NUTRITIONAL MANAGEMENT OF METABOLIC DISEASES

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As the understanding of equine metabolic diseases advances, so too must the management strategies implemented to care for horses diagnosed with these problems. For some of these syndromes, dietary changes can completely ameliorate clinical signs, thereby enhancing the lives of horses and ponies. For other syndromes, dietary modifications decrease the likelihood of laminitis and other life-threatening problems.

The endocrine system produces hormones that are distributed throughout the body by the blood. Most hormones have a complex cycle that regulates their activity, and many hormones affect the actions of others. The most important warning that a veterinarian can give the horse owner is to be prepared for some abnormal results (even in normal horses) when performing endocrine testing. Very often, the concentration of one or more hormones will be out of the normal range. The difficulty in interpreting endocrine tests lies in determining what abnormalities are significant and if some other factor (insulin resistance, drug administration, etc.) is influencing the hormone. The “big picture” of the horse’s laboratory results, clinical signs, history, and other coexisting conditions determines which, if any, endocrine abnormalities need to be addressed.

*Nutritional Management of Metabolic Diseases* provides an overview of the most common metabolic problems diagnosed in horses and ponies. Practical nutritional strategies are included for each disease. When appropriate, feed and supplement recommendations are provided.
Pituitary pars intermedia dysfunction (PPID) is caused by hypertrophy of the pars intermedia of the pituitary gland. The pituitary produces excessive amounts of adrenocorticotropic hormone (ACTH), which results in an increased secretion of cortisol from the adrenal glands. The intermediate lobe of the pituitary gland (pars intermedia) is located between the pars nervosa (posterior pituitary) and the pars distalis of the anterior pituitary and is the primary site of disease in horses. PPID is commonly referred to as equine Cushing’s disease.

Affected population
Primarily aged horses and ponies are affected, though younger animals can sometimes develop PPID; no breed or sex predilections are noted.

Clinical signs
- hirsutism (growth of a long, sometimes curly coat that often fails to shed in the spring)
- lethargy
- loss of appetite
- weight loss
- pot-bellied appearance
- recurrent laminitis
- hyperhidrosis (excessive sweating)
- polydipsia (excessive drinking)
- polyuria (excessive urination)
- increased supraorbital fat pads
- chronic infections, dental disease, hoof abscesses, etc.
- increased fat deposition at the crest of the neck and at the tailhead

**Less common clinical signs**
- osteoporosis
- delayed wound healing
- central nervous system dysfunction
- suspensory ligament breakdown
- persistent lactation
- infertility
- blindness

**Diagnosis**

- **Dexamethasone suppression test**
  This test remains the gold standard test for PPID. Blood cortisol is measured before and 19 hours after injection of dexamethasone (a steroid). Dexamethasone is usually injected in the muscle between 4 p.m. and 6 p.m., and cortisol is measured again the following morning. A normal horse should respond with a very low blood cortisol concentration after injection of dexamethasone. Horses with PPID lack this normal response and will continue to have normal to elevated cortisol concentrations (failure to suppress). Results of dexamethasone suppression tests are not reliable if the test is performed during the fall and early winter because cortisol concentrations are higher during that part of the year. There is a slightly increased chance for exacerbating any laminitis that may be present with dexamethasone administration, but this risk is relatively small and most clinicians consider it to be a safe test unless obvious laminitis is present.

- **Adrenocorticotropic hormone (ACTH) concentrations**
  ACTH is produced by the pars intermedia of the pituitary gland and signals the adrenal glands to produce cortisol. Adrenocorticotropic hormone may not be elevated in all horses with PPID, but consistently elevated concentrations probably confirm PPID. Storage and collection of serum for ACTH can be difficult because the hormone is not stable under normal conditions. Assay of ACTH is not reliable during the fall, because its concentration is normally increased during that time of year.

- **Fasting glucose and insulin concentrations**
  Insulin resistance is fairly common with PPID. Elevated cortisol antagonizes the actions of insulin and if insulin resistance is present, then blood concentrations of glucose (blood sugar) and insulin will be increased. Several other conditions can also cause insulin resistance; therefore diagnosis of insulin resistance is a symptom rather than a confirmation of PPID.

