

# 12 Muscle

## Mechanisms of Contraction and Neural Control

### Objectives

After studying this chapter, you should be able to . . .

1. describe the gross and microscopic structure of skeletal muscles.
2. describe the nature of a muscle twitch and explain how summation and tetanus are produced.
3. distinguish between isometric and isotonic contractions.
4. explain how the series-elastic component affects muscle contraction.
5. define the term *motor unit* and explain how motor units are used to control muscle contraction.
6. describe the structure of myofibrils and explain how it accounts for the striated appearance of skeletal muscle fibers.
7. explain what is meant by the sliding filament theory of contraction.
8. list the events that occur during cross-bridge cycles and describe the role of ATP in muscle contraction.
9. explain how tropomyosin and troponin control muscle contraction and relaxation, and describe the role of  $\text{Ca}^{2+}$  and the sarcoplasmic reticulum in excitation-contraction coupling.
10. describe the structure and functions of muscle spindles and explain the mechanisms involved in a stretch reflex.
11. describe the function of Golgi tendon organs and explain why a slow, gradual muscle stretch could avoid the spasm that may result from a rapid stretch.
12. explain what is meant by reciprocal innervation and describe the neural pathways involved in a crossed-extensor reflex.
13. explain the significance of gamma motoneurons in the neural control of muscle contraction and in the maintenance of muscle tone.
14. describe the neural pathways involved in the pyramidal and extrapyramidal systems.
15. explain the significance of the maximal oxygen uptake and describe the function of phosphocreatine in muscles.
16. explain how slow-twitch, fast-twitch, and intermediate fibers differ in structure and function.
17. describe skeletal muscle metabolism during exercise, and explain how muscles fatigue and how muscle fibers change as a result of physical training.
18. compare cardiac muscle and skeletal muscle in terms of structure and physiology.
19. describe the structure of smooth muscle and explain how its contraction is regulated.

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## Refresh Your Memory

Before you begin this chapter, you may want to review the following concepts from previous chapters:

- Cytoplasm and Its Organelles 56
- Glycolysis and the Lactic Acid Pathway 102
- Aerobic Respiration 107
- Electrical Activity in Axons 158
- Acetylcholine as a Neurotransmitter 169

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Visit the Online Learning Center (text website) and use the Essential Study Partner (ESP) CDs included with this text as additional study resources.

- Learning objectives
- Objective question quizzes
- Essay question quizzes
- ESP CD #1, Support and Movement: Muscular System
- Case studies
- Related websites
- Selected chapter references

[www.mhhe.com/fox](http://www.mhhe.com/fox)

Maria, an energetic 40-year-old who plays softball and has been active in athletics most of her life, complains that she is experiencing fatigue and muscle pain and that her body just doesn't seem as limber as it should. Upon exercise testing, she is found to have a high maximal oxygen uptake. Her muscles, though not large, are well toned—perhaps excessively so. Laboratory tests reveal a normal blood level of creatine phosphokinase but an elevated blood  $\text{Ca}^{2+}$  concentration. She has hypertension, which is well controlled with a calcium channel-blocking drug.

What might be responsible for Maria's fatigue and muscle pain?

**Table 12.1** Skeletal Muscle Actions

Category	Action
Extensor	Increases the angle at a joint
Flexor	Decreases the angle at a joint
Abductor	Moves limb away from the midline of the body
Adductor	Moves limb toward the midline of the body
Levator	Moves insertion upward
Depressor	Moves insertion downward
Rotator	Rotates a bone along its axis
Sphincter	Constricts an opening

## Skeletal Muscles

Skeletal muscles are composed of individual muscle fibers that contract when stimulated by a motor neuron. Each motor neuron branches to innervate a number of muscle fibers, and all of these fibers contract when their motor neuron is activated. Activation of varying numbers of motor neurons, and thus varying numbers of muscle fibers, results in gradations in the strength of contraction of the whole muscle.

Skeletal muscles are usually attached to bone on each end by tough connective tissue tendons. When a muscle contracts, it shortens, and this places tension on its tendons and attached bones. The muscle tension causes movement of the bones at a joint, where one of the attached bones generally moves more than the other. The more movable bony attachment of the muscle, known as its *insertion*, is pulled toward its less movable attachment known as its *origin*. A variety of skeletal movements are possible, depending on the type of joint involved and the attachments of the muscles (table 12.1 and fig. 12.1). When *flexor muscles* contract, for example, they decrease the angle of a joint. Contraction of *extensor muscles* increases the angle of their attached bones at the joint. The prime mover of any skeletal movement is called the **agonist muscle**; in flexion, for example, the flexor is the agonist muscle. Flexors and extensors that act on the same joint to produce opposite actions are **antagonistic muscles**.

The position of the limbs, for example, is determined by the actions of a variety of antagonistic muscles. In addition to the movements of flexion and extension, a limb can be moved away from the midline of the body by contraction of *abductor muscles*, and it can be brought inward toward the midline by contraction of *adductor muscles*. In all cases, these skeletal movements are produced by the shortening of the appropriate muscle groups—the agonists—while the antagonist muscles remain relaxed.

## Structure of Skeletal Muscles

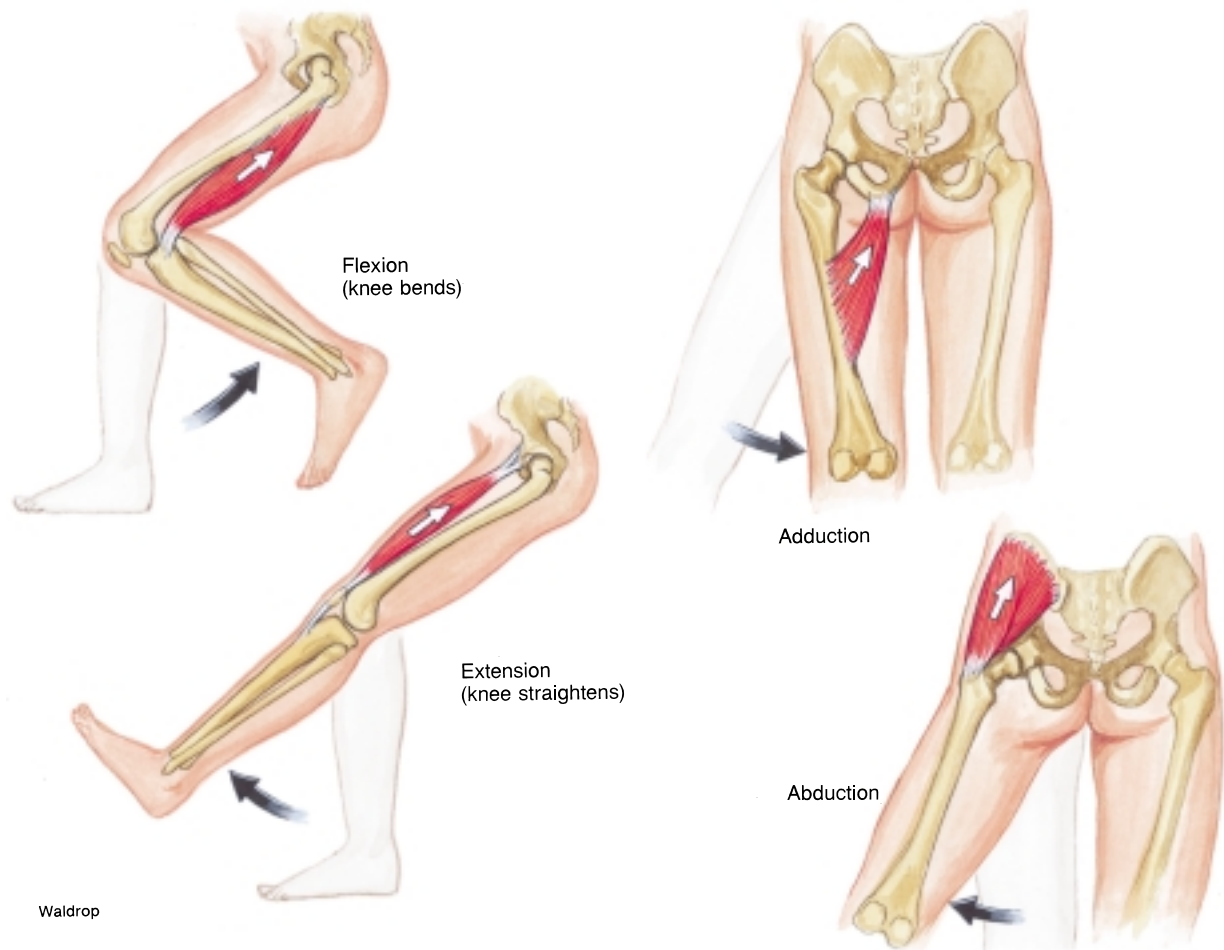
The fibrous connective tissue proteins within the tendons extend around the muscle in an irregular arrangement, forming a sheath known as the *epimysium* (*epi* = above; *my* = muscle). Connective tissue from this outer sheath extends into the body of the muscle, subdividing it into columns, or *fascicles* (these are the “strings” in stringy meat). Each of these fascicles is thus surrounded by its own connective tissue sheath, which is known as the *perimysium* (*peri* = around).

Dissection of a muscle fascicle under a microscope reveals that it, in turn, is composed of many **muscle fibers**, or *myofibers*. Each is surrounded by a cell membrane, or **sarcolemma**, enveloped by a thin connective tissue layer called an *endomysium* (fig. 12.2). Since the connective tissue of the tendons, epimysium, perimysium, and endomysium is continuous, muscle fibers do not normally pull out of the tendons when they contract.



**Duchenne's muscular dystrophy** is the most severe of the muscular dystrophies, afflicting 1 out of 3,500 boys each year. This disease, inherited as an X-linked recessive trait, involves progressive muscular

wasting and usually results in death by the age of 20. The product of the defective gene is a protein named *dystrophin*, which is associated with the plasma membrane of skeletal muscle fibers (the sarcolemma). Using this information, scientists have recently developed laboratory tests that can detect this disease in fetal cells obtained by amniocentesis. This research has been aided by the development of a strain of mice that exhibit an equivalent form of the disease. When the “good genes” for dystrophin are inserted into mouse embryos of this strain, the mice do not develop the disease. Insertion of the gene into large numbers of mature muscle cells, however, is more difficult, and so far has met with only limited success.



■ **Figure 12.1** Actions of antagonistic muscles that move the thigh and leg. Muscle contraction and shortening is responsible for all movements of the skeleton.

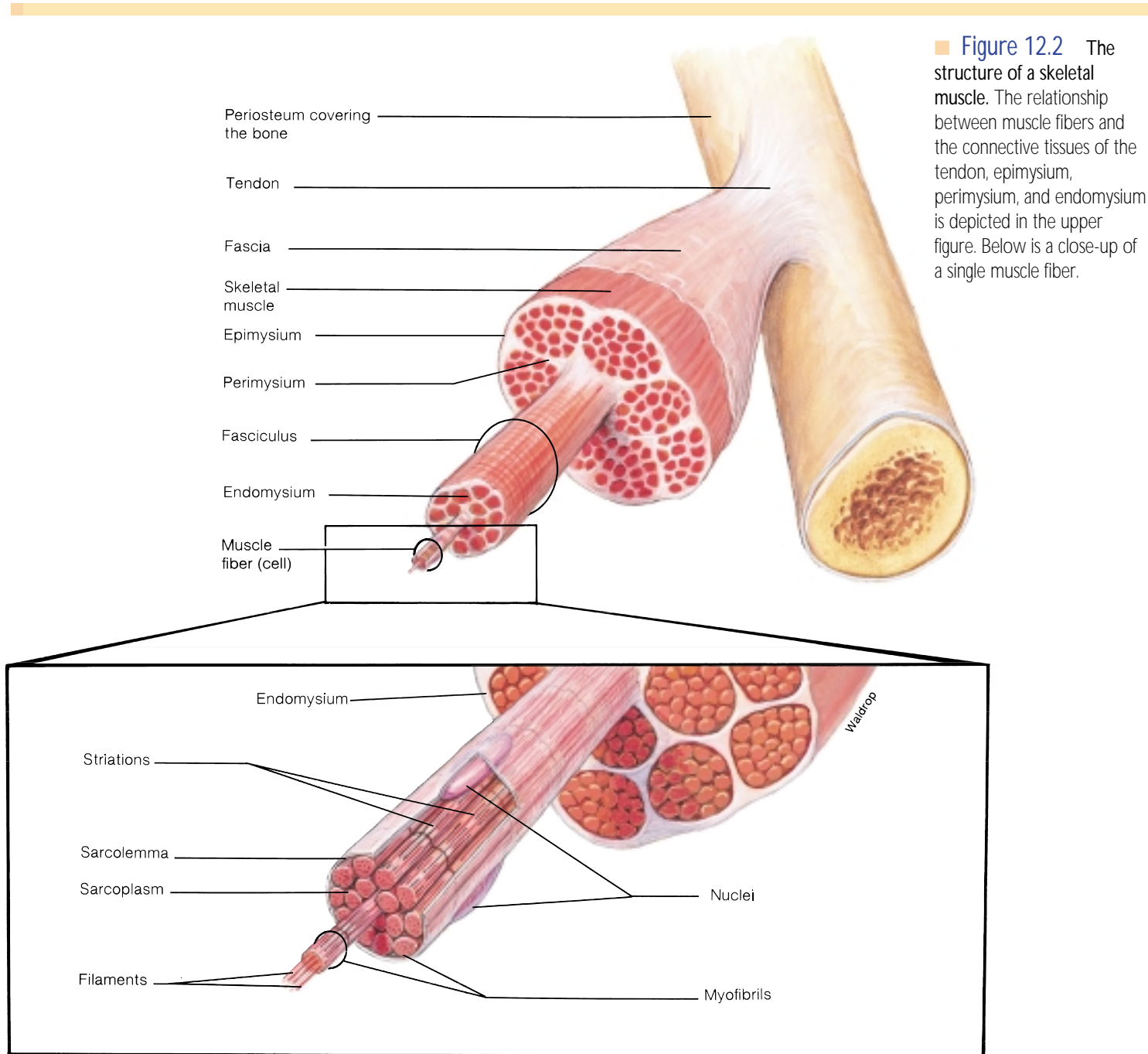
Despite their unusual elongated shape, muscle fibers have the same organelles that are present in other cells: mitochondria, intracellular membranes, glycogen granules, and others. Unlike most other cells in the body, skeletal muscle fibers are multinucleate—that is, they contain multiple nuclei. This is because, as described in chapter 1, each muscle fiber is a syncytial structure. That is, each muscle fiber is formed from the union of several embryonic myoblast cells. The most distinctive feature of skeletal muscle fibers, however, is their **striated** appearance when viewed microscopically (fig. 12.3). The striations (stripes) are produced by alternating dark and light bands that appear to span the width of the fiber.

