Facts About Myopathies
Friends:

When I was in my early teens, I was having an ice cream at the mall with some friends, and suddenly I couldn’t move a muscle. The paramedics and the fire department came, and I had to be wheeled out on a stretcher. The doctors, my parents and friends were baffled by what had happened. Many of the doctors doubted there was anything wrong with me. I had similar attacks over the years. Finally, it was found that I had hyperkalemic periodic paralysis, one of the myopathies described in this booklet.

If you’ve recently found out you have an inheritable myopathy, you understand what my family and I went through. Because of the rarity of these diseases, your primary physician may not be aware that many of these myopathies can be managed with medication or changes in diet and exercise. This is why it’s very important that you get all the information you can about your disorder. This booklet will help you get started.

Learning that you or your child has a rare myopathy can be frightening and confusing. Some people may think you’re lazy or mentally unbalanced, and that can hurt. One thing you can be sure of is that your disorder wasn’t caused by anything you or your parents did, and you didn’t catch it from anyone. As this pamphlet explains, each inheritable myopathy is caused by a very uncommon genetic defect that people often don’t even know they have. (Two of the myopathies aren’t inheritable; they’re caused by thyroid imbalances that can occur for no known reason.)

I’ve had to make many adjustments to living with my myopathy. I know what foods and activities can trigger an attack. With a balanced diet keeping my potassium down and my body hydrated, I can
exercise and live a full life. I have a career as a legal assistant and a supportive husband who’s knowledgeable about my hyperKPP, and we have two healthy children. These victories were possible with the assistance of caring doctors I found through the Muscular Dystrophy Association. MDA also has provided endless support and information.

Like my hyperKPP, many myopathies can be controlled so that they cause very little limitation on your life. But, if you have one of the myopathies that has more disabling effects, you can be sure that MDA is your best ally — helping you to find appropriate therapists, and to locate and purchase important assistive devices.

And today, people with disabilities have many opportunities to develop and use their abilities. Federal law guarantees us a public education, equal employment opportunity and access to public places.

Computers and other technological advances help us to move around, communicate and work.

“MDA is Here to Help You,” on page 26, tells of the Association’s services. MDA is also the world leader in research on neuromuscular diseases, and its scientists have made many discoveries about myopathies in recent years.

This booklet will give you the basic facts about the inherited and endocrine myopathies, and MDA will help you answer all your questions as they arise. As you face the challenges ahead, please be assured that we’re making rapid progress toward better treatments and a cure. And remember, you’re not alone.

Christine Swanson
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What are Myopathies?

The word myopathy means “disease of muscle.” More specifically, myopathies are diseases that cause problems with the tone and contraction of skeletal muscles (muscles that control voluntary movements.) These problems range from stiffness (called myotonia) to weakness, with different degrees of severity.

Some myopathies, especially when they’re present from birth, have life-threatening complications. But, with time and physical therapy, some people born with myopathies can gain muscle strength. Others often can manage their symptoms through medication, lifestyle modifications, or use of orthopedic and respiratory equipment.

This booklet focuses on the six types of inherited myopathy (a myopathy that can be passed from parent to child) in MDA’s program.

- **myotonia congenita** (Thomsen disease and Becker type)
- **paramyotonia congenita** (Eulenberg disease)

Myopathies can cause weakness or stiffness in all of the body’s voluntary muscles. Because muscles support the body’s posture, severe muscle weakness can lead to skeletal deformities.
In the inherited myopathies, genetic mutations cause defects in various proteins necessary for muscle tone and contraction.

What are muscle tone and contraction, and what controls them?

**Contraction** is the forceful shortening or tightening of muscle, which pulls on the joints to cause movement. In other words, when your brain “tells” a muscle to move, you cause it to contract, and it’s then able to do what you’re asking.

**Muscle tone** refers to a readiness for contraction that makes resting muscle resistant to stretching. A toned muscle holds its shape and elasticity and is able to respond by contracting when you want it to move. Bodies with poor muscle tone appear “floppy.” Good muscle tone is important for posture and coordination.

A skeletal muscle’s tone and contraction depend on its ability to respond to stimulation from nerve cells, which relay signals from the brain, such as the decision to move your hand or leg. A muscle is actually a bundle of individual muscle cells, and a cluster of muscle cells stimulated by a single nerve cell is called a **motor unit**.
The process of muscle contraction begins when the nerve cells release chemical signals onto the muscle cells. These signals cause the opening of **ion channels**, pores in each muscle cell’s outer surface that open and close to regulate the movements of charged atoms called ions. Different types of ion channels allow specific ions — sodium, calcium, potassium or chloride — to pass into and out of the muscle cell, creating electrical currents. Opening of sodium and calcium channels causes an **electrical excitation** that leads to contraction, while opening of potassium and chloride channels keeps the excitation from occurring.