- **Thyrotropin-releasing hormone (TRH) stimulation test**
  Some cells in the pars intermedia of the pituitary gland release ACTH in response
to TRH administration, and this will result in increased blood concentrations of cortisol. Unfortunately, the test is infrequently performed and TRH produced specifically for equine diagnostic testing is unavailable, but TRH can be purchased from chemical suppliers.

- **Combined dexamethasone suppression/thyrotropin-releasing hormone test**
  This test is a combination of the dexamethasone suppression and TRH stimulation tests. The combined dexamethasone suppression and TRH stimulation test has the advantage of better diagnosing PPID than when either test is used alone. The disadvantages of this test are the additional expense and collection of multiple blood samples. Both dexamethasone and TRH are administered and several blood samples are obtained at baseline, some hours later, and 24 hours after dexamethasone administration.

- **Domperidone response test**
  This is a relatively new diagnostic test for PPID. Domperidone (Equidone®) is most commonly used for the treatment of decreased milk production and tall fescue toxicosis. Domperidone inhibits dopamine, which allows the pituitary gland in horses with PPID to release even more ACTH. Horses with PPID will double their ACTH concentration 4-8 hours after domperidone administration, while normal horses will maintain normal concentrations of ACTH. As more horses are tested using the domperidone response test, its clinical value will become better known.

**Approach to diagnosis**

- The “big picture” of the horse’s laboratory results, clinical signs, history, and other coexisting conditions determines which, if any, endocrine abnormalities need to be addressed.
- Many, if not most, older horses have PPID. Horses that are not showing any adverse clinical signs of PPID usually do not need to be treated; however, if the horse is having chronic laminitis, weight loss, or recurrent infections, then treatment is likely to be beneficial.
- An aged horse or pony with hirsutism is often assumed to suffer from PPID.

**Treatment**

Pergolide is the drug of choice for treating PPID in horses and ponies.

**Nutritional management**

- For overweight horses with PPID, a ration of primarily hay should be fed. Most hays have low glycemic indexes compared to cereal grains and sweet feeds. Hay rations should be supplemented with a low-inclusion fortified balancer pellet to provide nutrients that may be deficient in the forage (see I.R. Pellet™ on page 9).
- If a horse or pony diagnosed with PPID is underweight or has difficulty maintaining body condition, its ration can be supplemented with additional calories
from a high-fat, low-starch product.

- In addition to providing a concentrated source of energy, vegetable oil has been shown to greatly reduce glycemic response to a grain meal, possibly by delaying gastric emptying. If beet pulp is added to the ration, it should be rinsed to reduce its glycemic index.

- Feeds that are designed for senior horses may not be desirable for horses and ponies with PPID because they may contain ingredients that produce a high glycemic response. Some senior feeds have a similar glycemic index to straight oats.

- Pasture represents a major challenge to those horses with PPID, laminitis, and insulin resistance. The nonstructural carbohydrate (NSC) content of pastures fluctuates with plant species and environmental conditions. Limited grazing should be considered for these horses, and can be accomplished by restricted access to pasture or through the use of a grazing muzzle. Some horses and ponies might not tolerate grazing at all.

### Products from Kentucky Equine Research

**Re·Leve® Original**

![Re·Leve® Original](image)

Appropriate for any health condition in which a low-starch diet is recommended. Specifically, Re·Leve® Original may be beneficial for horses or ponies with PPID that have trouble maintaining weight on an all-forage diet.

**EquiShure®**

![EquiShure®](image)

Supports the microbial population of the hindgut by moderating pH, thereby decreasing the risk of laminitis, either from grain overload or fructan-rich pastures.

**EO·3™**

![EO·3™](image)

Improves insulin sensitivity and decreases the inflammatory response from elevated cortisol levels, lowering the risk of laminitis associated with PPID.

**I.R. Pellet™**

![I.R. Pellet™](image)

A concentrated, low-calorie, low-starch source of vitamins and trace minerals ideal for horse and ponies diagnosed with PPID.
Equine metabolic syndrome (EMS) is believed to be an abnormal adaptation to storing body fat in preparation for times when food will be scarce. Fat acts as much more than an energy-storage tissue. Fat has an important role in stimulating inflammation and affecting insulin metabolism.