The dark bands are called **A bands**, and the light bands are called **I bands**. At high magnification in an electron microscope, thin dark lines can be seen in the middle of the I bands. These are called **Z lines**. The labels A, I, and Z—derived in the

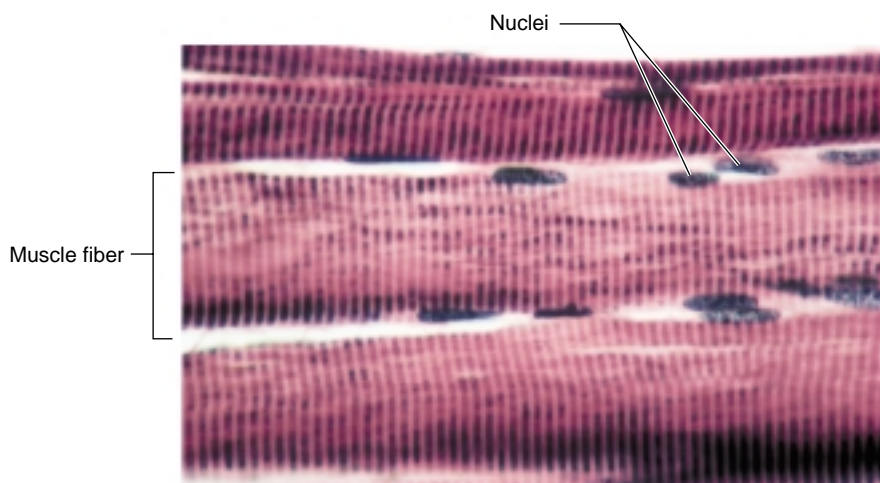
course of early muscle research—are useful for describing the functional architecture of muscle fibers. The letters *A* and *I* stand for *anisotropic* and *isotropic*, respectively, which indicate the behavior of polarized light as it passes through these regions; the letter *Z* comes from the German word *Zwischen-scheibe*, which translates to “between disc.” These derivations are of historical interest only.

## Motor Units

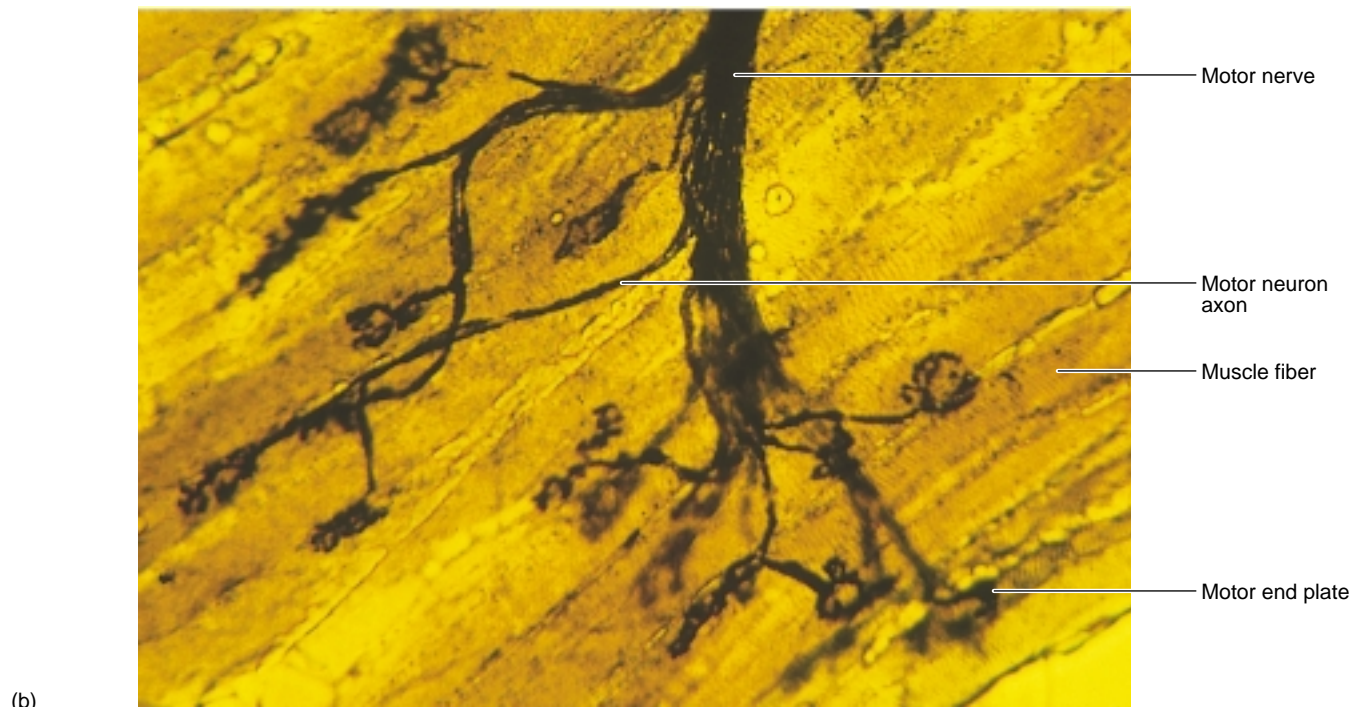
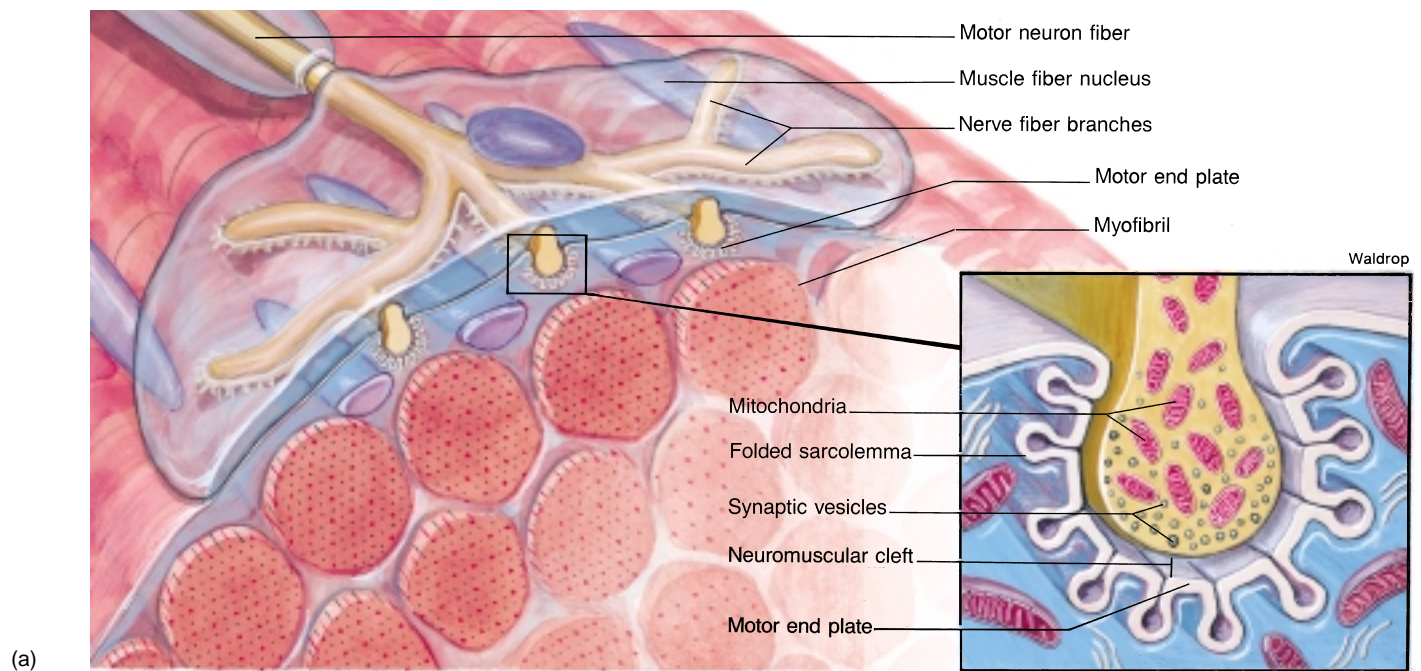
In vivo, each muscle fiber receives a single axon terminal from a somatic motor neuron. The motor neuron stimulates the muscle fiber to contract by liberating acetylcholine at the neuromuscular junction (described in chapter 7). The specialized region of the sarcolemma of the muscle fiber at the neuromuscular junction is known as a **motor end plate** (fig. 12.4).



■ **Figure 12.2** The structure of a skeletal muscle. The relationship between muscle fibers and the connective tissues of the tendon, epimysium, perimysium, and endomysium is depicted in the upper figure. Below is a close-up of a single muscle fiber.



■ **Figure 12.3** The appearance of skeletal muscle fibers through the light microscope. The striations are produced by alternating dark A bands and light I bands. (Note the peripheral location of the nuclei.)



**Figure 12.4** Motor end plates at the neuromuscular junction. The neuromuscular junction is the synapse between the nerve fiber and muscle fiber. The motor end plate is the specialized portion of the sarcolemma of a muscle fiber surrounding the terminal end of the axon. (a) An illustration of the neuromuscular junction. Notice the slight gap between the membrane of the axon and that of the muscle fiber. (b) A photomicrograph of muscle fibers and neuromuscular junctions.



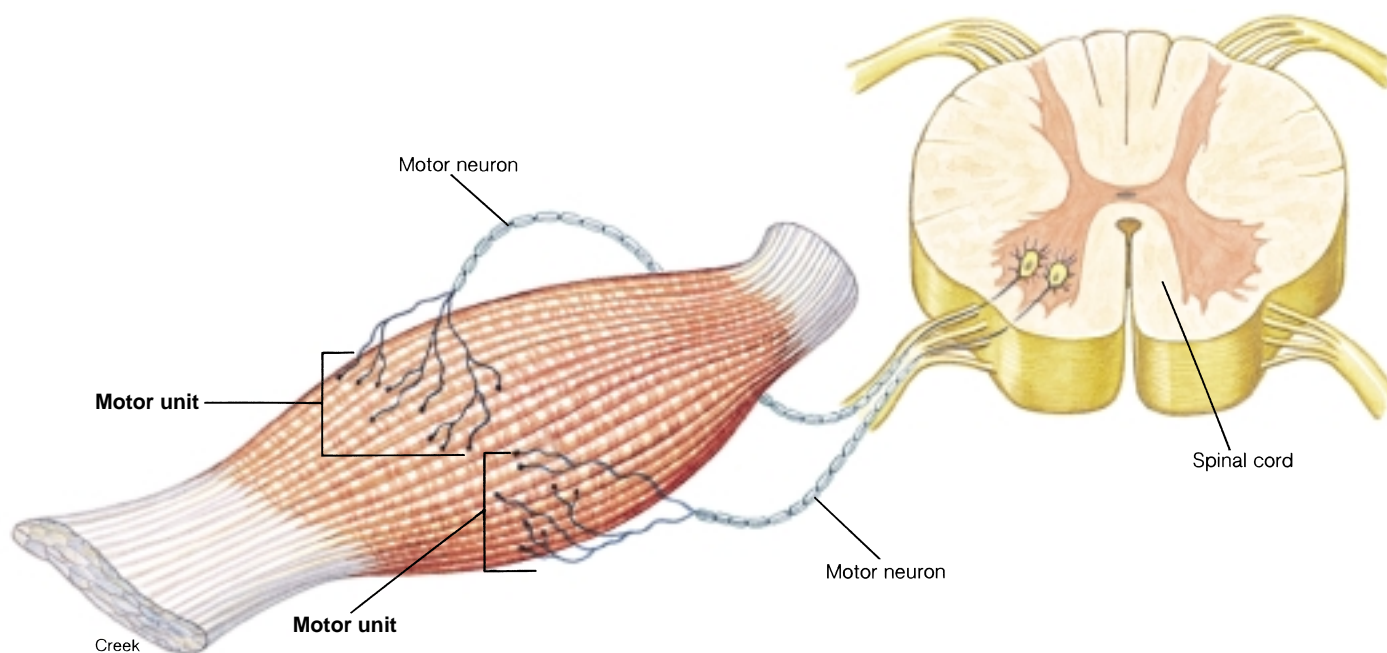
The acetylcholine (ACh) released by the axon terminals diffuses across the synaptic cleft and binds to ACh receptors in the plasma membrane of the end plate, thereby stimulating the muscle fiber. Prior to its release, the ACh is contained in synaptic vesicles that dock and fuse with the plasma membrane of the axon terminal and undergo exocytosis (see chapter 7, fig. 7.21). The potentially deadly **botulinum toxin**, produced by the bacteria *Clostridium botulinum*, is selectively taken into cholinergic nerve endings and cleaves the proteins needed for the exocytosis of the synaptic vesicles. This blocks nerve stimulation of the muscles, producing a flaccid paralysis. Interestingly, botulinum toxin is now used medically in certain cases to relieve muscle spasms due to excessive nerve stimulation. For example, it is injected into an affected extraocular muscle in order to help correct **strabismus** (deviation of the eye).

The cell body of a somatic motor neuron is located in the ventral horn of the gray matter of the spinal cord and gives rise to a single axon that emerges in the ventral root of a spinal nerve (chapter 8). Each axon, however, can produce a number of collateral branches to innervate an equal number of muscle fibers. Each somatic motor neuron, together with all of the muscle fibers that it innervates, is known as a **motor unit** (fig. 12.5).

Whenever a somatic motor neuron is activated, all of the muscle fibers that it innervates are stimulated to contract. In vivo, graded contractions of whole muscles are produced by variations in the number of motor units that are activated. In order for these graded contractions to be smooth and sustained, different motor units must be activated by rapid, asynchronous stimulation.

Fine neural control over the strength of muscle contraction is optimal when there are many small motor units involved. In the extraocular muscles that position the eyes, for example, the *innervation ratio* (motor neuron:muscle fibers) of an average motor unit is one neuron per twenty-three muscle fibers. This affords a fine degree of control. The innervation ratio of the gastrocnemius, by contrast, averages one neuron per thousand muscle fibers. Stimulation of these motor units results in more powerful contractions at the expense of finer gradations in contraction strength.

All of the motor units controlling the gastrocnemius, however, are not the same size. Innervation ratios vary from 1:100 to 1:2,000. A neuron that innervates fewer muscle fibers has a smaller cell body and is stimulated by lower levels of excitatory input than a larger neuron that innervates a greater number of muscle fibers. The smaller motor units, as a result, are the ones that are used most often. When contractions of greater strength are required, larger and larger motor units are activated in a process known as **recruitment** of motor units.



■ **Figure 12.5** **Motor units.** A motor unit consists of a motor neuron and the muscle fibers it innervates. This diagram illustrates the innervation of muscle fibers by different motor units. (Actually, many more muscle fibers would be included in a single motor unit than are shown here.)

## Test Yourself Before You Continue

1. Describe the actions of muscles when they contract, and define the terms *agonist* and *antagonist* in muscle action.
2. Describe the different levels of muscle structure, explaining how the muscle and its substructures are packaged in connective tissues.
3. Define the terms *motor unit* and *innervation ratio* as they relate to muscle function, and draw a simple diagram of a motor unit with a 1:5 innervation ratio.
4. Using the concept of recruitment, explain how muscle contraction can be graded in its strength.