The purpose of the electrical excitation is to rapidly spread the signal to contract throughout the entire muscle cell, and to stimulate the opening of still more channels that release calcium from internal compartments in the muscle cell.

Finally, the freed calcium ions trigger muscle contraction by stimulating the sliding action of **filament proteins**. These rodlike proteins run lengthwise within the muscle cell and are anchored at opposite ends by scaffolds called **z-discs**. When the filament proteins slide past each other — in a ratchet-like mechanism that is fueled by cellular energy sources — they cause shortening of the muscle cell and shortening (contraction) of the whole muscle.

A muscle cell is stimulated to contract by chemical signals sent from an adjoining nerve cell (1). Those signals open ion channels at the muscle cell’s surface, causing an inward/outward flow of ions that acts as an electrical current (2). Inside the muscle cell, the current spreads and causes opening of ion channels that line calcium storage compartments, releasing the calcium ions trapped within (3). The freed calcium ions trigger nearby filament proteins to slide past each other, pulling the Z-discs closer together and shortening the muscle cell (4).
If this process is disrupted at any stage between the nerve’s signaling the muscle and the filament proteins’ action, the muscle loses its normal capacity for tone and contraction. At one extreme, the muscle might be limp and weak, and at the other extreme, the muscle may be involuntarily active and unable to relax.

What goes wrong in inherited myopathies?

Many of the inherited myopathies are caused by mutations that interfere with ion channels, causing either too much or too little current from flowing through the muscle cells. These disorders (myotonia congenita, paramyotonia congenita, periodic paralysis and central core disease) are sometimes called channelopathies.

Central core disease seems to damage, and thus weaken, muscles by causing an excess release of calcium from internal storage compartments.

A fifth myopathy, nemaline myopathy, is caused by mutations that affect filament proteins. When the filament proteins fail to do their jobs, muscles can’t contract properly, causing a loss of tone and strength.

At least one myopathy (a type of myotubular myopathy) is caused by mutations in a muscle protein required for normal muscle development. When this protein is absent or inactive, the muscles don’t form properly.

Some of the inherited myopathies are congenital, meaning they cause problems from the time of birth. Others have a later onset, with symptoms appearing in childhood or adulthood.

Myopathies aren’t contagious, and they aren’t caused by overexertion. However, exercise can aggravate some of the myopathies, because of mutations that change the way muscles respond to activity.

What happens to someone with an inheritable myopathy?

Some of the congenital inheritable myopathies cause severe, general muscle weakness that creates problems with basic activities like swallowing and breathing. These problems can be fatal if not dealt with, but they can be treated with assistive medical devices like feeding tubes and mechanical ventilators.
Other inheritable myopathies cause episodes of muscle weakness or stiffness that are milder and more localized and temporary in nature. These episodes often can be managed through medication, or by careful control of exercise and diet.

Unlike muscular dystrophies, myopathies usually don’t cause muscles to die but just keep them from working properly. Also, myopathies are usually nonprogressive — that is, a myopathy usually doesn’t grow worse over a person’s lifetime. In fact, some children with myopathies gain strength as they grow older.

Finally, some myopathies can give people a listless facial expression, caused by weakness of muscles in the face. Myopathies have no effect on intelligence.

Special issues in inheritable myopathies

- **Anesthesia**: People with myopathies can experience a range of adverse reactions to certain anesthetic drugs used during surgery. Although these drugs sometimes just aggravate the myopathy, they also can produce potentially fatal reactions, such as *malignant hyperthermia*, which refers to a dangerously high increase in body temperature. People with central core disease are especially at risk for malignant hyperthermia because the two conditions are sometimes caused by the same ion channel defects.

Malignant hyperthermia is triggered by certain inhaled anesthetics (like halothane) and certain muscle relaxants (like succinylcholine). These drugs can intensify ion channel defects and boost muscle metabolism — the set of chemical reactions that provides energy to muscle. The increased metabolism raises body temperature, and causes excessive contraction and *rhabdomyolysis* — a process of acute muscle breakdown. The resulting leakage of ions and muscle proteins into the circulatory system can cause life-threatening damage to the heart, lungs and kidneys.

