop). It is found in Parson Russell Terriers, Springer Spaniels, and Smooth Fox Terriers. They are born without the acetylcholine receptors (AChRs) at the neuromuscular junction. Puppies that are born with congenital MG present similar to acquired MG with general muscle weakness to the striated skeletal muscles and megaesophagus. Measurement of acetylcholine receptor antibody titer is not a diagnostic tool for congenital MG. A muscle biopsy is needed to diagnose this condition. The outcome for congenital myasthenia gravis is not good since traditional treatment for MG is not effective. Read more by clicking the tab for myasthenia gravis from the home page.

Acquired Megaesophagus
Secondary acquired megaesophagus can be brought on by any disease that interferes with the esophageal peristalsis (movement) by disrupting central, efferent, or afferent nerve pathways or by any disease of the esophagus muscle system, including immune-mediated, infectious, and preneoplastic (pre stages of a tumor) causes. The wide spread causes of acquired ME can be the result of a variety of neuromuscular, immune-mediated, endocrine, gastrointestinal, paraneoplastic, and toxic diseases.
**Neuromuscular and Immune-Mediated Causes**

**Acquired Myasthenia Gravis**
This disease is said to be the cause of 25% or more of canine megaesophagus. Early diagnosis and treatment is the key to success with this disease. Since 90% of dogs with MG have ME also as a symptom, controlling the ME is most important. Aspiration pneumonia continues to be the biggest threat here. Read more about MG in the tab section from our home page. Important to remember: Dogs with MG can go into remission on average in 6-8 months and sometimes their ME can resolve as well. ALL DOGS THAT ARE NOT BORN WITH ME AND ARE DIAGNOSED WITH ME SHOULD BE TESTED FOR MG. ALL DOGS THAT ARE DIAGNOSED WITH MG SHOULD BE TESTED FOR ME!

Possible Neuromuscular and Immune-Mediated causes include:

- Botulism
- Canine Distemper
- Dermatomyositis
- Dysautonomia
- Glycogen storage disease (GSD)
- Laryngeal paralysis-polyneuropathy complex (LP-PNC)
- Myasthenia Gravis
- Neosporosis
- Polymyositis
- Polyneuropathy
- Tetanus
Endocrine Causes

Hypoadrenocorticism (Addison’s Disease) and Hypothyroidism

These are two diseases that are associated with reversible megaesophagus. And a good reason to test!

Dogs with Addison’s disease may have megaesophagus due to electrolyte imbalances and a cortisol deficiency. Electrolyte imbalances cause altered membrane potentials, which result in decreased neuromuscular functions. Muscle weakness is a symptom of deficient cortisol.

Addison’s Disease

Addison’s Disease is the common name for hypoadrenocorticism, or adrenal insufficiency. There are three forms of Addison's disease: primary, secondary and atypical. Primary and atypical Addison's are usually the result of immune mediated damage to the glands. Secondary hypoadrenocorticism is from failure of the pituitary to stimulate the adrenals with adrenocorticotropic hormone (ACTH). It is important for you to know which type of Addison's disease your dog is being treated for.

Symptoms of Addison disease may include weakness, depression, loss of appetite, vomiting, diarrhea, and sometimes increased thirst and urination and megaesophagus. The confirming test for Addison's disease is an ACTH stimulation test. In this test blood samples are taken before and after the dog is injected with an adrenal-stimulating hormone. If your dog has primary Addison’s, which means both layers of the adrenal gland are under-producing hormones, fludrocortisone is usually prescribed. There is also an injectable drug called DOCP, brand name Percorten V. This is a long-acting drug given by injection every 25 days. The drug is given primarily to regulate electrolytes, so cortisol replacement will also be needed. This means both an injection and an oral daily cortisone drug. For dogs with atypical or secondary Addison's, only glucocorticoid replacement (Prednisone, for example) is given.
Hypothyroidism

This is a disease of thyroid deficiency. Dogs with Hypothyroidism have metabolic rates below normal. The most common symptom is hair loss that involves the front of the neck down to the chest, the sides of the body, the backs of the thighs, and the top of the tail. The hair is excessively dry and brittle, and falls out easily. The exposed skin is dry, thick, puffy, and darkly pigmented. Some dogs develop secondary seborrhea. Other signs include weight gain, intolerance to cold, slow heart rate, absence of heat cycle, lethargy, corneal ulcers, deafness and megaesophagus. It is easy to test for and easy to treat. Your vet will take a simple blood test to check the levels of T3 and T4. If this blood test is low it is a red flag for hypothyroidism. It is effectively treated with a daily or twice a day thyroid hormone replacement using synthetic L-thyroxine (L-T4)(Levathyroxine). Many times once the thyroid is well regulated the ME resolves as well. Megaesophagus occurs in 3% of hypothyroid dogs.
Gastrointestinal (GI) Tract consist of mouth, teeth, tongue, pharynx, esophagus, stomach, small and large intestines.