Normally, insulin stimulates blood vessels to relax and dilate, which improves blood flow. Recent research has shown, however, that high blood concentrations of insulin (as occurs in EMS) can actually do the opposite, constricting blood vessels and reducing blood flow. Laminitis often occurs with lush pasture growth during late spring to early summer, when pastures have high sugar content (nonstructural carbohydrates).

Affected population
Almost always, horses with EMS can be described as easy keepers that gain or maintain weight with very little feed. Breeds most commonly affected with EMS include ponies, Morgans, Paso Finos, Arabians, American Saddlebreds, Spanish Mustangs, and Warmbloods. Other light breeds can also be affected, although less often. Most horses with EMS are between 5-15 years of age.

Clinical signs
- insulin resistance
- generalized obesity
- regional adiposity or abnormal fat deposits on the body (fat around the tail head that make the tail look inset into the body and fat pads around the shoulder, sheath, or udder)
- recurrent or seasonal laminitis
- hyperlipemia (elevated triglycerides in the blood)
- abnormal estrous cycles
- easy keeper

**Diagnosis**

Diagnosis of EMS can be difficult and usually begins with ruling out pituitary pars intermedia dysfunction (PPID). Occasionally, horses can be affected with both conditions, and some researchers speculate that EMS may predispose some horses to develop PPID and at an earlier age.

**Blood insulin and glucose concentrations**

Affected horses have insulin resistance (similar to type II diabetes mellitus in humans). Blood work may show elevated insulin and blood glucose (sugar) concentrations. Horses with EMS may have both high blood insulin and glucose concentrations, but usually blood glucose is normal and toward the higher end of normal ranges. Insulin results are most reliable if the horse has been completely fasted for at least 12 hours before drawing blood. The higher the fasting insulin level, the more likely that the horse has EMS. Normal horses can have elevated insulin concentration with corticosteroid therapy, pain, or stress. Diabetes mellitus is uncommon in horses. A glucose to insulin ratio of less than 10:1 is considered abnormal. In cases of EMS, a low glucose to insulin ratio is usually due to increased insulin concentration and not elevated blood glucose concentration.

**Thyroid hormones**

Horses with EMS can have normal to low thyroid hormone concentrations. Thyroid hormone levels alone are of little benefit to diagnose EMS.

**Dynamic glucose and insulin tests**

The most reliable tests to accurately determine glucose and insulin responses involve intravenous infusion of glucose and insulin followed by measuring their concentrations over time. However, these tests are complicated and not commonly performed, even in hospital settings.

**Treatment**

- Medical therapy for EMS can involve levothyroxine (thyroid hormone) supplementation and/or metformin.
- Thyroid hormone supplementation helps affected horses lose weight by increasing their basal metabolic rate and the number of calories that they use. Researchers have shown that low-dose levothyroxine supplementation has not produced any adverse effects.
- Metformin is a drug that improves the action of insulin and inhibits glucose production and its metabolism for energy. Research has shown that intestinal absorption of metformin is low in horses and that metformin did not improve insulin resistance in insulin-resistant ponies.
Nutritional management

- The primary goals are to reduce obesity, improve insulin sensitivity, and lower the risk of laminitis through nutritional management.
- Dietary management for EMS is critical. Like humans, horses will lose weight if they are fed less and exercised more. Owners of affected horses often do not realize that they are killing their horses with kindness unless they significantly restrict or eliminate feeding of highly digestible sugars in grain.
- The diet should be forage-based, but pasture intake should be restricted with a grazing muzzle or limited turnout, or completely avoided during periods of lush growth. Lush, quickly growing, early-season grasses often bring about signs of laminitis and cause significant problems for EMS horses. For this reason, limiting turnout, drylot turnout, or wearing a grazing muzzle are important when pastures are rapidly growing. Some EMS horses cannot tolerate unrestricted access to pasture at any time of the year.
- Ideally EMS-affected horses are fed only mature hay at 1.5% of their body weight per day (15 pounds [7 kg] per 1000 pounds [455 kg] of body weight).
- Of primary importance is avoidance of feeds that exacerbate insulin resistance. If any concentrate is fed, it should be low starch, high fiber, and derive most of its calories from fat or other sources.
- It is important to supplement or have ready access to minerals and vitamins because overall feed intake is limited. Most feeds designed for weight loss in horses will have added minerals and vitamins to make up for decreased intake. Because calorie restriction is important, a concentrated balancer pellet should be used for vitamin and mineral supplementation (see I.R. Pellet™ on page 13).
- If affected horses are sound, then as much turnout as possible will encourage exercise and improve insulin sensitivity.
- Though most horses with EMS are easy keepers, there are others that are not, specifically those asked to perform routine exercise. To fuel performance and maintain body condition, these horses might have to be fed a concentrated energy source. A low-starch feed rich in fat and fermentable fiber is appropriate (see Re·Leve® on page 13).
- Limited pasture access, especially when pasture grasses are most likely to contain large concentrations of starches, sugars, and fructans, is essential to reduce the development or flare-ups of pasture-associated laminitis. Pastures have the highest amounts of carbohydrates in the spring during rapid growth when temperatures are cool and from midmorning onward. It is safest to have insulin resistant horses graze pasture from early to midmorning.
- A source of the omega-3 fatty acids eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) may be helpful in reducing insulin resistance (see EO-3™ below).
Products from Kentucky Equine Research