People with central core disease aren’t always susceptible to malignant hyperthermia. Those who are susceptible won’t experience malignant hyperthermia unless they’re exposed to triggering anesthetics.

Before having surgery, people who have a personal or family history of central core disease — or any other myopathy — should consult their doctors about the risks of anesthesia and
about the availability of “nontriggering” anesthetics.

**Respiratory care:** Nemaline myopathy and congenital (X-linked) myotubular myopathy may cause weakening of the respiratory muscles (those that control the lungs). Therefore, people with either of these diseases may need to use mechanical ventilation to support breathing, and should have their breathing monitored regularly by a specialist. Also, weak lungs are susceptible to infection, so signs of respiratory illness should be taken seriously.

**Cardiac care:** With the exception of Andersen-Tawil syndrome, the myopathies almost never affect heart muscle directly. However, sometimes they can cause indirect damage to the heart.

In nemaline myopathy and congenital myotubular myopathy, an inadequate oxygen supply to the body during severe bouts of respiratory weakness can lead to heart problems.

In one form of periodic paralysis (the hypokalemic form), attacks of weak-

ness are associated with a decrease in blood potassium level, whereas another type of periodic paralysis (the hyperkalemic form) causes an increase. Either change can indirectly cause an irregularity in with the rhythmic contraction of the heart.

A rare type of periodic paralysis, Andersen-Tawil syndrome, is caused by an inherited defect in a potassium channel that is found in both the heart and skeletal muscle. As a result, these patients may have heart rhythm disturbances, even if their blood potassium level is normal.

People with these diseases should be wary of potential heart problems and have their cardiac function checked by a specialist.
On being told they have a genetic disorder such as an inheritable myopathy, patients often ask, “But it doesn’t run in the family, so how could it be genetic?” Inheritable myopathies can run in a family, even if only one “blood relative” in the family has it.

This is because genetic diseases like inheritable myopathies can be inherited in a variety of ways: X-linked, autosomal dominant and autosomal recessive. Or, a new spontaneous mutation, may occur for the first time in a child.

X-linked means that the genetic mutation (or defect) is located on the X chromosome. For many X-linked diseases, a normal copy of the gene can compensate for the defective copy. Because males have only one X chromosome while females have two, X-linked diseases almost always affect males.

Autosomal means the mutation occurs on a chromosome other than the X or Y. Therefore, autosomal diseases affect males and females equally.

Autosomal recessive means that two copies of a defective gene are required for the full-blown disease. One copy is inherited from each parent, neither of whom would normally have the disease but would be a “carrier.”

Autosomal dominant means that one copy of a defective gene is enough to cause disease. So, a person who inherits the defective gene from a parent will have the disease, as will the parent.

Inheritable myopathies passed on in an autosomal dominant pattern can be easy to trace through the family tree. By contrast, X-linked or autosomal recessive disorders might seem to occur “out of the blue.” But in reality, one or both parents might be carriers who silently harbor a genetic mutation. Many parents have no idea they’re carriers of a disease until they have a child with the disease.

Spontaneous mutations literally come “out of the blue,” when a new mutation occurs during the child’s conception. After they occur, these mutations can be passed on to the next generation.

A good way to find out more about your risk of inheriting or passing on an inheritable myopathy is to talk to your MDA clinic physician or a genetic counselor. Also, see MDA’s pamphlet “Facts About Genetics & Neuromuscular Diseases.”
What are the symptoms and treatments for each inheritable myopathy?

Myotonia congenita

Cause:
This disease is caused by mutations in the gene for a chloride channel that’s necessary for shutting off the electrical excitation that causes muscle contraction.

Inheritance:
autosomal recessive (Becker type), autosomal dominant (Thomsen)

Onset:
early to late childhood

Symptoms:
The main problems faced by people with this disease are delayed muscle relaxation and muscle stiffness, typically provoked by sudden movements after rest. The stiffness can interfere with simple activities like walking, grasping and chewing, but is usually manageable by doing warm-up movements. The disease doesn’t cause any muscle wasting; instead, it sometimes can cause muscle enlargement and increased muscle strength. Becker-type myotonia is the most common form of myotonia congenita, while Thomsen disease is a very rare, relatively mild form.

Treatment:
Someone who has myotonia congenita can lead a long, productive life, and can even excel at sports where strength is more important than agility. Your MDA clinic director can tell you about appropriate exercises, and if necessary, appropriate medications for dealing with muscle stiffness.