## Mechanisms of Contraction

The A bands within each muscle fiber are composed of thick filaments and the I bands contain thin filaments. Movement of cross bridges that extend from the thick to the thin filaments causes sliding of the filaments, and thus muscle tension and shortening. The activity of the cross bridges is regulated by the availability of  $\text{Ca}^{2+}$ , which is increased by electrical stimulation of the muscle fiber. Electrical stimulation produces contraction of the muscle through the binding of  $\text{Ca}^{2+}$  to regulatory proteins within the thin filaments.

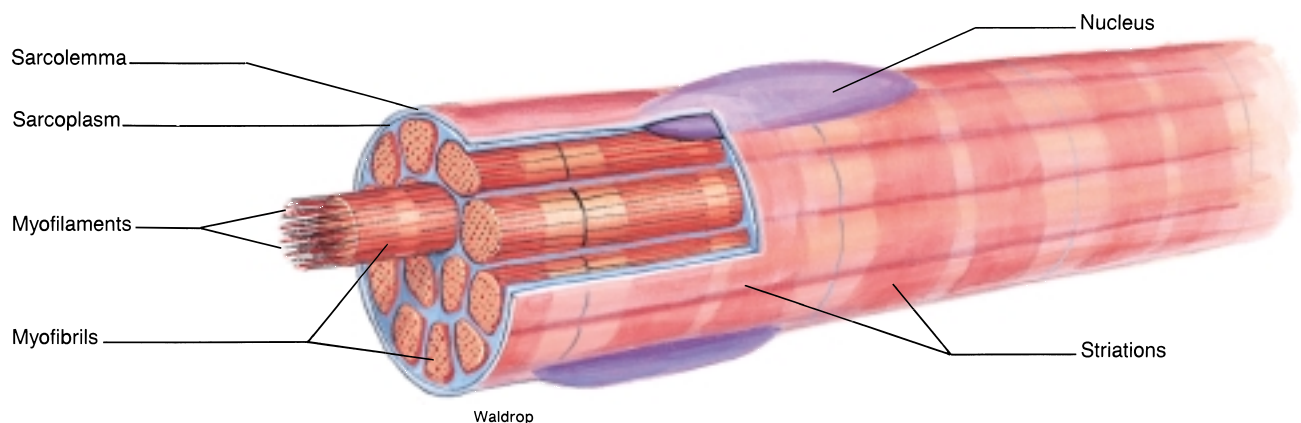
When muscle cells are viewed in the electron microscope, which can produce images at several thousand times the magnification possible in an ordinary light microscope, each cell is seen

to be composed of many subunits known as **myofibrils** (*fibrils* = little fibers) (fig. 12.6). These myofibrils are approximately 1 micrometer ( $1\ \mu\text{m}$ ) in diameter and extend in parallel rows from one end of the muscle fiber to the other. The myofibrils are so densely packed that other organelles, such as mitochondria and intracellular membranes, are restricted to the narrow cytoplasmic spaces that remain between adjacent myofibrils.

With the electron microscope, it can be seen that the muscle fiber does not have striations that extend from one side of the fiber to the other. It is the myofibrils that are striated with dark *A bands* and light *I bands* (fig. 12.7). The striated appearance of the entire muscle fiber when seen with a light microscope is an illusion created by the alignment of the dark and light bands of the myofibrils from one side of the fiber to the other. Since the separate myofibrils are not clearly seen at low magnification, the dark and light bands appear to be continuous across the width of the fiber.

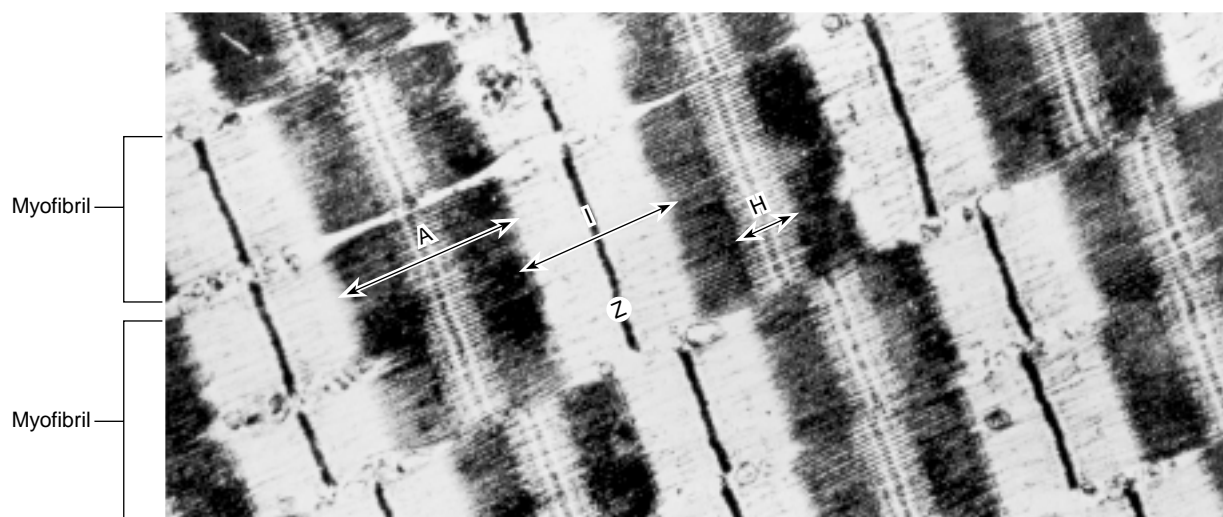
Each myofibril contains even smaller structures called **myofilaments**, or simply **filaments**. When a myofibril is observed at high magnification in longitudinal section (side view), the A bands are seen to contain **thick filaments**. These are about 110 angstroms thick ( $110\ \text{\AA}$ , where  $1\ \text{\AA} = 10^{-10}\ \text{m}$ ) and are stacked in register. It is these thick filaments that give the A band its dark appearance. The lighter I band, by contrast, contains **thin filaments** (from 50 to 60  $\text{\AA}$  thick). The thick filaments are primarily composed of the protein **myosin**, and the thin filaments are primarily composed of the protein **actin**.

The I bands within a myofibril are the lighter areas that extend from the edge of one stack of thick filaments to the edge of the next stack of thick filaments. They are light in appearance because they contain only thin filaments. The thin filaments, however, do not end at the edges of the I bands. Instead, each thin filament extends partway into the A bands on each side (between the stack of thick filaments on each side of an I band). Since thick and thin filaments overlap at the edges of each A



■ **Figure 12.6** The components of a skeletal muscle fiber. A skeletal muscle fiber is composed of numerous myofibrils that contain myofilaments of actin and myosin. Overlapping of the myofilaments produces a striated appearance. Each skeletal muscle fiber is multinucleated.





■ **Figure 12.7** An electron micrograph of a longitudinal section of myofibrils. The A, H, and I bands are clearly seen. Notice how the dark and light bands of each myofibril are stacked in register.

band, the edges of the A band are darker in appearance than the central region. These central lighter regions of the A bands are called the *H bands* (for *helle*, a German word meaning “bright”). The central H bands thus contain only thick filaments that are not overlapped by thin filaments.

In the center of each I band is a thin dark Z line. The arrangement of thick and thin filaments between a pair of Z lines forms a repeating pattern that serves as the basic subunit of striated muscle contraction. These subunits, from Z to Z, are known as **sarcomeres** (fig. 12.8a). A longitudinal section of a myofibril thus presents a side view of successive sarcomeres.

This side view is, in a sense, misleading; there are numerous sarcomeres within each myofibril that are out of the plane of the section (and out of the picture). A better appreciation of the three-dimensional structure of a myofibril can be obtained by viewing the myofibril in cross section. In this view, it can be seen that the Z lines are actually **Z discs**, and that the thin filaments that penetrate these Z discs surround the thick filaments in a hexagonal arrangement (fig. 12.8b, right). If we concentrate on a single row of dark thick filaments in this cross section, the alternating pattern of thick and thin filaments seen in longitudinal section becomes apparent.

## Sliding Filament Theory of Contraction

When a muscle contracts it decreases in length as a result of the shortening of its individual fibers. Shortening of the muscle fibers, in turn, is produced by shortening of their myofibrils, which occurs as a result of the shortening of the distance from Z line to Z line. As the sarcomeres shorten in length, however, the A bands do *not* shorten but instead move closer together. The I

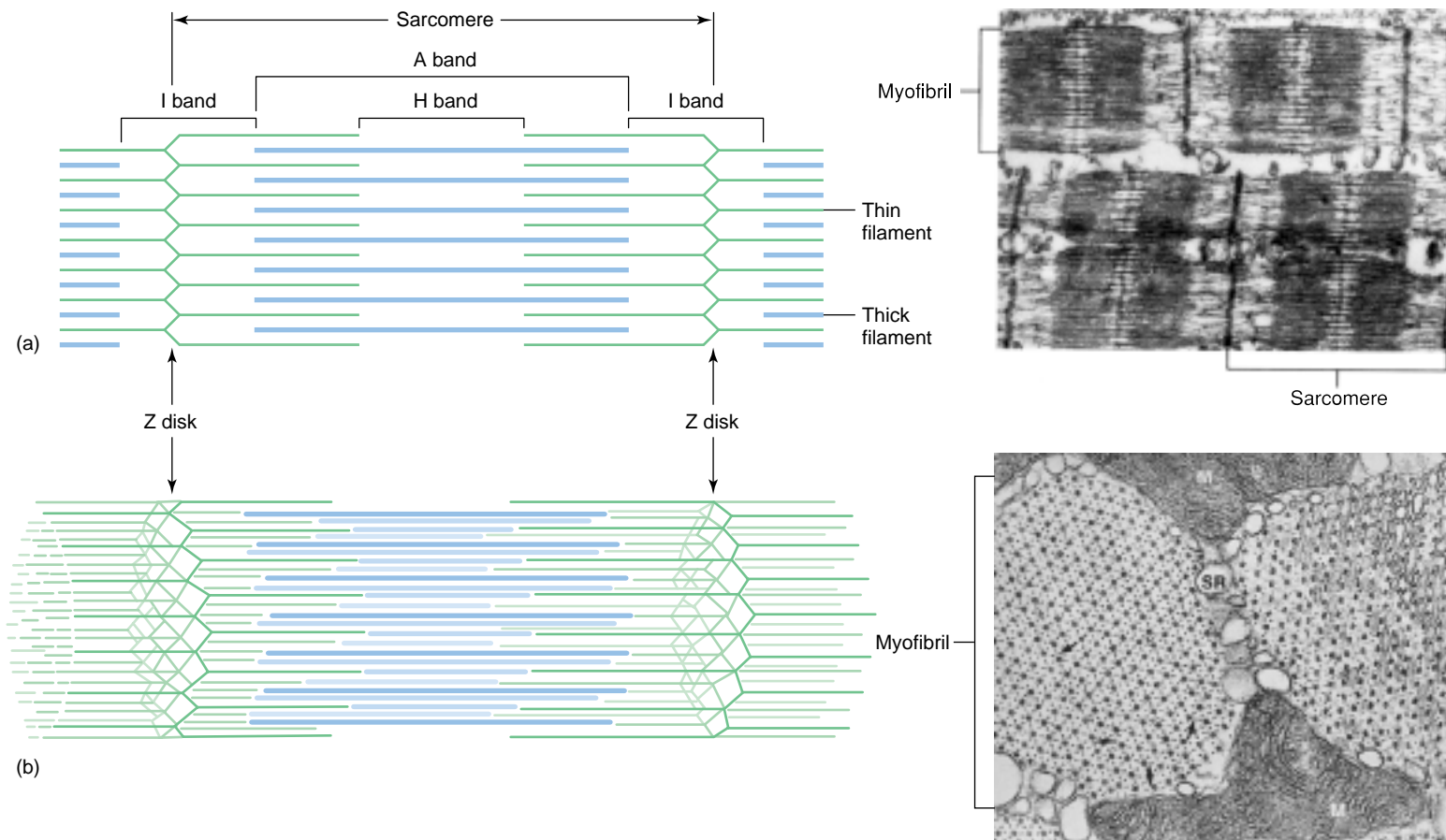
bands—which represent the distance between A bands of successive sarcomeres—decrease in length (table 12.2).

The thin filaments composing the I band, however, do not shorten. Close examination reveals that the thick and thin filaments remain the same length during muscle contraction. Shortening of the sarcomeres is produced not by shortening of the filaments, but rather by the *sliding* of thin filaments over and between the thick filaments. In the process of contraction, the thin filaments on either side of each A band slide deeper and deeper toward the center, producing increasing amounts of overlap with the thick filaments. The I bands (containing only thin filaments) and H bands (containing only thick filaments) thus get shorter during contraction (fig. 12.9).

### Cross Bridges

Sliding of the filaments is produced by the action of numerous **cross bridges** that extend out from the myosin toward the actin. These cross bridges are part of the myosin proteins that extend from the axis of the thick filaments to form “arms” that terminate in globular “heads” (fig. 12.10). A myosin protein has two globular heads that serve as cross bridges. The orientation of the myosin heads on one side of a sarcomere is opposite to that of the other side, so that, when the myosin heads form cross bridges by attaching to actin on each side of the sarcomere, they can pull the actin from each side toward the center.

Isolated muscles are easily stretched (although this is opposed in the body by the stretch reflex, described in a later section), demonstrating that the myosin heads are not attached to actin when the muscle is at rest. Each globular myosin head of a cross bridge contains an ATP-binding site closely associated with an actin-binding site (fig. 12.10). The globular heads function as **myosin ATPase** enzymes, splitting ATP into ADP and P<sub>i</sub>.



**Figure 12.8** Arrangement of thick and thin filaments in a striated muscle fiber. (a) In a longitudinal section, the thick and thin filaments are seen to form repeating units called sarcomeres. The banding patterns of the sarcomeres are labeled I, A, and H, as shown. A corresponding electron micrograph (53,000 $\times$ ) is shown to the right of the illustration. (b) The three-dimensional structure of the sarcomeres is illustrated. This three-dimensional structure can be seen in a cross section of a myofibril taken through a region of overlapping thick and thin filaments. In the electron micrograph, the arrows point to cross bridges between the thick filaments (dark dots) and thin filaments (light dots). (SR = sarcoplasmic reticulum; M = mitochondria).