Re•Leve® Original

Appropriate for any health condition in which a low-starch diet is recommended. Specifically, Re•Leve® Original may be beneficial for horses or ponies with EMS that have trouble maintaining weight on an all-forage diet, including performance horses.

EquiShure®

Supports the microbial population of the hindgut by moderating pH, thereby decreasing the risk of laminitis, either from grain overload or fructan-rich pastures.

EO-3™

Improves insulin sensitivity and decreases the inflammatory response from elevated cortisol levels, lowering the risk of laminitis associated with EMS.

I.R. Pellet™

A concentrated, low-calorie, low-starch source of vitamins and trace minerals ideal for horse and ponies diagnosed with EMS.
Hyperkalemic periodic paralysis (HPP or HYPP) is a genetic defect that causes abnormal function of the sodium channel in muscle cells, leading to unbalanced sodium and potassium concentrations and abnormalities in electrical signals that control muscle contractions. HPP is inherited as an autosomal dominant trait (males and females are equally affected and only one copy of the gene is needed to be affected). Horses that are homozygous (have two copies of the gene) are more severely affected than heterozygotes (have just one copy of the gene).

Clinical signs
- sporadic muscle twitching and tremors
- general weakness or trembling
- twitching of the third eyelid
- difficulty chewing or swallowing
- collapse (especially of the hind end) and recumbency
- occasional death

There is a wide range of symptom severity. Some horses with HPP are completely asymptomatic. Horses that are homozygous are more severely affected. Some heterozygous horses may have fewer attacks as they get older and more accustomed to the stimuli that can precipitate attacks.

Affected population
The condition is most often associated with Quarter Horses, but any equine that has the Quarter Horse stallion Impressive in its pedigree has a chance of being affected (Paint Horses, Appaloosas, etc.).
**Diagnosis**

Genetic testing is available through the Veterinary Genetics Laboratory at the University of California-Davis, Mann Equitest, and Vita-Tech Canada. See page 25 for details.

**Treatment**

Acetazolamide is a diuretic that promotes potassium loss in the urine and insulin production. Not only is insulin important to move glucose into cells, it also promotes intracellular movement of potassium.

**Nutritional management**

- Limiting potassium intake is one of the most effective management steps. Although forage is typically rich in potassium, steps can be taken to minimize potassium consumption. Grazing is ideal because the high water content of fresh grass keeps ingested potassium at a low level. Grass or oat hay is preferable to legume hay, although potassium level in any hay can be influenced by forage type, maturity, application of fertilizer, and rainfall. Nutrient analysis can help owners choose hay with low potassium content.
- The feeding schedule for affected horses should provide two to three small grain meals per day to decrease fluctuations in blood insulin concentrations. A more level and constant concentration of insulin helps to maintain more normal blood glucose and potassium concentrations.
- Soybean meal, soybean oil, canola oil, and molasses contain high levels of potassium and should be avoided. The carbohydrate portion of the diet should be built around a combination of oats, corn, or barley.
- Beet pulp-based feeds have been recommended as a readily fermentable fiber source and to reduce dietary carbohydrates. Beet pulp is also low in potassium.
- Keeping dietary potassium no higher than 1% of total intake and limiting consumption of potassium to less than 30 g per meal offers the best chance of minimizing problems for horses with HPP.
- Fasting can precipitate HPP attacks, and feeding schedules should be as regular as possible. Changes in feeding should be made very gradually.
- Steer clear of commercial electrolyte preparations, as many contain potassium. Consider using plain salt or a potassium-free electrolyte product.