Paramyotonia congenita

Also called:
Eulenberg disease. (Some researchers regard paramyotonia congenita as a form of periodic paralysis.)

Cause:
Sodium channels normally open to cause muscle excitation, and then close to end the excitation. In paramyotonia congenita, mutations in the muscle sodium channel gene prolong the channel’s opening, causing higher-than-normal muscle excitation.

Inheritance:
autosomal dominant

Onset:
congenital (existing at birth)

Symptoms:
Paramyotonia congenita causes episodes of muscle stiffness and weak-
ness — mostly in the face, neck, and upper extremities — that can last from minutes to hours. The stiffness is sensitive to exercise and cold. During brief exercise, overexcitation of muscles can cause stiffness, and with prolonged exercise, the overexcitation can occasionally lead to a fatigue-like weakness or even complete paralysis. Cold exposure can have similar effects, but some people experience muscle stiffness, weakness or, sometimes, temporary paralysis even when they’re warm.

**Treatment:**
By avoiding strenuous exercise and cold, most people with this condition can largely escape disability. But medications can be beneficial, especially for those who experience symptoms independent of exercise and cold. Your MDA clinic director can give you more information about these medications.

**PERIODIC PARALYSES**
In these diseases, faulty ion channels cause “attacks” of temporary muscle weakness that can result in temporary paralysis when severe.

There are different types of periodic paralysis, distinguished by what happens to potassium levels in the blood (specifically the serum, or fluid portion of the blood). In the hyperkalemic type (hyperKPP), high serum potassium levels can cause attacks. In the hypokalemic type (hypoKPP), low serum potassium levels can trigger attacks. (Kalemic refers to potassium; hyper means too much and hypo too little.) Total potassium in the body is usually normal. With HyperKPP, the serum potassium is high because it has moved out of muscle into the blood. Conversely, in episodes of hypoKPP, potassium moves from the blood into muscle cells, where it is trapped. Unlike the case for most myopathies, many people with hypoKPP and some people with hyperKPP experience progressive, permanent muscle damage that occurs independently of the attacks.

Andersen-Tawil syndrome affects ion channels present not only in skeletal muscles but also in the heart. This
makes control of episodes even more urgent than in other forms of periodic paralysis.

(Severe bouts of weakness related to a disturbance of thyroid function sometimes are confused with inherited periodic paralysis. Your doctor may order blood tests to check your thyroid function, especially if a family history of this condition is lacking.)

**Hyperkalemic periodic paralysis**

*Cause*: This disease is caused by distinct mutations in the muscle sodium channel gene. These mutations may either disrupt the closing of the sodium channel that normally helps end the brief excitation to trigger contraction (a process called channel inactivation), or cause channels to open too easily (activation). These changes may enhance excitability to produce stiffness (myotonia), or completely overwhelm the muscle fiber, rendering it inexcitable, causing weakness or paralysis. In all of us, higher potassium in the blood will promote muscle excitation. For individuals with hyperKPP, the sodium channel defect amplifies this effect to produce stiffness or weakness.

*Inheritance*: autosomal dominant

*Onset*: childhood

*Symptoms*: Attacks of weakness usually last 15 minutes to an hour, but can last for a day or more. They can recur daily in severe cases. The attacks commonly occur after vigorous exercise followed by rest, and can be aggravated by stress, pregnancy or foods high in potassium. During attacks not caused by excess potassium intake, a person can become hyperkalemic or remain normokalemic (with no change in serum potassium levels). The frequency of attacks usually declines after middle age.

*Treatment*: To keep hyperKPP attacks to a minimum, stick to a diet rich in carbo-
hydrates and low in potassium, and avoid strenuous exercise. When you do exercise, be sure to “cool down” with mild activity before resting.

During an attack, certain prescription drugs can be used to alleviate symptoms. Your MDA clinic director can give you more specific information on how to manage hyperKPP through appropriate exercise, diet and medication.

**Hypokalemic periodic paralysis**

*Causes:*  
As in all forms of periodic paralysis, episodes of weakness in hypoKPP are caused by a temporary loss of muscle excitability. Interestingly, this disease may be caused by genetic defects in either the calcium channel or the sodium channel. The mutations cause a loss of muscle excitability when the serum potassium is low.

*Inheritance:*  
autosomal dominant

*Onset:*  
early childhood to adulthood

*Symptoms:*  
Attacks of weakness can occur daily and usually happen in the morning (during waking) or at night. Some people with the disease might experience only a few mild attacks in their lifetime. But the most severe attacks cause nearly full-blown paralysis. Slowly progressive permanent weakness in the legs often develops after age 50.