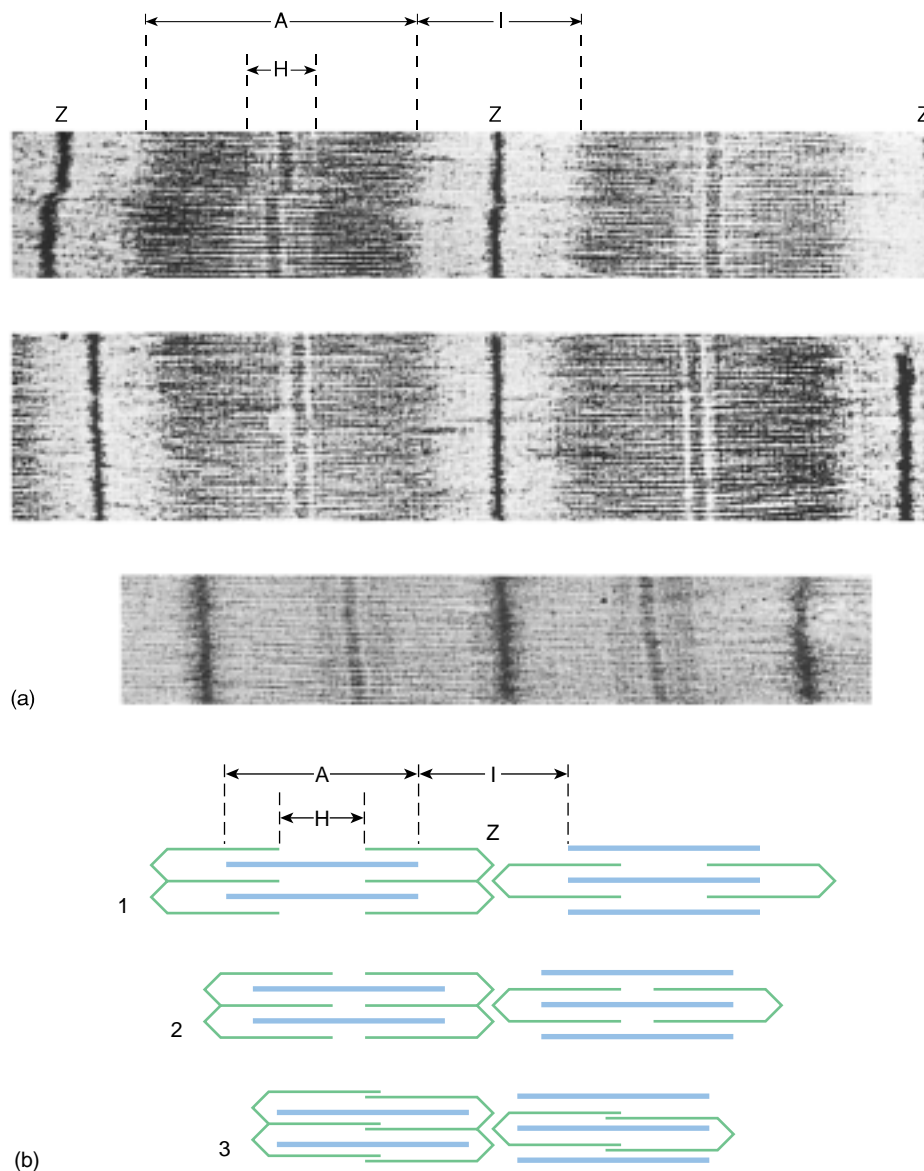
Electron micrographs (right) from R. G. Kessel and R. H. Kardon, *Tissues and Organs: A Text-Atlas of Scanning Electron Microscopy*, 1979, W. H. Freeman & Company.

## Table 12.2 Summary of the Sliding Filament Theory of Contraction

1. A myofiber, together with all its myofibrils, shortens by movement of the insertion toward the origin of the muscle.
2. Shortening of the myofibrils is caused by shortening of the sarcomeres—the distance between Z lines (or discs) is reduced.
3. Shortening of the sarcomeres is accomplished by sliding of the myofilaments—the length of each filament remains the same during contraction.
4. Sliding of the filaments is produced by asynchronous power strokes of myosin cross bridges, which pull the thin filaments (actin) over the thick filaments (myosin).
5. The A bands remain the same length during contraction, but are pulled toward the origin of the muscle.
6. Adjacent A bands are pulled closer together as the I bands between them shorten.
7. The H bands shorten during contraction as the thin filaments on the sides of the sarcomeres are pulled toward the middle.

This reaction occurs before the myosin heads combine with actin, and indeed is required for activating the myosin heads so that they can attach to actin. The ADP and  $P_i$  remain bound to the myosin heads until the cross bridges attach to the actin.

The myosin heads are able to bind to specific attachment sites in the actin subunits. When the cross bridges bind to actin, they release the  $P_i$ . This causes a conformational change in the myosin protein, resulting in a *power stroke* that pulls the thin filaments toward the center of the A bands. The ADP is released when the cross bridges bind to a fresh ATP at the end of the power stroke. This release of ADP upon bonding to a new ATP is required in order for the cross bridges to break their bond with actin at the end of the power stroke. The myosin ATPase will then split ATP and become activated as in the previous cycle. Note that the splitting of ATP is required *before* a cross bridge can attach to actin and undergo a power stroke, and that the attachment of a *new* ATP is needed for the cross bridge to release from actin at the end of a power stroke (fig. 12.11).



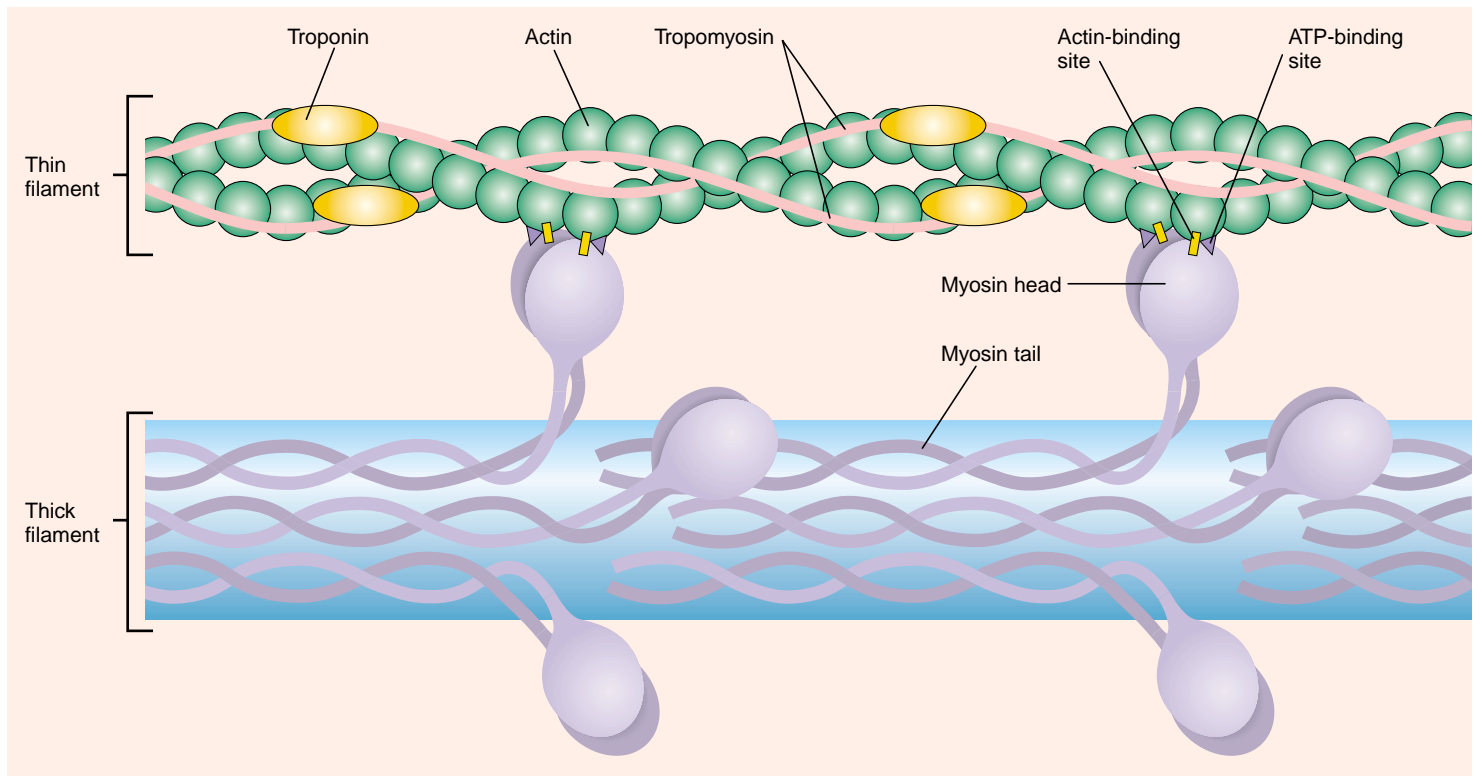
**Figure 12.9** The sliding filament model of muscle contraction. (a) An electron micrograph and (b) a diagram of the sliding filament model of contraction. As the filaments slide, the Z lines are brought closer together and the sarcomeres get shorter. (1) Relaxed muscle; (2) partially contracted muscle; (3) fully contracted muscle.



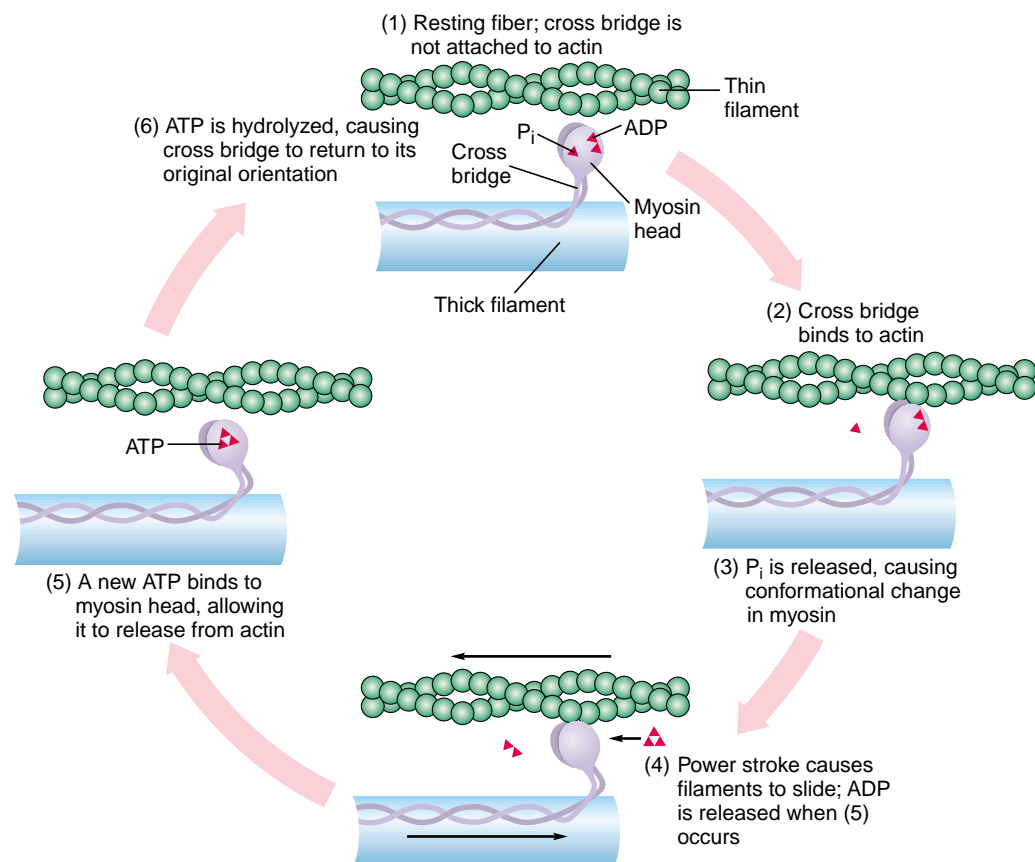
The detachment of a cross bridge from actin at the end of a power stroke requires that a new ATP molecule bind to the myosin ATPase. The importance of this process is illustrated by the muscular contracture called **rigor mortis** that occurs due to lack of ATP when the muscle dies. Without ATP, the ADP remains bound to the cross bridges, and the cross bridges remain tightly bound to actin. This results in the formation of “rigor complexes” between myosin and actin that cannot detach. In rigor mortis, the muscles remain stiff until the myosin and actin begin to decompose.

Because the cross bridges are quite short, a single contraction cycle and power stroke of all the cross bridges in a muscle would shorten the muscle by only about 1% of its resting length. Since muscles can shorten up to 60% of their resting lengths, it is obvious that the contraction cycles must be repeated many times. In order for this to occur, the cross bridges must detach from the actin at the end of a power stroke, reassume their resting orientation, and then reattach to the actin and repeat the cycle.

During normal contraction, however, only a portion of the cross bridges are attached at any given time. The power strokes are thus not in synchrony, as the strokes of a competitive rowing team would be. Rather, they are like the actions of a team engaged



■ **Figure 12.10** The structure of myosin, showing its binding sites for ATP and actin. The myosin heads can bind only to actin when a muscle has been stimulated to contract.



■ **Figure 12.11** The cross-bridge cycle that causes sliding of the filaments and muscle contraction. Hydrolysis of ATP is required for activation of the cross bridge, and the binding of a new ATP is required for the cross bridge to release from the actin at the end of a cycle.

in tug-of-war, where the pulling action of the members is asynchronous. Some cross bridges are engaged in power strokes at all times during the contraction.

## Regulation of Contraction

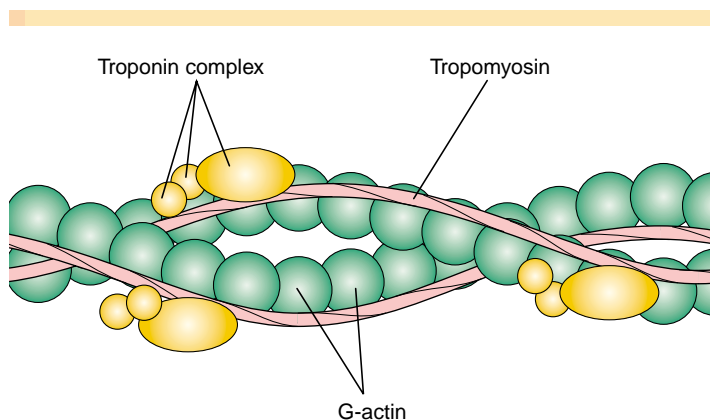
When the cross bridges attach to actin, they undergo power strokes and cause muscle contraction. In order for a muscle to relax, therefore, the attachment of myosin cross bridges to actin must be prevented. The regulation of cross-bridge attachment to actin is a function of two proteins that are associated with actin in the thin filaments.

The actin filament—or *F-actin*—is a polymer formed of 300 to 400 globular subunits (*G-actin*), arranged in a double row and twisted to form a helix (fig. 12.12). A different type of protein, known as **tropomyosin**, lies within the groove between the double row of *G-actin* monomers. There are forty to sixty tropomyosin molecules per thin filament, with each tropomyosin spanning a distance of approximately seven actin subunits.

Attached to the tropomyosin, rather than directly to the actin, is a third type of protein called **troponin** (actually a complex of three proteins—see fig. 12.12). Troponin and tropomyosin work together to regulate the attachment of cross bridges to actin, and thus serve as a switch for muscle contraction and relaxation. In a relaxed muscle, the position of the tropomyosin in the thin filaments is such that it physically blocks the cross bridges from bonding to specific attachment sites in the actin. Thus, in order for the myosin cross bridges to attach to actin, the tropomyosin must be moved. This requires the interaction of troponin with  $\text{Ca}^{2+}$ .