**Product from Kentucky Equine Research**

**I.R. Pellet™**

A concentrated, low-calorie, low-starch source of vitamins and trace minerals ideal for horse and ponies diagnosed with HPP that can maintain their weight on forage-only diets.
Recurrent exertional rhabdomyolysis (RER) is suggested to be an autosomal dominant heritable trait. The underlying defect is believed to be abnormal calcium release or regulation. The sarcoplasmic reticulum surrounds muscle fibers, and stores and releases calcium. Calcium is integral in triggering and sustaining muscle contraction. Muscle biopsies taken from horses with RER have been found to have enhanced calcium release when exposed to caffeine or halothane (an anesthetic gas).

Affected population
About 5-10% of Thoroughbreds and possibly Standardbreds and Arabians are affected. RER is more common in fillies, especially those that are high-strung or nervous. Thoroughbreds used in eventing competition may show signs after the steeplechase or at the beginning of the cross-country phase of a three-day event.

Clinical signs
- firm, painful muscles
- lameness
- stiffness
- sweating
- short stride and reluctance to move after moderate exercise

RER is a syndrome of repeated episodes of tying-up. In contrast to typical tying-up, RER occurs in fit horses and clinical signs are observed when horses are training at less than maximal effort. Episodes of RER seem to occur more often during training and not after racing or breezing, and may become more frequent as fitness improves.
Diagnosis
Diagnosis is based on breed, clinical signs, and lack of histological evidence of polysaccharide storage myopathy in muscle biopsy samples. Muscle biopsy testing is available from the Neuromuscular Diagnostic Laboratory at the University of Minnesota. See page 25 for details.

Nutritional management
- Replacing much of the grain in the diet with a low-starch, high-fat feed will significantly decrease the likelihood of an episode in affected horses. Most horses with RER have medium to high energy requirements and need significant calories supplied above those found in the forage portion of the ration. An appropriate feed should be fortified to be fed at fairly high levels of intake (9-13 lb or 4-6 kg per day). The feed should be low in NSC (<10%), high in fat (>10%), and supply a significant portion of its energy from fermentable fiber.
- Increased dietary fat as an energy source can reduce the clinical signs and number of episodes of tying-up. Switching part of the calories to fat may help calm some nervous, fractious horses. Muscle enzyme activities improve within a week once the diet is changed to a low-starch, high-fat diet (see Re-Leve® below).
- Ensure that dietary vitamin E and selenium are adequate. Supplementation of vitamin E and selenium is recommended for horses that have repeated episodes of tying-up or muscle disease. Vitamin E and selenium both function as antioxidants and can each offset a deficiency of the other, as long as both are not deficient. Several studies have shown that water-soluble, natural vitamin E (d-α-tocopherol) is better absorbed than synthetic vitamin E and will more quickly achieve higher concentrations in the tissues (see Nano-E® on page 18).
- Efforts should be made to reduce stress in horses with RER, such as maximizing turnout, low-dose tranquilization, equine companionship, and feeding and training before other horses.

Products from Kentucky Equine Research
Re-Leve® Original
Appropriate for any health condition in which a low-starch, high-fat diet is recommended. Specifically, Re-Leve® Original may be beneficial for horses with RER that require significant dietary energy to perform optimally.
As a water-soluble, natural vitamin E supplement, Nano-E® provides fast-acting antioxidant protection to working muscles.

Preserve™ provides antioxidant protection through vitamin E, organic selenium, and vitamin C, supporting proper muscle function and recovery after intense exercise.

B·Quiet™ supports horses with low thiamine levels, which can cause excitability. The paste form contains magnesium, an essential mineral for digestion of protein and starch.
When examined under a microscope, the muscle of horses with polysaccharide storage myopathy (PSSM) contains amylase-resistant inclusions of abnormal polysaccharide (glycogen, the large starch molecule used by muscle for energy). This abnormal polysaccharide is less branched than normal glycogen because the enzyme that links glucose molecules together is more active than the enzyme that creates branch points in horses with PSSM. Studies have also shown that horses with PSSM are more sensitive to the effects of insulin, and they take up glucose into muscle more efficiently than unaffected horses. There are two forms of polysaccharide storage myopathy, type 1 and type 2.