*Treatment:*  
As in hyperKPP, attacks of hypoKPP can be prevented by avoiding strenuous activity and alleviated by medications. The dietary precautions, however, are nearly opposite. High-carbohydrate foods can trigger hypokalemia and contribute to an attack, while potassium intake can restore serum potassium levels and stem an oncoming attack. Ask your MDA clinic director for specific recommendations about diet, exercise and medications.

**Andersen-Tawil syndrome**

*Causes:*  
This disease is caused by defects in a potassium
channel normally present in skeletal and cardiac muscles. Mutations in this potassium channel gene interfere with the ability of a muscle to stay poised, ready to contract. As a result, periodic episodes of paralysis may occur, and the heartbeat can become irregular.

**Inheritance:**
autosomal dominant

**Onset:**
childhood to adolescence

**Symptoms:**
Periodic episodes of weakness lasting hours to days occur, as can severe heartbeat irregularities, sometimes accompanied by loss of consciousness.

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**Central core disease/malignant hyperthermia susceptibility**

**Causes:**
This rare disease appears to have multiple origins. But it’s commonly caused by defects in a channel that acts like a gate to internal calcium stores. The defect causes leakage of calcium from the stores, which appears to damage muscle cells.

**Inheritance:**
autosomal dominant, possibly autosomal recessive in rare cases

**Onset:**
congenital

**Symptoms:**
The disease is named for damaged areas within muscle cells (the cores), where the filament proteins are disorganized, and mitochondria (the tiny energy-producing factories that power muscle contraction) are missing. The impact of the cores on disease severity isn’t clear.

This disease causes poor muscle tone (hypotonia) and persistent muscle weakness in infants. In rare cases,
How Ion Channels Regulate Muscle Contraction

1. Acetylcholine leaves the nerve fiber and docks on receptors in the muscle membrane, causing that area of the muscle fiber to become slightly more positive (“depolarized”).

2. Sodium channels open in response to this small depolarization, permitting a huge flow of positively charged sodium ions to enter the muscle fiber. The depolarization is greatly amplified, and a brief electrical impulse (“action potential”) spreads throughout the fiber.
Depolarization of the muscle fiber is sensed by calcium channels and triggers the release of calcium ions from internal storage areas. This flood of released internal calcium is the chemical signal that causes the thick and thin filaments of the muscle fiber to slide past each another (contract).

The sodium channels spontaneously close, potassium channels open, and positively charged potassium ions exit the fiber. Chloride channels also stay open, and negatively charged chloride ions enter the fiber. All these actions cause the inside of the fiber to become more negative (“repolarized”).
toddlers with the disease fail to walk at all, but usually they’re just late in reaching motor milestones. Older children and adults typically experience mild disabilities that worsen slowly with time, if at all. Due to chronic muscle weakness, many people develop skeletal deformities, including joint dislocations and scoliosis, or curvature of the spine that can compress vital internal organs.

People with this disease should be cautious about surgery because they face an especially high risk of malignant hyperthermia, a potentially fatal reaction to certain anesthetic drugs (see “Anesthesia,” page 8).

**Treatment:**
Someone with a severe form of central core disease might need a walker or other support devices for mobility, but many people require none. Unlike the case for other myopathies, people with this disease can benefit from exercise. Scoliosis and other skeletal problems can usually be corrected by use of orthopedic devices or by surgery. Your doctor or MDA clinic director can tell you more about the risks of surgery, and about anesthetic drugs that are safe.

**Nemaline myopathy**

*Also called:*
rod body disease

**Causes:**
This disease is caused by a variety of genetic defects, each one affecting one of the filament proteins required for muscle tone and contraction.

**Inheritance:**
autosomal recessive, autosomal dominant

**Onset:**
birth to adulthood

**Symptoms:**
The disease gets its name from the fact that the muscle cells contain abnormal clumps of threadlike material — probably disorganized filament proteins — called nemaline bodies (*nema* is Greek for “thread”). It causes weakness and poor tone (hypotonia) in the muscles of the face, neck and upper limbs, and often affects the respiratory muscles (those that control breathing).

The infantile-onset cases tend to be the most severe. Usually, infants with the disease lack the muscle strength and tone required for simple postures and movements. They also have serious difficulties with feeding and respiration. Although many infants with the disease die from respiratory failure or lung infections, some survive to adulthood. Affected children usually attain motor...
Children and adults also can benefit from respiratory support, since respiratory failure during sleep can be a persistent danger. Mobility and strength can be improved significantly by physical and orthopedic therapies. If you or your child has nemaline myopathy, your MDA clinic director can provide further information about treatments.