**Gastrointestinal Causes**
Gastrointestinal disorders associated with acquired megaesophagus include esophagitis, esophageal obstruction, gastric dilatation-volvulus (GDV) or commonly referred to as “bloat”, and hiatal hernia.
**Esophagitis**

Esophagitis can lead to megaesophagus if not treated properly. Acid reflux is the most common cause of esophagitis. Acid reflux is the result of stomach acids passing into the esophageal canal, causing irritation to the tissue lining of the esophagus. Intubation for anesthesia for surgery can also cause esophagitis. Chronic vomiting, dog swallowed a foreign body, retaining ingested pills or capsules, infections and accidental ingestion of irritant chemicals can also cause esophagitis. Megaesophagus comes into play if the condition has progressed to the point that the esophagus has formed a stricture (narrowing) or a pocket where food will lodge itself and the dog regurgitates. Endoscopy (a long thin tube with a tiny camera on the end) may be recommended as a diagnostic tool to differential between esophagitis and megaesophagus or to rule out an obstructive lesion or a tumor. Carafate/Sucralfate to soothe the esophagus is usually prescribed plus a pain reliever such as Tramadol.

**Esophageal obstruction**

Esophageal obstruction can be caused by esophageal foreign bodies, neoplasia (tumors), strictures (narrowing of the esophagus sometimes caused by scar tissue), or vascular ring anomalies.

Foreign bodies or chronic gastroesophageal reflux (GERD) can cause esophageal strictures (scar tissue). The buildup of gastric stomach acid (pepsin and bile salts) and other components of the gastrointestinal juices can cause damage to the protective mucus lining of the esophagus. This can lead to esophagitis (inflammation of the esophagus) and later lead to scar tissue forming in the esophagus. An antacid is often times prescribed to reduce acid reflux.
Esophageal obstruction can also be caused by **vascular ring anomalies**. In 95% of the cases secondary megaesophagus is caused by **PRAA (Persistent Right Aortic Arch)**. Other vascular anomalies associated with secondary megaesophagus included persistent right or left subclavian arteries, double aortic arch, persistent right dorsal aorta, left aortic arch, right ligamentum arteriosum, aberrant intercostals arteries, and persistent left cranial vena cava.

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**Gastric dilatation-volvulus (GDV)**

Dog with chronic "**bloat**" (GDV) with or without volvulus (the stomach rotates or twist in the abdomen), have an increase risk of developing megaesophagus. In these dogs, megaesophagus is due to a weak **lower esophageal sphincter (LES)** caused by a combination of esophagitis from chronic GERD or vomiting; chronic intermittent obstruction of the LES; increased intragastric and intra-abdominal pressures; and delayed gastric emptying. **Promotility drugs** that may be effective in emptying the stomach are **Metroclopamidine (Reglan)** and **Propulsid (Cisapride)**.

Bloat occurs when the dog’s stomach fills with gas and twists, cutting off blood and oxygen to the stomach. As the swollen stomach pushes against surrounding organs and large blood vessels, blood flow to the heart eventually stops. Without immediate treatment, shock and a painful death occurs within minutes. Any dog can have GDV, larger deep chested dogs are more at risk. It is suggested that you avoid strenuous exercise after eating or drinking, eating slower and more frequent meals may help as well.

Dogs with chronic GDV may need a surgery called “**gastropexy**”. This surgery returns the stomach to the normal position permanently by surgically attaching the stomach to the abdominal wall to prevent twisting.
Hiatal Hernia
Any time an internal body part pushes into an area where it does not belong, it’s called a hernia. In the case of a hiatal hernia the esophagus is essentially obstructed. There are two types of hernias that canines can have; a Type 1 or “sliding” hernia or Type II a paraesophageal or “rolling” hernia. Xrays and CT imaging are used as diagnostic tools. Fluoroscopy, which is a real-time radiographic procedure that utilizes a low dose, continual x-ray transmitted through the body onto a fluorescent screen and generates a short dynamic movie-like study, may show motility in motion. Temporary relief of symptoms may be achieved with drugs such as omeprazole (Prilosec) and cimetidine (Tagamet). In most cases surgery to correct the hernia is required, usually involving surgical reduction of the hernia followed by placation of the esophagus hiatus and left flank gastropexy.