Clinical signs
- firm, painful muscles
- skin twitching
- sweating
- weakness
- reluctance to move with light exercise
- gait abnormalities
- mild colic
- muscle wasting
- serum creatine kinase (CK) and aspartate aminotransferase (AST) activities are elevated
Type 1 polysaccharide storage myopathy

Cause
A genetic disorder, suggested to be an autosomal dominant trait (males and females equally affected, only one copy of the gene needed to be affected). Most horses affected with PSSM have a genetic mutation in their glycogen synthase-1 enzyme (type 1 PSSM).

Affected population
At least 20 breeds are known to be affected, including Quarter Horses and related breeds, Belgians, Percherons, Morgans, Mustangs, Tennessee Walking Horses, and some Warmblood breeds. Though rare in Clydesdales and Shires, 36-50% of Belgians and Percherons are affected, and 8% of Quarter Horse-related breeds are affected. Clinical signs usually begin by two or three years of age but may occur in weanlings.

Diagnosis
Muscle biopsy samples are examined for the presence of amylase-resistant crystalline polysaccharide. Genetic testing is available. Both tests are available from the Neuromuscular Diagnostic Laboratory at the University of Minnesota. See page 25 for details.

Type 2 polysaccharide storage myopathy

Cause
Horses with type 2 PSSM have another, yet unidentified, mechanism.

Affected population
Quarter Horse-related breeds, a few Arabians, Thoroughbreds, and possibly other light breeds are affected. Clinical signs usually begin by two or three years of age but may occur in weanlings.

Diagnosis
Muscle biopsy samples are examined for the presence of abnormal, amylase-resistant polysaccharide. Muscle biopsy testing is available from the Neuromuscular Diagnostic Laboratory at the University of Minnesota. See page 25 for details.

Nutritional management of type 1 and type 2 PSSM
- Replacing much of the grain in the diet with a low-starch, high-fat feed will significantly decrease the incidence of clinical signs in affected horses. Most of the horses diagnosed with PSSM have lower energy requirements than those with recurrent exertional rhabdomyolysis (RER). Therefore, the concentration of other
nutrients needs to be greater than in feeds designed for horses diagnosed with RER in order to provide similar nutrient intake at a lower energy level (see Re·Leve® below). Hay that is high in NSC should be avoided for horses with PSSM. Most grass hays will be sufficiently low enough in NSC (less than 24%).

- Ensure that dietary vitamin E and selenium are adequate. Supplementation of vitamin E and selenium is recommended for horses that have repeated episodes of tying-up or muscle disease. Vitamin E and selenium both function as antioxidants and can each offset a deficiency of the other, as long as both are not deficient. Several studies have shown that water-soluble, natural vitamin E (d-α-tocopherol) is better absorbed than synthetic vitamin E and will more quickly achieve higher concentrations in the tissues (see Nano·E® below).

- Frequent, regular exercise and reduction in dietary carbohydrates coupled with increased fat are essential to manage PSSM. Horses with PSSM should have a regular exercise program if clinical signs of muscle soreness allow and ideally should be turned out all the time.

Products from Kentucky Equine Research

**Re·Leve® Concentrate**

Appropriate for any health condition in which a low-starch, high-fat diet is recommended. Specifically, Re·Leve® Concentrate provides essential protein, vitamins, and minerals for horses with lower energy requirements.

**Re·Leve® Original**

Appropriate for any health condition in which a low-starch, high-fat diet is recommended. Specifically, Re·Leve® Original may be beneficial for horses with PSSM that require significant dietary energy to perform optimally.

**Nano·E®**

As a water-soluble, natural vitamin E supplement, Nano·E® provides fast-acting antioxidant protection to working muscles.

**Preserve™**

Provides antioxidant support through vitamin E, organic selenium, and vitamin C, supporting proper muscle function and recovery after intense exercise. Also includes magnesium, a mineral essential for proper digestion of protein and starch.
Malignant hyperthermia (MH) is a genetic disorder suggested to be an autosomal dominant trait (only one copy of the gene needed to be affected). Males and females are equally affected. MH can occur with type 1 polysaccharide storage myopathy (PSSM).