**CENTRONUCLEAR MYOPATHIES**

The centronuclear myopathies are named for the mislocation of cell nuclei in the muscle fibers. Normally, these nuclei are arranged around the periphery of the fiber. In these disorders, many of them are centrally located instead.

Myotubular myopathy is a very severe form of centronuclear myopathy. Because of its X-linked inheritance pattern, it affects boys far more commonly than girls. If girls are affected, the disease is usually much less severe than in boys.

The other forms of centronuclear myopathies are less severe and are called *autosomal*, which is a reference to their inheritance pattern. The genetic mutations that underlie these myopathies are not on the X chromosome; they’re on *autosomes*, the numbered chromosomes.

**Myotubular myopathy**

**Causes:**
The most common and severe form of centronuclear myopathy is myo-

milestones slowly, and at puberty they might experience further weakening, necessitating use of a wheelchair.

For adults, even noncongenital forms of the disease can cause life-threatening respiratory problems. Adults also might experience swallowing and speech problems, and those with restricted mobility might develop scoliosis. However, even people who have had the disease since birth can lead active lives.

**Treatment:**
An infant with nemaline myopathy usually requires a feeding tube to deliver nutrition and mechanical ventilation to support respiration.
**Endocrine Myopathies**

Endocrine myopathies can occur when a gland produces too much or too little of a hormone. Hormones travel through the bloodstream and affect metabolism (a set of vital chemical reactions) in a variety of tissues, including muscle. Overproduction of thyroid hormones, known as T3 and T4, by the thyroid gland causes **hyperthyroid myopathy**, while underproduction causes **hypothyroid myopathy**.

A common cause of both endocrine myopathies is autoimmunity, a condition in which the immune system turns against part of the body — in this case, the thyroid.

Both myopathies can be almost completely alleviated by restoring normal thyroid activity, called the **euthyroid state**.

**Hyperthyroid myopathy**

*Also called:* thyrotoxic myopathy

**Symptoms:**
This disease commonly involves weakness and wasting of muscles around the shoulders and sometimes the hips. There also can be weakness in muscles of the face and throat, and in the respiratory muscles. Severe cases can cause **rhabdomyolysis** (acute muscle breakdown). Some people with hyperthyroid myopathy develop Grave’s disease, damage to muscles that control movement of the eye and eyelids, which can lead to vision loss. Others develop thyrotoxic periodic paralysis, which involves temporary, but severe attacks of muscle weakness in association with low serum potassium. If your doctor is concerned about the possibility of hypoKPP, it is important to check the thyroid function with a blood test to exclude the possibility of an endocrine myopathy (see “Periodic Paralyses,” page 12).

**Treatment:**
Restoring normal levels of the thyroid hormones can be achieved with anti-thyroid drugs, but sometimes requires partial or complete surgical removal of the thyroid (thyroidectomy).
Hypothyroid myopathy

**Symptoms:**
The most common symptoms include weakness around the hips and sometimes the shoulders, and a slowing of reflexes. Some people also experience muscle stiffness and painful muscle cramps. Severe cases can cause rhabdomyolysis (see previous page).

**Treatment:**
Thyroid hormone levels can be brought up to normal with oral medication.

Sometimes, the disease causes muscle enlargement along with muscle weakness.
tubular myopathy, which is caused by defects or deficiencies of myotubularin, a protein thought to promote normal muscle development.

**Inheritance:**
X-linked

**Onset:**
congenital (present at birth)

**Symptoms:**
X-linked myotubular myopathy usually affects only boys, and causes severe muscle weakness and hypotonia noticeable at birth and sometimes before. The weakness and hypotonia interfere with posture and movement, and cause life-threatening difficulties with feeding and respiration. Sometimes, failure or infection of the lungs causes death in early infancy, but others survive into childhood. Usually, these boys require a feeding tube and assisted ventilation.

Contractures (joints frozen in place because of muscle weakness) may develop, particularly in the hips and knees. Spinal curvature (scoliosis) may develop in childhood.