### Role of $\text{Ca}^{2+}$ in Muscle Contraction

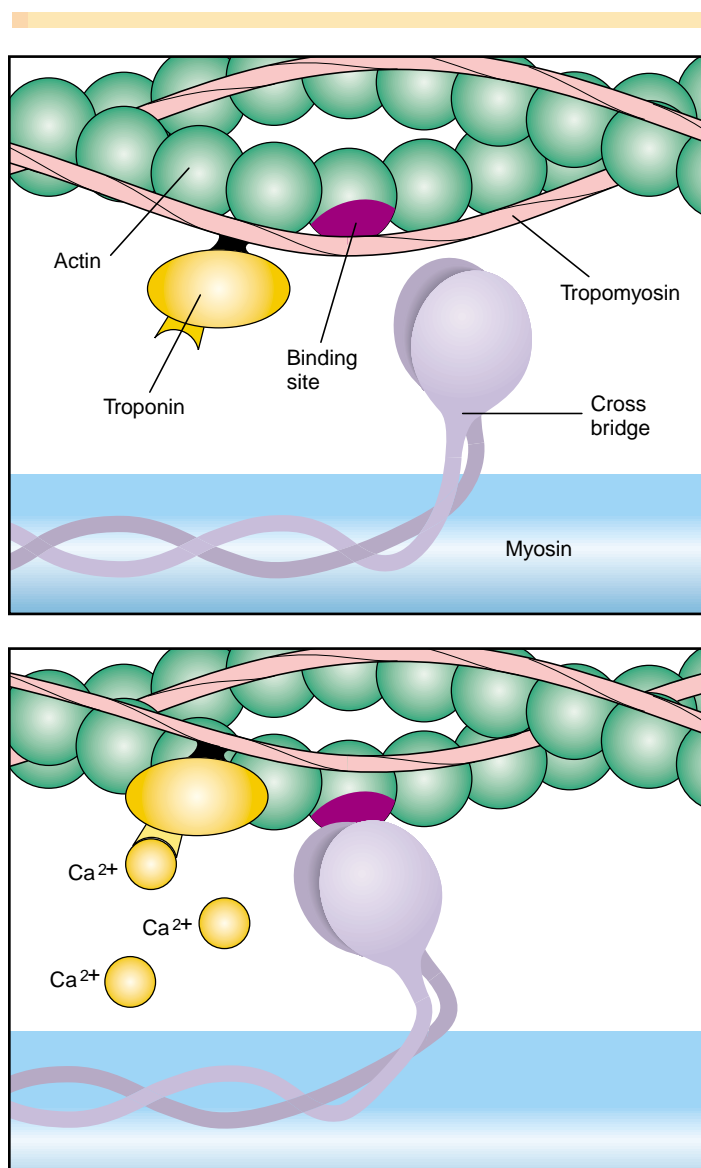
In a relaxed muscle, when tropomyosin blocks the attachment of cross bridges to actin, the concentration of  $\text{Ca}^{2+}$  in the sarcoplasm (cytoplasm of muscle cells) is very low. When the muscle cell is stimulated to contract, mechanisms that will be discussed shortly cause the concentration of  $\text{Ca}^{2+}$  in the sarcoplasm to quickly rise. Some of this  $\text{Ca}^{2+}$  attaches to troponin,



■ **Figure 12.12** The structural relationship between troponin, tropomyosin, and actin. The tropomyosin is attached to actin, whereas the troponin complex of three subunits is attached to tropomyosin (not directly to actin).

causing a conformational change that moves the troponin complex *and* its attached tropomyosin out of the way so that the cross bridges can attach to actin (fig. 12.13). Once the attachment sites on the actin are exposed, the cross bridges can bind to actin, undergo power strokes, and produce muscle contraction.

The position of the troponin-tropomyosin complexes in the thin filaments is thus adjustable. When  $\text{Ca}^{2+}$  is not attached to troponin, the tropomyosin is in a position that inhibits attachment of cross bridges to actin, preventing muscle contraction. When  $\text{Ca}^{2+}$  attaches to troponin, the troponin-tropomyosin complexes shift position. The cross bridges can then attach to actin, produce a power stroke, and detach from actin. Moreover, these contraction cycles can continue as long as  $\text{Ca}^{2+}$  is attached to troponin.



■ **Figure 12.13** The role of  $\text{Ca}^{2+}$  in muscle contraction. The attachment of  $\text{Ca}^{2+}$  to troponin causes movement of the troponin-tropomyosin complex, which exposes binding sites on the actin. The myosin cross bridges can then attach to actin and undergo a power stroke.

## Clinical Investigation Clues

Remember that Maria has muscle pain and fatigue, and that her body seems stiff. Further, she has an elevated blood concentration of  $\text{Ca}^{2+}$ .

How might the high blood  $\text{Ca}^{2+}$  be related to Maria's symptoms?

What might cause an elevated blood  $\text{Ca}^{2+}$  (hint—see chapter 11 or 19).

### Excitation-Contraction Coupling

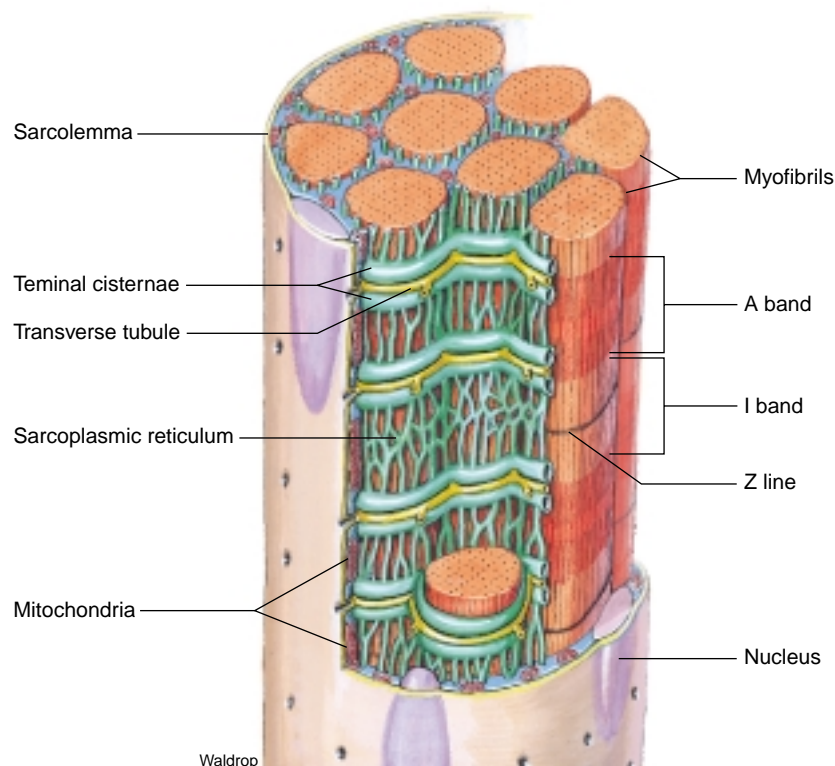
Muscle contraction is turned on when sufficient amounts of  $\text{Ca}^{2+}$  bind to troponin. This occurs when the  $\text{Ca}^{2+}$  concentration of the sarcoplasm rises above  $10^{-6}$  molar. In order for muscle relaxation to occur, therefore, the  $\text{Ca}^{2+}$  concentration of the sarcoplasm must be lowered to below this level. Muscle relaxation is produced by the active transport of  $\text{Ca}^{2+}$  out of the sarcoplasm into the **sarcoplasmic reticulum** (fig. 12.14). The sarcoplasmic reticulum is a modified endoplasmic reticulum, consisting of interconnected sacs and tubes that surround each myofibril within the muscle cell.

Most of the  $\text{Ca}^{2+}$  in a relaxed muscle fiber is stored within expanded portions of the sarcoplasmic reticulum known as *terminal cisternae*. When a muscle fiber is stimulated to contract by either a motor neuron in vivo or electric shocks in vitro, the

stored  $\text{Ca}^{2+}$  is released from the sarcoplasmic reticulum so that it can attach to troponin. When a muscle fiber is no longer stimulated, the  $\text{Ca}^{2+}$  from the sarcoplasm is actively transported back into the sarcoplasmic reticulum. Now, in order to understand how the release and uptake of  $\text{Ca}^{2+}$  is regulated, one more organelle within the muscle fiber must be described.

The terminal cisternae of the sarcoplasmic reticulum are separated only by a very narrow gap from **transverse tubules** (or **T tubules**). These are narrow membranous “tunnels” formed from and continuous with the sarcolemma (muscle cell membrane). The transverse tubules thus open to the extracellular environment through pores in the cell surface and are capable of conducting action potentials. The stage is now set to explain exactly how a motor neuron stimulates a muscle fiber to contract.

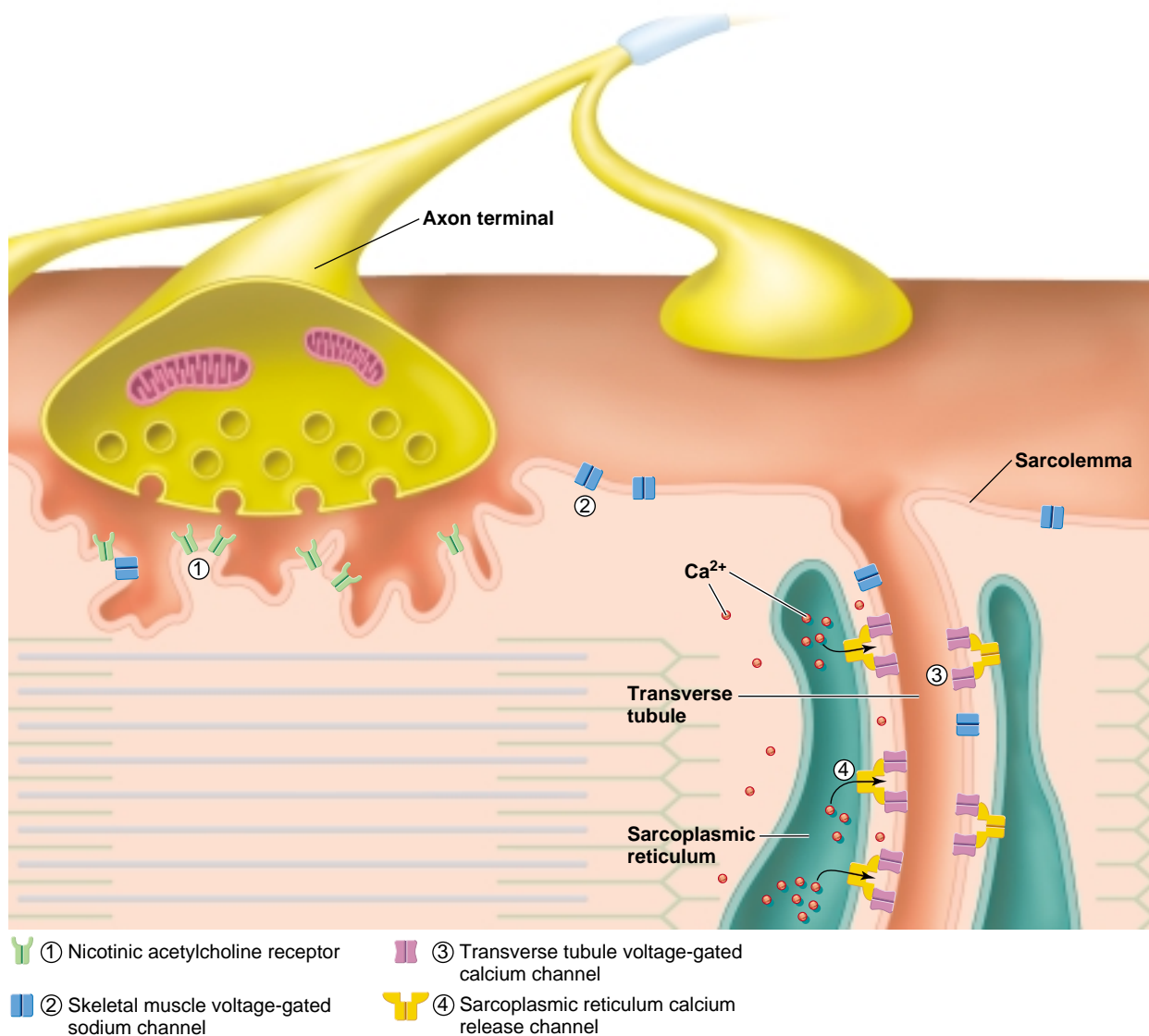
The release of acetylcholine from axon terminals at the neuromuscular junctions (motor end plates), as previously described, causes electrical activation of skeletal muscle fibers. End-plate potentials (analogous to EPSPs—chapter 7) are produced that generate action potentials. Action potentials in muscle cells, like those in nerve cells, are all-or-none events that are regenerated along the plasma membrane. It must be remembered that action potentials involve the flow of ions between the extracellular and intracellular environments across a plasma membrane that separates these two compartments. In muscle cells, therefore, action potentials can be conducted into the interior of the fiber across the membrane of the transverse tubules.



■ **Figure 12.14** The sarcoplasmic reticulum. This figure depicts the relationship between myofibrils, the transverse tubules, and the sarcoplasmic reticulum. The sarcoplasmic reticulum (green) stores  $\text{Ca}^{2+}$  and is stimulated to release it by action potentials arriving in the transverse tubules (yellow).

### Table 12.3 Summary of Events in Excitation-Contraction Coupling

1. Action potentials in a somatic motor neuron cause the release of acetylcholine neurotransmitter at the myoneural junction (one myoneural junction per myofiber).
2. Acetylcholine, through its interaction with receptors in the muscle cell membrane (sarcolemma), produces action potentials that are regenerated across the sarcolemma.
3. The membranes of the transverse tubules (T tubules) are continuous with the sarcolemma and conduct action potentials deep into the muscle fiber.
4. Action potentials in the T tubules, acting through a mechanism that is incompletely understood, stimulate the release of  $\text{Ca}^{2+}$  from the terminal cisternae of the sarcoplasmic reticulum.
5.  $\text{Ca}^{2+}$  released into the sarcoplasm attaches to troponin, causing a change in its structure.
6. The shape change in troponin causes its attached tropomyosin to shift position in the actin filament, thus exposing binding sites for the myosin cross bridges.
7. Myosin cross bridges, previously activated by the hydrolysis of ATP, attach to actin.
8. Once the previously activated cross bridges attach to actin, they undergo a power stroke and pull the thin filaments over the thick filaments.
9. Attachment of fresh ATP allows the cross bridges to detach from actin and repeat the contraction cycle as long as  $\text{Ca}^{2+}$  remains attached to troponin.
10. When action potentials stop being produced, the sarcoplasmic reticulum actively accumulates  $\text{Ca}^{2+}$  and tropomyosin returns to its inhibitory position.