Affected population
MH is found in Quarter Horse-related bloodlines, though less than 1% of the Quarter Horse population has the disease. MH is present in high frequency in two Quarter Horse bloodlines. Mature horses are usually affected.

Clinical signs
- high body temperature (can be life-threatening), either under general anesthesia or with an episode of tying-up
- metabolic failure
- death after an episode of tying-up, especially when type 1 PSSM is also present

Diagnosis
Genetic testing is available from the Neuromuscular Diagnostic Laboratory at the University of Minnesota. See page 25 for details.

Nutritional management
- If this horse is also diagnosed with polysaccharide storage myopathy (PSSM), the horse may be fed in such a way that minimizes PSSM symptoms, namely a diet that is low in starch and high in fermentable fiber and fat.
Ensure that dietary vitamin E and selenium are adequate. Supplementation of vitamin E and selenium is recommended for horses that have repeated episodes of tying-up or muscle disease. Vitamin E and selenium both function as antioxidants and can each offset a deficiency of the other, as long as both are not deficient. Several studies have shown that water-soluble, natural vitamin E (d-α-tocopherol) is better absorbed than synthetic vitamin E and will more quickly achieve higher concentrations in the tissues.

Horses with both MH and PSSM are more difficult to manage and may not respond as well to changes in diet and exercise.

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Re-Leve® Concentrate

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Nano-E®

As a water-soluble, natural vitamin E supplement, Nano-E® provides fast-acting antioxidant protection to working muscles.

Preserve™

Provides antioxidant support through vitamin E, organic selenium, and vitamin C, supporting proper muscle function and recovery after intense exercise. Also includes magnesium, a mineral essential for proper digestion of protein and starch.
Selected references


Testing resources

For polysaccharide storage myopathy (PSSM), recurrent exertional rhabdomyolysis (RER), and malignant hyperthermia (MH) testing:

■ Neuromuscular Diagnostic Laboratory
University of Minnesota
1333 Gortner Ave.
St. Paul, MN  55108
612-625-8787
800-605-8787
www.cvm.umn.edu/umec/lab/home.html

For hyperkalemic periodic paralysis (HPP) testing:

■ University of California-Davis
Veterinary Genetics Laboratory
P.O. Box 1102
Davis, CA 95617-1102
530-752-2211
www.vgl.ucdavis.edu/

■ Mann Equitest
335 Laird Road
Guelph, ON
N1G 4P7 Canada
519-836-2400

■ Vita-Tech Canada
1345 Denison Street
Markham, ON
L3R 5V2 Canada
905-475-6499

Useful websites

■ Kentucky Equine Research
www.ker.com
www.Equinews.com

■ University of Minnesota
Neuromuscular Diagnostic Laboratory
www.cvm.umn.edu/umec/lab/home.html

■ University of California-Davis
Veterinary Genetics Laboratory
www.vgl.ucdavis.edu/
Kentucky Equine Research (KER) was founded in 1988 when Joe Pagan, Ph.D., realized that information generated from research did not reach the individuals who needed it most: feed manufacturers and horse owners. Since then the primary focus of the company has been bridging the gap between the research community and horsemen.

The company accomplishes this through research, consultation, and nutritional solutions.

KER is one of the most prolific private equine nutrition and exercise physiology research companies in the world. The quantity of published research derived from studies conducted at KER rivals that of leading universities. In addition to its own research, KER collaborates with prominent universities to develop and patent products and diagnostic techniques that target specific problems in horses of all ages and uses.

Aside from its research efforts, KER serves as an industry-wide consultant. At the core of the KER consultation services rests its Team Members, a collection of feed manufacturers dedicated to the production of high-quality feeds. The roster of Team Members continues to grow as feed manufacturers around the world recognize the value added to their equine products through KER’s research, technology, and credibility.

The advantages of purchasing a feed manufactured by a Team Member are numerous: expert formulation of feed designed specifically in the region or country in which it is to be fed, a complete understanding of trends in equine nutrition, and on-site consultation by leading equine nutritionists, just to name a few.

As a result of its research and consultation endeavors, KER has developed nutritional supplements to assist horse owners and managers in overcoming health problems. Many of these products are completely unique to the industry and provide nutritional solutions to common management problems.

For more information on Kentucky Equine Research, visit ker.com.