**Treatment:**
Until recently, nearly all infants with X-linked myotubular myopathy died within their first few months of life. But it’s now clear that intensive, continuous support of feeding and ventilation can significantly improve their life expectancy and allow a high qual-

**Autosomal centronuclear myopathies**

**Causes:**
As of 2009, two genes have been found that, when flawed, cause an autosomal form of centronuclear myopathy. One gene is for the amphiphysin 2 protein, which normally is involved in maintenance of the membrane surrounding muscle fibers. The other gene is for the dynamin 2 protein, which is part of the transportation system for substances inside cells.

**Inheritance:**
autosomal recessive (amphiphysin 2 type); autosomal dominant (dynamin 2 type)
Onset:
childhood, adolescence or adulthood; recessive type tends to have an earlier onset than the dominant type.

Symptoms:
Autosomal centronuclear myopathies are rarely fatal in childhood and do not seriously weaken the respiratory muscles the way the X-linked form does. Weakness is diffuse but generally has a preference for either the proximal (near the center of the body) or distal (away from the center of the body) muscles.

Autosomal dominant centronuclear myopathy often has a very gradual onset and may not cause a person to seek medical attention until young adulthood. It’s slowly progressive, and most people are able to walk independently well into adulthood. Drooping of the upper eyelids is common. Contractures may occur.

Autosomal recessive centronuclear myopathy tends to begin earlier and be more severe. Children typically have some neuromuscular impairment as they grow and develop. They also may develop weakness of the eye muscles. Some may have an elongated face and a high-arched palate. Contractures may occur.

Treatment:
There is no specific treatment. Supportive care, such as physical therapy to minimize contracture development, can be helpful.

How are these six inherited myopathies diagnosed?

Usually, diagnosis begins with evaluation of the patient’s personal and family history, and proceeds with physical and neurological examinations that test reflexes and strength. The exams can detect problems with muscle tone and contraction, and the histories can bring to light patterns of inheritance and conditions that might have aggravated the muscle problems in the past.

Given this information, a doctor can sometimes distinguish an inherited myopathy from other diseases that affect muscle function, such as muscular dystrophies and neurological disorders. But to accurately identify the myopathy and plan an appropriate course of treatment, the doctor can use several specialized tests:

Genetic (DNA-based) tests, usually performed on a blood sample, are available for many of the genetic mutations that underlie various myopathies.
A muscle biopsy, the removal of a small piece of muscle tissue, is sometimes performed. The sample is examined for physical signs of muscle disease. Under the microscope, muscles affected by various myopathies have fairly distinct appearances.

Also, muscle biopsy can be used to see how isolated muscles respond to different potentially harmful conditions. For example, to determine a patient’s susceptibility to malignant hyperthermia, a biopsied muscle can be tested for its reaction to potentially dangerous anesthetic drugs.

A muscle’s activity can be measured in the body by electromyography (EMG), which involves observing the electrical signals that a muscle produces during contraction. A needlelike electrode inserted into the muscle “reads” the electrical signals and sends them to a monitor called an oscilloscope. The technique usually causes some discomfort, but is useful for diagnosing channelopathies, which can show telltale abnormal signals on the oscilloscope.

What about myopathies that aren’t inherited?

MDA covers noninherited myopathies that are caused by an excess or a deficiency of hormones made by the thyroid gland, which is part of the endocrine system. These myopathies are known as endocrine myopathies. Fortunately, they can almost always be successfully treated by restoring normal levels of thyroid hormones with medication or surgery.

Assessing thyroid function is often part of the diagnostic process for a myopathy, especially if there is no family history of the condition.

Doctors frequently measure the level of thyroid-stimulating hormone (TSH) in a blood sample to determine whether the myopathy is endocrine-related or to rule out the endocrine myopathies.

Alan Beggs at Harvard University has had MDA support to study the molecular genetics of inheritable myopathies.
With MDA’s support, scientists have made significant progress toward understanding and treating the inherited myopathies.

Not long ago, many people with myopathies that cause temporary symptoms were told they had psychological problems or were accused of being lazy. MDA-funded scientists helped show that these are treatable, physiological medical conditions.

In the 1990s, researchers discovered that ion channel defects were at the root of several myopathies; that defective filament proteins give rise to nemaline myopathy; and that defects in or loss of a previously unknown protein called myotubularin cause X-linked myotubular myopathy.

The following decade, an MDA research grant to Rabi Tawil at the University of Rochester (N.Y.) helped lead to the identification of potassium channel abnormalities as the basis of Andersen-Tawil syndrome.

More recently, MDA-supported researchers have found that centronuclear myopathies encompass a larger group of diseases than just X-linked myotubular myopathy, and that these often have a better prognosis than the X-linked disease.