■ **Figure 12.15** The structures involved in excitation-contraction coupling. The acetylcholine released from the axon binds to its nicotinic receptors in the motor end plate. This stimulates the production of a depolarization, which causes the opening of voltage-gated  $\text{Na}^+$  channels and the resulting production of action potentials along the sarcolemma. The spread of action potentials into the transverse tubules stimulates the opening of their voltage-gated  $\text{Ca}^{2+}$  channels, which (directly or indirectly) causes the opening of voltage-gated  $\text{Ca}^{2+}$  channels in the sarcoplasmic reticulum. Calcium diffuses out of the sarcoplasmic reticulum, binds to troponin, and stimulates contraction.

Action potentials in the transverse tubules cause the release of  $\text{Ca}^{2+}$  from the sarcoplasmic reticulum. This process is known as **excitation-contraction coupling** (table 12.3). Since the transverse tubules are not physically continuous with the sarcoplasmic reticulum, however, there must be some mechanism to permit communication between these two organelles. It is currently believed that there may be a direct coupling, on a molecular level, between voltage-regulated  $\text{Ca}^{2+}$  channels in the transverse tubules and the  $\text{Ca}^{2+}$  release channels in the sarcoplasmic reticulum. The  $\text{Ca}^{2+}$  release channel proteins of the sarcoplasmic reticulum have a part that extends into the cytoplasm. This part, which has a footlike appearance in the electron microscope, may be able to interact directly with the  $\text{Ca}^{2+}$  channel proteins of the transverse tubules (fig 12.15).

This arrangement has been described as an *electromechanical release* mechanism, because changes in membrane voltage (action potentials) in the transverse tubules cause a change in protein conformation of calcium channels, which are mechanically linked to other calcium channels in the sarcoplasmic reticulum. There is also evidence that the  $\text{Ca}^{2+}$  flow through the channels in the transverse tubules may stimulate the opening of other calcium channels in the sarcoplasmic reticulum. This is termed a  *$\text{Ca}^{2+}$ -induced  $\text{Ca}^{2+}$  release mechanism*, and has been shown to be the major mechanism for excitation-contraction coupling in heart muscle. By these mechanisms,  $\text{Ca}^{2+}$  can be released from the sarcoplasmic reticulum, bind to troponin, and stimulate muscle contraction.

As long as action potentials continue to be produced—which is as long as the neural stimulation of the muscle is maintained— $\text{Ca}^{2+}$  will remain attached to troponin and cross bridges will be able to undergo contraction cycles. When neural activity and action potentials in the muscle fiber cease, the sarcoplasmic reticulum actively accumulates  $\text{Ca}^{2+}$  and muscle relaxation occurs. Note that the return of  $\text{Ca}^{2+}$  to the sarcoplasmic reticulum involves active transport and thus requires the hydrolysis of ATP. ATP is therefore needed for muscle relaxation as well as for muscle contraction.

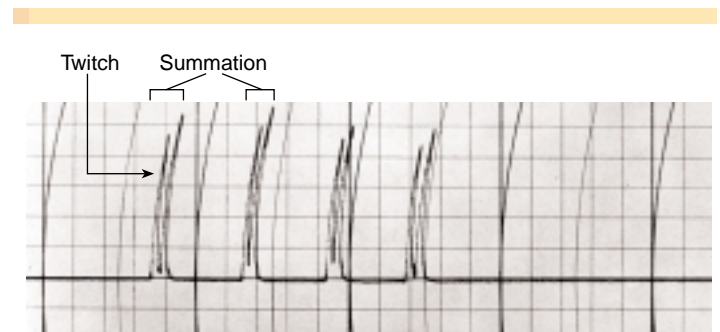
### Test Yourself Before You Continue

1. With reference to the sliding filament theory, explain how the lengths of the A, I, and H bands change during contraction.
2. Describe a cycle of cross-bridge activity during contraction and discuss the role of ATP in this cycle.
3. Draw a sarcomere in a relaxed muscle and a sarcomere in a contracted muscle and label the bands in each. What is the significance of the differences in your drawings?
4. Describe the molecular structure of myosin and actin. How are tropomyosin and troponin positioned in the thin filaments and how do they function in the contraction cycle?
5. Use a flowchart to show the sequence of events from the time ACh is released from a nerve ending to the time  $\text{Ca}^{2+}$  is released from the sarcoplasmic reticulum.
6. Explain the requirements for  $\text{Ca}^{2+}$  and ATP in muscle contraction and relaxation.

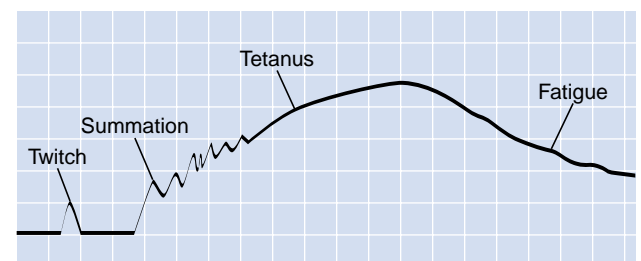
## Contractions of Skeletal Muscles

Contraction of muscles generates tension, which allows muscles to shorten and thereby perform work. The contraction strength of skeletal muscles must be sufficiently great to overcome the load on a muscle in order for that muscle to shorten.

The contractions of skeletal muscles generally produce movements of bones at joints, which act as levers to move the loads against which the muscle's force is exerted. The contractile behavior of the muscle, however, is more easily studied *in vitro* (outside the body) than *in vivo* (within the body). When a muscle—for example, the gastrocnemius (calf muscle) of a frog—is studied *in vitro*, it is usually mounted so that one end is fixed and the other is movable. The mechanical force of the muscle contraction is transduced (changed) into an electric current, which can be amplified and displayed as pen deflections in a multichannel recorder (fig. 12.16). In this way, the contractile behavior of the whole muscle in response to experimentally administered electric shocks can be studied.



(a)



(b)

**Figure 12.16** Recording muscle contractions. (a) Actual recorder tracings demonstrating twitch and summation of an isolated frog gastrocnemius muscle. (b) Illustration of a recording that demonstrates twitch, summation, tetanus, and fatigue. The process of fatigue was produced in this instance by maintaining the electrical stimulation of the muscle, and the mechanisms involved are described in a later section.



## Twitch, Summation, and Tetanus

When the muscle is stimulated with a single electric shock of sufficient voltage, it quickly contracts and relaxes. This response is called a **twitch**. Increasing the stimulus voltage increases the strength of the twitch, up to a maximum. The strength of a muscle contraction can thus be *graded*, or varied—an obvious requirement for the proper control of skeletal movements. If a second electric shock is delivered immediately after the first, it will produce a second twitch that may partially “ride piggy-back” on the first. This response is called **summation**.

Stimulation of fibers within a muscle *in vitro* with an electric stimulator, or *in vivo* by motor axons, usually results in the full contraction of the individual fibers. Stronger muscle contractions are produced by the stimulation of greater numbers of muscle fibers. Skeletal muscles can thus produce **graded contractions**, the strength of which depends on the number of fibers stimulated rather than on the strength of the contractions of individual muscle fibers.

If the stimulator is set to deliver an increasing frequency of electric shocks automatically, the relaxation time between successive twitches will get shorter and shorter as the strength of contraction increases in amplitude. This effect is known as **incomplete tetanus**. Finally, at a particular “fusion frequency” of stimulation, there is no visible relaxation between successive twitches (fig. 12.16). Contraction is smooth and sustained, as it is during normal muscle contraction *in vivo*. This smooth, sustained contraction is called **complete tetanus**. (The term *tetanus* should not be confused with the disease of the same name, which is accompanied by a painful state of muscle contracture, or *tetany*.) The tetanus produced *in vitro* by the asynchronous twitches of muscle fibers simulates the normal, smooth contraction produced *in vivo* by the asynchronous activation of motor units.

### Treppe

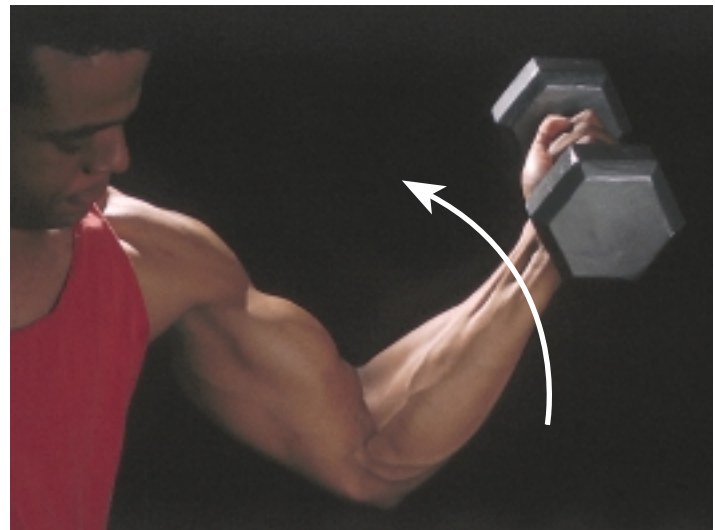
If the voltage of the electrical shocks delivered to an isolated muscle *in vitro* is gradually increased from zero, the strength of the muscle twitches will increase accordingly, up to a maximal value at which all of the muscle fibers are stimulated. This demonstrates the graded nature of the muscle contraction. If a series of electrical shocks at this maximal voltage is given to a fresh muscle so that each shock produces a separate twitch, each of the twitches evoked will be successively stronger, up to a higher maximum. This demonstrates **treppe**, or the *staircase effect*. Treppe may represent a warmup effect, and is believed to be due to an increase in intracellular  $\text{Ca}^{2+}$ , which is needed for muscle contraction.

## Isotonic and Isometric Contractions

In order for muscle fibers to shorten when they contract, they must generate a force that is greater than the opposing forces that act to prevent movement of the muscle’s insertion. When a weight is lifted by flexing the elbow joint, for example, the force produced by contraction of the biceps brachii muscle is greater



(a)



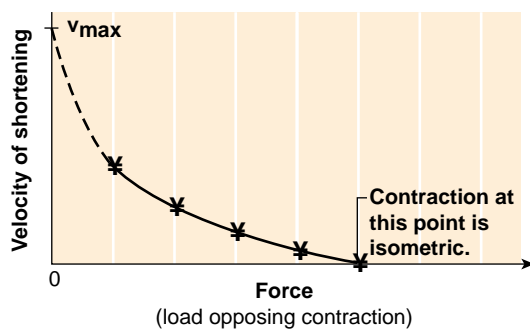
(b)

■ **Figure 12.17** Photographs of isometric and isotonic contractions. (a) An isometric contraction, where the muscle stays the same length, and (b) an isotonic contraction, where the muscle shortens.

than the force of gravity on the object being lifted (fig. 12.17). The tension produced by the contraction of each muscle fiber separately is insufficient to overcome the opposing force, but the combined contractions of numerous muscle fibers may be sufficient to overcome the opposing force and flex the forearm. In this case, the muscle and all of its fibers shorten in length.

This process can be seen by examining the **force-velocity curve**. This graph shows the inverse relationship between the force opposing muscle contraction (the load against which the muscle must work) and the velocity of muscle shortening (fig. 12.18). The tension produced by the shortening muscle is just greater than the force (load) at each value, causing the muscle to shorten. Since the contraction strength is constant at each load, a muscle contraction during shortening is called an **isotonic contraction** (*iso* = same; *tonic* = strength).

If the load is zero, a muscle contracts and shortens with its maximum velocity. As the load increases, the velocity of muscle



■ **Figure 12.18** Force-velocity curve. This graph illustrates the inverse relationship between the force opposing muscle contraction (the load against which the muscle must work) and the velocity of muscle shortening. A force that is sufficiently great prevents muscle shortening, so that the contraction is isometric. If there is no force acting against the muscle contraction, the velocity of shortening is maximal ( $V_{\max}$ ). Since this cannot be measured (because there will always be some load), the estimated position of the curve is shown with a dashed line.

shortening decreases. When the force opposing contraction (the load) becomes sufficiently great, the muscle is unable to shorten when it exerts a given tension. That is, its velocity of shortening is zero. At this point, where muscle tension does not cause muscle shortening, the contraction is called an **isometric** (literally, “same length”) **contraction**.

Isometric contraction can be voluntarily produced, for example, by lifting a weight and maintaining the forearm in a partially flexed position. We can then increase the amount of muscle tension produced by recruiting more muscle fibers until the muscle begins to shorten; at this point, isometric contraction is converted to isotonic contraction (see fig 12.17).

## Series-Elastic Component

In order for a muscle to shorten when it contracts, and thus to move its insertion toward its origin, the noncontractile parts of the muscle and the connective tissue of its tendons must first be pulled tight. These structures, particularly the tendons, have elasticity—they resist distension, and when the distending force is released, they tend to spring back to their resting lengths. Tendons provide what is called a **series-elastic component** because they are somewhat elastic and in line (in series) with the force of muscle contraction. The series-elastic component absorbs some of the tension as a muscle contracts, and it must be pulled tight before muscle contraction can result in muscle shortening.

When the gastrocnemius muscle was stimulated with a single electric shock as described earlier, the amplitude of the twitch was reduced because some of the force of contraction was used to stretch the series-elastic component. Quick delivery of a second shock thus produced a greater degree of muscle shortening than the first shock, culminating at the fusion fre-

quency of stimulation with complete tetanus, in which the strength of contraction was much greater than that of individual twitches.

Some of the energy used to stretch the series-elastic component during muscle contraction is released by elastic recoil when the muscle relaxes. This elastic recoil, which helps the muscles return to their resting length, is of particular importance for the muscles involved in breathing. As we will see in chapter 16, inspiration is produced by muscle contraction and expiration is produced by the elastic recoil of the thoracic structures that were stretched during inspiration.

## Length-Tension Relationship

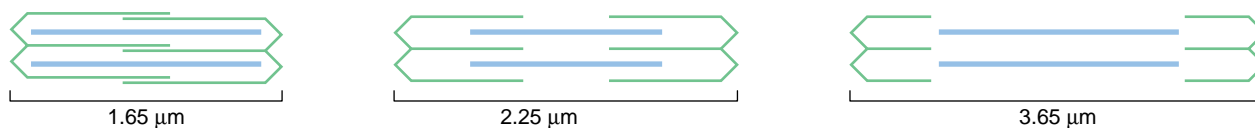
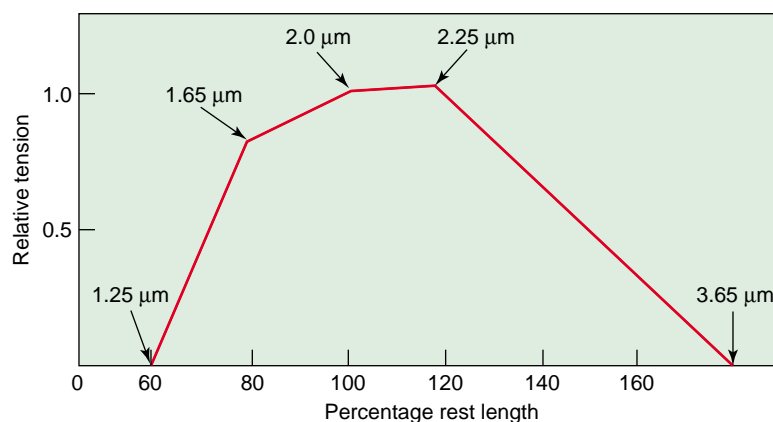
The strength of a muscle’s contraction is influenced by a variety of factors. These include the number of fibers within the muscle that are stimulated to contract, the frequency of stimulation, the thickness of each muscle fiber (thicker fibers have more myofibrils and thus can exert more power), and the initial length of the muscle fibers when they are at rest.