Paraneoplastic Causes
Megaesophagus is present in 40% of dogs with a thymoma. A thymoma is a tumor of the thymus organ. The thymus is located in front of the heart in the rib cage. It is part of the lymphatic system. It produces lymphocytes (white blood cells) called T-cells. These white blood cells are important because they kill foreign cells, and activate other immune cells to fight infection. They also stop the immune response after a foreign cell has been eliminated. T-cells migrate away from the thymus to other parts of the body such as the lymph nodes, spleen and the bloodstream. As the pet matures, the thymus has served its function and it will wither away. The incidence of a thymoma in dogs with MG is about 3%. The prognosis of a dog with MG with a thymoma is poor. However, removing the tumor can result in resolution of megaesophagus and a decreased AChR antibody titer.
**Toxic Causes**
Toxic substances that can cause megaesophagus include **lead, organophosphates, snake venom and tick-bite paralysis**

**Lead**
Low-level lead exposure causes severe abdominal pain, vomiting, diarrhea and megaesophagus. Lead intoxication can occur from ingestion of batteries, fishing line weights, lead-based paint, linoleum, and plumbing or solder supplies. **A serum blood test to confirm toxicity is available commercially.**

**Organophosphates**
**Organophosphate toxicosis** should be suspected if a dog presents with concurrent weakness and cerebellar signs. Organophosphates exist in **flea collars and insecticides**. Organophosphate toxicity may lead to chronic anorexia, muscle weakness and muscle twitching which may last for days or even weeks. Organophosphate insecticides inhibit cholinesterases and acetylcholinesterase, essential enzymes in the body. Cholinesterases are enzymes which break down acetylcholine, which is a neurotransmitter. Consequently, acetylcholine remains attached to the postsynaptic receptors of the neurons causing continuous, unending nervous transmission to nervous tissue, organs and muscles (smooth and skeletal). This causes seizures and shaking. A blood test to confirm toxicity is available.

**Snake Venom**
**Australian Tiger Snake venom** causes a rapidly progressing myopathy (disease resulting in muscle weakness) of skeletal muscle. If not lethal, Australian Tiger snake venom has a 75% recovery rate for normal esophageal function. **This snake is found in southern regions of Australia, including its coastal islands of Tasmania.**

**Tick Bite Paralysis**
Tick paralysis can occur when a dog is bitten by a certain species of female tick that releases a potent toxin. The toxins released cause **lower motor neuron paralysis**. **The muscles stay in an apparent state of relaxation with this type of paralysis.** Symptoms are gradual in nature and usually begin around 6-9 days after being bitten. Symptoms may include: Vomiting, regurgitation, unsteadiness, high blood pressure, fast heart rate, weakness-especially in hind legs, respiratory muscle paralysis, dysphonia-bark change, drooling, and megaesophagus. Removing and identifying the tick is the first step to prevent further release of toxins. Treatment will depend on the type of tick found and severity of the symptoms.

**Incidental or Temporary ME**
Incidental esophageal dilation is associated with excitement, aerophagia (intestinal gas) general anesthesia and vomiting. In cases of anesthesia and esophagitis associated megaesophagus, **it should resolve within 2 to 14 days.**
**Medications**

Please make sure to check the medications you are giving your dog. For example **Potassium Bromide**, used to control seizure disorders, carries a risk of megaesophagus.

**Idiopathic**

An **idiopathic (cause unknown)** diagnosis is made when all possible scenarios have been eliminated and all testing is complete. This can be the hardest diagnosis to accept. There is no magic pill to make it go away. Your dogs megaesophagus may stay about the same, may improve or it may get worse.

The veterinarian community many times will recommend euthanasia. It can be a bleak, lonely time for an ME pet parent. There is so much hope in this community despite the doom and gloom. There are so many success stories of dogs that have **lived a full life** with the proper management in place. Please visit our home page and review the tabs on megaesophagus for further instructions on how your dog can live a healthy, happy life with MegaE!

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