Today, MDA continues to support scientists in their quest to understand the molecular bases of the myopathies and to find effective treatments. As of 2009, the Association is helping to support a large-scale trial to see whether either of two drugs, acetazolamide or dichlorphenamide, helps decrease attacks of weakness in hyperkalemic and hypokalemic periodic paralysis.
The Muscular Dystrophy Association offers a vast array of services to help you and your family deal with inheritable and endocrine myopathies. Whether you’re an adult who’s just received a diagnosis, or the parent of a child with a myopathy, the staff at your local MDA office is there to assist you in many ways. The Association’s services include:

- A nationwide network of clinics staffed by top neuromuscular disease specialists
- MDA summer camps for kids with neuromuscular diseases
- Professionally facilitated support groups for those affected, spouses, parents or other caregivers
- Assistance with purchase and repair of wheelchairs, leg braces and communication devices
- Evaluations for physical, occupational, speech and respiratory therapy
- Flu shots to help protect the respiratory system
- Equipment loan closets

MDA’s public health education program helps you stay abreast of research news, medical findings and disability information, through educational speakers, seminars, videos, newsletters and more. Be sure and ask your local office for MDA’s latest brochures, including “Services for the Individual, Family and Community.” You’ll also want to get copies of “Breathe Easy: Respiratory Care in Neuromuscular Disorders,” “Learning to Live with Neuromuscular Disease: A Message for Parents” and some publications geared to children. Many MDA booklets are available in Spanish.

Everyone registered with MDA also receives Quest, MDA’s award-winning national magazine. Quest publishes detailed articles about research findings; medical and day-to-day care; helpful products and devices; psychological, social and family issues related to living with a disability; and much more.

If you have any questions about inheritable and endocrine myopathies, someone at MDA will help you find the answers.

Adam Foye, born in 2001, was at first given a diagnosis of X-linked myotubular myopathy, with an uncertain prognosis. His diagnosis has since been changed to centronuclear myopathy.
The Muscular Dystrophy Association fights neuromuscular diseases through an unparalleled worldwide research effort. The following diseases are included in MDA’s program:

**Diseases of Neuromuscular Junction**
- Myasthenia gravis
- Lambert-Eaton (myasthenic) syndrome
- Congenital myasthenic syndromes

**Diseases of Peripheral Nerve**
- Charcot-Marie-Tooth disease
- Friedreich’s ataxia
- Dejerine-Sottas disease

**Muscular Dystrophies**
- Myotonic dystrophy
  
  (*Steinert disease*)
- Duchenne muscular dystrophy
- Becker muscular dystrophy
- Limb-girdle muscular dystrophy
- Facioscapulohumeral muscular dystrophy
- Congenital muscular dystrophy
- Oculopharyngeal muscular dystrophy
- Distal muscular dystrophy
- Emery-Dreifuss muscular dystrophy

**Motor Neuron Diseases**
- Amyotrophic lateral sclerosis (ALS)
- Infantile progressive spinal muscular atrophy
  
  (*Type 1, Werdnig-Hoffmann disease*)
- Intermediate spinal muscular atrophy
  
  (*Type 2*)
- Juvenile spinal muscular atrophy
  
  (*Type 3, Kugelberg-Welander disease*)
- Adult spinal muscular atrophy
  
  (*Type 4*)
- Spinal-bulbar muscular atrophy
  
  (*Kennedy disease*)

**Metabolic Diseases of Muscle**
- Phosphorylase deficiency
  
  (*McArdle disease*)
- Acid maltase deficiency
  
  (*Pompe disease*)
- Phosphofructokinase deficiency
  
  (*Tarui disease*)
- Debrancher enzyme deficiency
  
  (*Cori or Forbes disease*)
- Mitochondrial myopathy
- Carnitine deficiency
- Carnitine palmityl transferase deficiency
- Phosphoglycerate kinase deficiency
- Phosphoglycerate mutase deficiency
- Lactate dehydrogenase deficiency
- Myoadenylate deaminase deficiency

**Myopathies Due to Endocrine Abnormalities**
- Hyperthyroid myopathy
- Hypothyroid myopathy

**Other Myopathies**
- Myotonia congenita
- Paramyotonia congenita
- Central core disease
- Nemaline myopathy
- Myotubular myopathy
- Periodic paralysis

**Inflammatory Myopathies**
- Polymyositis
- Dermatomyositis
- Inclusion-body myositis
MDA’s Web site is constantly updated with the latest information about the diseases in its program. Go to www.mda.org.