There is an “ideal” resting length for striated muscle fibers. This is the length at which they can generate maximum force. When the resting length exceeds this ideal, the overlap between actin and myosin is so small that few cross bridges can attach. When the muscle is stretched to the point that there is no overlap of actin with myosin, no cross bridges can attach to the thin filaments and the muscle cannot contract. When the muscle is shortened to about 60% of its resting length, the Z lines abut the thick filaments so that further contraction cannot occur.

The strength of a muscle’s contraction can be measured by the force required to prevent it from shortening. Under these isometric conditions, the strength of contraction, or *tension*, can be measured when the muscle length at rest is varied. Maximum tension of skeletal muscle is produced when the muscle is at its normal resting length in vivo (fig. 12.19). If the muscle were any shorter or longer than its normal length, in other words, its strength of contraction would be reduced. This resting length is maintained by reflex contraction in response to passive stretching, as described in a later section of this chapter.

### Test Yourself Before You Continue

1. Explain how graded contractions and smooth, sustained contractions can be produced in vitro and in vivo.
2. Distinguish between isotonic and isometric contractions, and describe what factors determine if a contraction will be isometric or isotonic.
3. Identify the nature and physiological significance of the series-elastic component of muscle contraction.
4. Describe the relationship between the resting muscle length and the strength of its contraction.



**Figure 12.19** The length-tension relationship in skeletal muscles. Maximum relative tension (1.0 on the y axis) is achieved when the muscle is 100% to 120% of its resting length (sarcomere lengths from 2.0 to 2.25  $\mu\text{m}$ ). Increases or decreases in muscle (and sarcomere) lengths result in rapid decreases in tension.

## Energy Requirements of Skeletal Muscles

Skeletal muscles generate ATP through aerobic and anaerobic respiration and through the use of phosphate groups donated by creatine phosphate. The aerobic and anaerobic abilities of skeletal muscle fibers differ according to muscle fiber type, which are described according to their speed of contraction, color, and major mode of energy metabolism.

Skeletal muscles at rest obtain most of their energy from the aerobic respiration of fatty acids. During exercise, muscle glycogen and blood glucose are also used as energy sources (fig. 12.20). Energy obtained by cell respiration is used to make ATP, which serves as the immediate source of energy for (1) the movement of the cross bridges for muscle contraction and (2) the pumping of  $\text{Ca}^{2+}$  into the sarcoplasmic reticulum for muscle relaxation.

## Metabolism of Skeletal Muscles

Skeletal muscles respire anaerobically for the first 45 to 90 seconds of moderate-to-heavy exercises because the cardiopulmonary system requires this amount of time to sufficiently increase the oxygen supply to the exercising muscles. If exercise

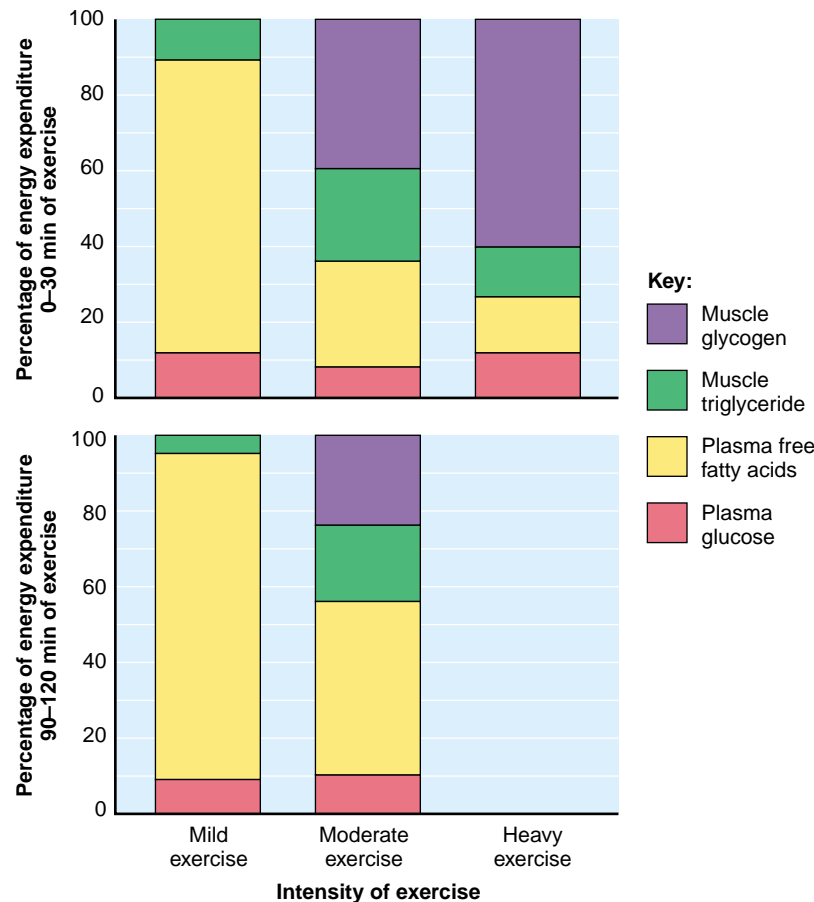
is moderate, aerobic respiration contributes the major portion of the skeletal muscle energy requirements following the first 2 minutes of exercise.

### Maximal Oxygen Uptake

Whether exercise is light, moderate, or heavy for a given person depends on that person's maximal capacity for aerobic exercise. The maximum rate of oxygen consumption (by aerobic respiration) in the body is called the **maximal oxygen uptake**, or the **aerobic capacity**, and is often expressed in abbreviated form as the  $\dot{V}_{\text{O}_2\text{max}}$ . The maximal oxygen uptake is determined primarily by a person's age, size, and sex. It is from 15% to 20% higher for males than for females and highest at age 20 for both sexes. The  $\dot{V}_{\text{O}_2\text{max}}$  ranges from about 12 ml of  $\text{O}_2$  per minute per kilogram body weight for older, sedentary people to about 84 ml per minute per kilogram for young, elite male athletes. Some world-class athletes have maximal oxygen uptakes that are twice the average for their age and sex—this appears to be due largely to genetic factors, but training can increase the maximum oxygen uptake by about 20%.

The intensity of exercise can also be defined by the **lactate (or anaerobic) threshold**. This is the percentage of the maximal oxygen uptake at which a significant rise in blood lactate levels occurs. For average healthy people, for example, a significant amount of blood lactate appears when exercise is performed at about 50% to 70% of the  $\dot{V}_{\text{O}_2\text{max}}$ .

During light exercise (at about 25% of the  $\dot{V}_{\text{O}_2\text{max}}$ ), most of the exercising muscle's energy is derived from the aerobic respiration of fatty acids. These are derived mainly from stored



■ **Figure 12.20** Muscle fuel consumption during exercise. The relative contributions of plasma glucose, plasma free fatty acids, muscle glycogen, and muscle triglycerides to the energy consumption of exercising muscles. These are shown during mild exercise (25% of  $\dot{V}_{O_2max}$ ), moderate exercise (65% of  $\dot{V}_{O_2max}$ ), and heavy exercise (85% of  $\dot{V}_{O_2max}$ ).

fat in adipose tissue, and to a lesser extent from triglycerides stored in the muscle (fig. 12.20). When a person exercises just below the lactate threshold, where the exercise can be described as moderately intense (at 50% to 70% of the  $\dot{V}_{O_2max}$ ), the energy is derived almost equally from fatty acids and glucose (obtained from stored muscle glycogen and the blood plasma). By contrast, glucose from these sources supplies two-thirds of the energy for muscles during heavy exercise above the lactate threshold.

During exercise, the carrier protein for the facilitated diffusion of glucose (GLUT4—chapter 6) is moved into the muscle cell membrane, so that the cell can take up an increasing amount of blood glucose. The uptake of plasma glucose contributes 15% to 30% of the muscle's energy needs during moderate exercise and up to 40% of the energy needs during very heavy exercise. This would produce hypoglycemia if the liver failed to increase its output of glucose. The liver increases its output of glucose primarily through hydrolysis of its stored glycogen, but gluconeogenesis (the production of glucose from amino acids, lactate, and glycerol) contributes increasingly to the liver's glucose production as exercise is prolonged.

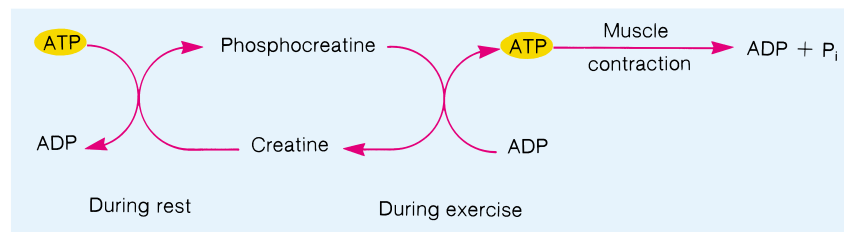
### Clinical Investigation Clue

Remember that Maria has a high maximal oxygen uptake, consistent with her athletic lifestyle.

Is it possible, likely, or unlikely that Maria's muscle pain and fatigue are caused by her playing softball?

### Oxygen Debt

When a person stops exercising, the rate of oxygen uptake does not immediately return to pre-exercise levels; it returns slowly (the person continues to breathe heavily for some time afterward). This extra oxygen is used to repay the **oxygen debt** incurred during exercise. The oxygen debt includes oxygen that was withdrawn from savings deposits—hemoglobin in blood and myoglobin in muscle (see chapter 16); the extra oxygen required for metabolism by tissues warmed during exercise; and the oxygen needed for the metabolism of the lactic acid produced during anaerobic respiration.

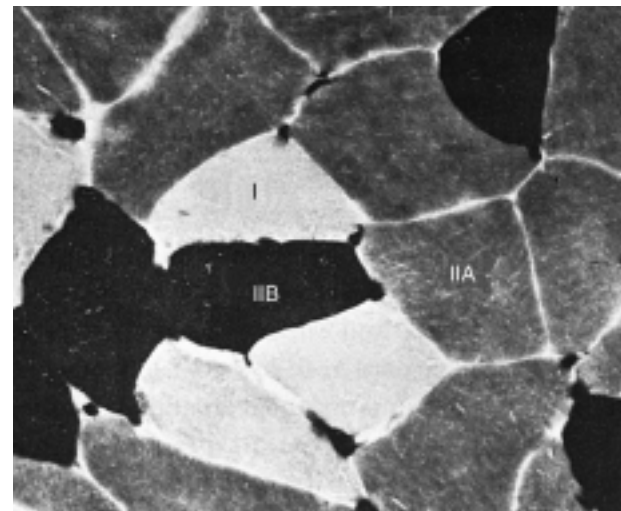


■ **Figure 12.21** The production and utilization of phosphocreatine in muscles. Phosphocreatine serves as a muscle reserve of high-energy phosphate, used for the rapid formation of ATP.

### Phosphocreatine

During sustained muscle activity, ATP may be used faster than it can be produced through cell respiration. At these times, the rapid renewal of ATP is extremely important. This is accomplished by combining ADP with phosphate derived from another high-energy phosphate compound called **phosphocreatine**, or **creatine phosphate**.

Within muscle cells, the phosphocreatine concentration is more than three times the concentration of ATP and represents a ready reserve of high-energy phosphate that can be donated directly to ADP (fig. 12.21). Production of ATP from ADP and phosphocreatine is so efficient that, even though the rate of ATP breakdown rapidly increases from rest to heavy exercise, muscle ATP concentrations hardly change! During times of rest, the depleted reserve of phosphocreatine can be restored by the reverse reaction—phosphorylation of creatine with phosphate derived from ATP.



■ **Figure 12.22** A skeletal muscle (of a cat) stained to indicate the activity of myosin ATPase. ATPase activity is greater in the type II (fast twitch) fibers than in the type I (slow twitch). Among the fast-twitch fibers, ATPase activity produces a darker stain in type II B (fast-glycolytic) fibers than in type II A (fast-oxidative) fibers.

## Slow- and Fast-Twitch Fibers

Skeletal muscle fibers can be divided on the basis of their contraction speed (time required to reach maximum tension) into **slow-twitch**, or **type I, fibers**, and **fast-twitch**, or **type II, fibers**. These differences are associated with different myosin ATPase isoenzymes, which can also be designated as “slow” and “fast.” The two fiber types can be distinguished by their ATPase isoenzyme when they are appropriately stained (fig. 12.22). The extraocular muscles that position the eyes, for example, have a high proportion of fast-twitch fibers and reach maximum tension in about 7.3 msec (milliseconds—thousandths of a second). The soleus muscle in the leg, by contrast, has a high proportion of slow-twitch fibers and requires about 100 msec to reach maximum tension (fig. 12.23).

Muscles like the soleus are *postural muscles*; they are able to sustain a contraction for a long period of time without

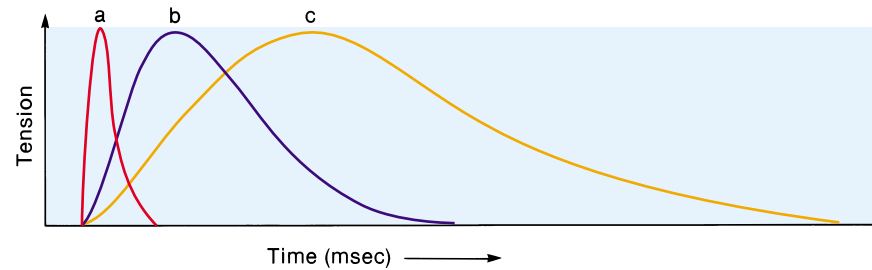


The enzyme that transfers phosphate between creatine and ATP is called **creatine kinase**, or **creatine phosphokinase**. Skeletal muscle and heart muscle have two different forms of this enzyme (they have different isoenzymes, as described in chapter 4). The skeletal muscle isoenzyme is found to be elevated in the blood of people with muscular dystrophy (degenerative disease of skeletal muscles). The plasma concentration of the isoenzyme characteristic of heart muscle is elevated as a result of myocardial infarction (damage to heart muscle), and measurements of this enzyme are thus used as a means of diagnosing heart disease.

### Clinical Investigation Clue

Remember that Maria had a normal blood level of creatine phosphokinase.

■ *What does this suggest about the health of her muscles and heart?*



■ **Figure 12.23** A comparison of the rates at which maximum tension is developed in three muscles. These are (a) the relatively fast-twitch extraocular and (b) gastrocnemius muscles, and (c) the slow-twitch soleus muscle.

**Table 12.4** Characteristics of Muscle Fiber Types

Feature	Slow Oxidative/Red (Type I)	Fast Oxidative/White (Type II A)	Fast Glycolytic/White (Type II B)
Diameter	Small	Intermediate	Large
Z-line thickness	Wide	Intermediate	Narrow
Glycogen content	Low	Intermediate	High
Resistance to fatigue	High	Intermediate	Low
Capillaries	Many	Many	Few
Myoglobin content	High	High	Low
Respiration	Aerobic	Aerobic	Anaerobic
Oxidative capacity	High	High	Low
Glycolytic ability	Low	High	High
Twitch rate	Slow	Fast	Fast
Myosin ATPase content	Low	High	High

fatigue. The resistance to fatigue demonstrated by these muscles is aided by other characteristics of slow-twitch (type I) fibers that endow them with a high oxidative capacity for aerobic respiration. Hence, the type I fibers are often referred to as **slow oxidative fibers**. These fibers have a rich capillary supply, numerous mitochondria and aerobic respiratory enzymes, and a high concentration of *myoglobin*. Myoglobin is a red pigment, similar to the hemoglobin in red blood cells, that improves the delivery of oxygen to the slow-twitch fibers. Because of their high myoglobin content, slow-twitch fibers are also called **red fibers**.

The thicker, fast-twitch (type II) fibers have fewer capillaries and mitochondria than slow-twitch fibers and not as much myoglobin; hence, these fibers are also called **white fibers**. Fast-twitch fibers are adapted to respire anaerobically by a large store of glycogen and a high concentration of glycolytic enzymes. In addition to the type I (slow-twitch) and type II (fast-twitch) fibers, human muscles have an intermediate fiber type. These intermediate fibers are fast-twitch but also have a high oxidative capacity; therefore, they are relatively resistant to fatigue. They are called **type IIA fibers**, or **fast oxidative fibers**, to distinguish them from the anaerobically adapted fast-twitch **type IIB fibers**, or **fast glycolytic fibers**. The low oxidative capacity causes these fibers to fatigue rapidly. The three fiber types are compared in table 12.4.

Interestingly, the conduction rate of motor neurons that innervate fast-twitch fibers is faster (80–90 meters per second)

than the conduction rate to slow-twitch fibers (60–70 meters per second). The fiber type indeed seems to be determined by the motor neuron. When the motor neurons to different fiber types are switched in experimental animals, the previously fast-twitch fibers become slow and the slow-twitch fibers become fast. As expected from these observations, all of the muscle fibers innervated by the same motor neuron (that are part of the same motor unit) are of the same type.

A muscle such as the gastrocnemius contains both fast- and slow-twitch fibers, although fast-twitch fibers predominate. A given somatic motor axon, however, innervates muscle fibers of one type only. The sizes of these motor units differ; the motor units composed of slow-twitch fibers tend to be smaller (have fewer fibers) than the motor units of fast-twitch fibers. As mentioned earlier, motor units are recruited from smaller to larger when increasing effort is required; thus, the smaller motor units with slow-twitch fibers would be used most often in routine activities. Larger motor units with fast-twitch fibers, which can exert a great deal of force but which respire anaerobically and thus fatigue quickly, would be used relatively infrequently and for only short periods of time.

## Muscle Fatigue

**Muscle fatigue** may be defined as the inability to maintain a particular muscle tension when the contraction is sustained or to reproduce a particular tension during rhythmic contraction over

time. Fatigue during a sustained maximal contraction, when all the motor units are used and the rate of neural firing is maximal—as when lifting an extremely heavy weight—appears to be due to an accumulation of extracellular  $K^+$  (Remember that  $K^+$  leaves axons and muscle fibers during the repolarization phase of action potentials.) This reduces the membrane potential of muscle fibers and interferes with their ability to produce action potentials. Fatigue under these circumstances lasts only a short time, and maximal tension can again be produced after less than a minute's rest.

Fatigue during moderate exercise occurs as the slow-twitch fibers deplete their reserve glycogen and fast-twitch fibers are increasingly recruited. Fast-twitch fibers obtain their energy through anaerobic respiration, converting glucose to lactic acid, and this results in a rise in intracellular  $H^+$  and a fall in pH. The decrease in muscle pH, in turn, promotes muscle fatigue, but the exact physiological mechanisms by which this occurs are not well understood. One possibility is that there may be a reduced ability of the sarcoplasmic reticulum to accumulate  $Ca^{2+}$  by active transport, or there may be a reduced ability of the sarcoplasmic reticulum to release  $Ca^{2+}$  in response to stimulation. By either mechanism, the decrease in cellular pH would produce muscle fatigue by interfering with excitation-contraction coupling.

## Adaptations of Muscles to Exercise Training

The maximal oxygen uptake, obtained during very strenuous exercise, averages 50 ml of  $O_2$  per minute per kilogram body weight in males between the ages of 20 and 25 (females average 25% lower). For trained endurance athletes (such as swimmers and long-distance runners), maximal oxygen uptakes can be as high as 86 ml of  $O_2$  per minute per kilogram. These considerable differences affect the lactate threshold, and thus the amount of exercise that can be performed before lactic acid production contributes to muscle fatigue. In addition to having a higher aerobic capacity, well-trained athletes also have a lactate threshold that is a higher percentage of their  $\dot{V}_{O_2\max}$ . The lactate threshold of an untrained person, for example, might be 60% of the  $\dot{V}_{O_2\max}$ , whereas the lactate threshold of a trained athlete can be up to 80% of the  $\dot{V}_{O_2\max}$ . These athletes thus produce less lactic acid at a given level of exercise than the average person, and therefore they are less subject to fatigue than the average person.

Since the depletion of muscle glycogen places a limit on exercise, any adaptation that spares muscle glycogen will improve physical endurance. This is achieved in trained athletes by an increased proportion of energy that is derived from the aerobic respiration of fatty acids, resulting in a slower depletion of their muscle glycogen. The greater the level of physical training, the higher the proportion of energy derived from the oxidation of fatty acids during exercise below the  $\dot{V}_{O_2\max}$ .

All fiber types adapt to endurance training by an increase in mitochondria, and thus in aerobic respiratory enzymes. In fact, the maximal oxygen uptake can be increased by as much as

20% through endurance training. There is a decrease in type IIB (fast glycolytic) fibers, which have a low oxidative capacity, accompanied by an increase in type IIA (fast oxidative) fibers, which have a high oxidative capacity. Although the type IIA fibers are still classified as fast-twitch, they show an increase in the slow myosin ATPase isoenzyme form, indicating that they are in a transitional state between the type II and type I fibers. A summary of the changes that occur as a result of endurance training is presented in table 12.5.

Endurance training does not increase the size of muscles. Muscle enlargement is produced only by frequent periods of high-intensity exercise in which muscles work against a high resistance, as in weightlifting. As a result of resistance training, type II muscle fibers become thicker, and the muscle therefore grows by hypertrophy (an increase in cell size rather than number of cells). This happens first because the myofibrils within a muscle fiber thicken because of the synthesis of actin and myosin proteins and the addition of new sarcomeres. Then, after a myofibril has attained a certain thickness, it may split into two myofibrils, each of which may become thicker as a result of the addition of sarcomeres. Muscle hypertrophy, in short, is associated with an increase in the size of the myofibrils, and then in the number of myofibrils within the muscle fibers.

The decline in physical strength of older people is associated with a reduced muscle mass, which is due to a decrease in the size of fast-twitch muscle fibers. Aging is also associated with a reduced density of blood capillaries surrounding the muscle fibers, leading to a decrease in oxidative capacity. These changes are partially caused by a more sedentary lifestyle and can be largely reversed through physical training. Resistance training has been shown to increase muscle mass in older people, and endurance training to increase the density of blood capillaries in the muscles. The muscle glycogen of older people can also be increased by endurance training, but it cannot be raised to the levels present in youth.

**Table 12.5** Effects of Endurance Training on Skeletal Muscles

1. Improved ability to obtain ATP from oxidative phosphorylation
2. Increased size and number of mitochondria
3. Less lactic acid produced per given amount of exercise
4. Increased myoglobin content
5. Increased intramuscular triglyceride content
6. Increased lipoprotein lipase (enzyme needed to utilize lipids from blood)
7. Increased proportion of energy derived from fat; less from carbohydrates
8. Lower rate of glycogen depletion during exercise
9. Improved efficiency in extracting oxygen from blood
10. Decreased number of type IIB (fast glycolytic) fibers; increased number of type IIA (fast oxidative) fibers

## Test Yourself Before You Continue

1. Draw a figure illustrating the relationship between ATP and creatine phosphate, and explain the physiological significance of this relationship.
2. Describe the characteristics of slow- and fast-twitch fibers (including intermediate fibers). Explain how the fiber types are determined and list the functions of different fiber types.
3. Explain the different causes of muscle fatigue with reference to the various fiber types.
4. Describe the effects of endurance training and resistance training on the fiber characteristics of muscles.



The disease known as **amyotrophic lateral sclerosis (ALS)** involves degeneration of the lower motor neurons, leading to muscle paralysis. This disease is sometimes called Lou Gehrig's disease, after the baseball player who suffered from it, and also includes the famous physicist Steven Hawking among its victims. Scientists have recently learned that the inherited form of this disease is caused by a defect in the gene for a specific enzyme—*superoxide dismutase*. This enzyme is responsible for eliminating superoxide free radicals, which are highly toxic products that can damage the motor neurons. The mutant gene produces an enzyme that has a different, and in fact destructive, action.

## Neural Control of Skeletal Muscles

Skeletal muscles contain stretch receptors called muscle spindles that stimulate the production of impulses in sensory neurons when a muscle is stretched. These sensory neurons can synapse with alpha motoneurons, which stimulate the muscle to contract in response to the stretch. Other motor neurons, called gamma motoneurons, stimulate the tightening of the spindles and thus increase their sensitivity.

Motor neurons in the spinal cord, or **lower motor neurons** (often shortened to *motoneurons*), are those previously described that have cell bodies in the spinal cord and axons within nerves that stimulate muscle contraction (table 12.6). The activity of these neurons is influenced by (1) sensory feedback from the muscles and tendons and (2) facilitatory and inhibitory effects from **upper motor neurons** in the brain that contribute axons to descending motor tracts. Lower motor neurons are thus said to be the *final common pathway* by which sensory stimuli and higher brain centers exert control over skeletal movements.

The cell bodies of lower motor neurons are located in the ventral horn of the gray matter of the spinal cord (chapter 8). Axons from these cell bodies leave the ventral side of the spinal cord to form the *ventral roots* of spinal nerves (see fig. 8.23). The *dorsal roots* of spinal nerves contain sensory fibers whose cell bodies are located in the *dorsal root ganglia*. Both sensory (*afferent*) and motor (*efferent*) fibers join in a common connective tissue sheath to form the spinal nerves at each segment of the spinal cord. In the lumbar region there are about 12,000 sensory and 6,000 motor fibers per spinal nerve.

About 375,000 cell bodies have been counted in a lumbar segment—a number far larger than can be accounted for by the number of motor neurons. Most of these neurons do not contribute fibers to the spinal nerve. Rather, they serve as *interneurons*, whose fibers conduct impulses up, down, and across the central nervous system. Those fibers that conduct impulses to higher spinal cord segments and the brain form *ascending tracts*, and those that conduct to lower spinal segments contribute to *descending tracts*. Those fibers that cross the midline of the CNS to synapse on the opposite side are part of *commissural tracts*. Interneurons can thus conduct impulses up and down on the same, or *ipsilateral*, side, and can affect neurons on the opposite, or *contralateral*, side of the central nervous system.

**Table 12.6** A Partial Listing of Terms Used to Describe the Neural Control of Skeletal Muscles

Term	Description
1. Lower motoneurons	Neurons whose axons innervate skeletal muscles—also called the “final common pathway” in the control of skeletal muscles
2. Higher motoneurons	Neurons in the brain that are involved in the control of skeletal movements and that act by facilitating or inhibiting (usually by way of interneurons) the activity of the lower motoneurons
3. Alpha motoneurons	Lower motoneurons whose fibers innervate ordinary (extrafusal) muscle fibers
4. Gamma motoneurons	Lower motoneurons whose fibers innervate the muscle spindle fibers (intrafusal fibers)
5. Agonist/antagonist	A pair of muscles or muscle groups that insert on the same bone, the agonist being the muscle of reference
6. Synergist	A muscle whose action facilitates the action of the agonist
7. Ipsilateral/contralateral	Ipsilateral—located on the same side, or the side of reference; contralateral—located on the opposite side
8. Afferent/efferent	Afferent neurons—sensory; efferent neurons—motor



**Table 12.7 Spindle Apparatus Content of Selected Skeletal Muscles**

Muscle	Muscle Weight (g)	Average Number of Spindles	Number of Spindles per Gram Muscle
Gastrocnemius	7.6	35	5
Rectus femoris	8.36	104	12
Tibialis anterior	4.57	71	15
Semitendinosus	6.41	114	18
Soleus	2.49	56	23
Fifth interossei—foot	0.33	29	88
Fifth interossei—hand	0.21	25	119

## Muscle Spindle Apparatus

In order for the nervous system to control skeletal movements properly, it must receive continuous sensory feedback information concerning the effects of its actions. This sensory information includes (1) the tension that the muscle exerts on its tendons, provided by the **Golgi tendon organs**, and (2) muscle length, provided by the **muscle spindle apparatus**. The spindle apparatus, so called because it is wider in the center and tapers toward the ends, functions as a length detector. Muscles that require the finest degree of control, such as the muscles of the hand, have the highest density of spindles (table 12.7).

Each spindle apparatus contains several thin muscle cells, called *intrafusal fibers* (*fusus* = spindle), packaged within a connective tissue sheath. Like the stronger and more numerous “ordinary” muscle fibers outside the spindles—the *extrafusal fibers*—the spindles insert into tendons on each end of the muscle. Spindles are therefore said to be in parallel with the extrafusal fibers.

Unlike the extrafusal fibers, which contain myofibrils along their entire length, the contractile apparatus is absent from the central regions of the intrafusal fibers. The central, noncontracting part of an intrafusal fiber contains nuclei. There are two types of intrafusal fibers. One type, the *nuclear bag fibers*, have their nuclei arranged in a loose aggregate in the central regions of the fibers. The other type of intrafusal fibers, called *nuclear chain fibers*, have their nuclei arranged in rows. Two types of sensory neurons serve these intrafusal fibers. **Primary**, or **annulospiral, sensory endings** wrap around the central regions of the nuclear bag and chain fibers (fig. 12.24), and **secondary**, or **flower-spray, endings** are located over the contracting poles of the nuclear chain fibers.

Since the spindles are arranged in parallel with the extrafusal muscle fibers, stretching a muscle causes its spindles to stretch. This stimulates both the primary and secondary sensory endings. The spindle apparatus thus serves as a length detector because the frequency of impulses produced in the primary and secondary endings is proportional to the length of the muscle. The primary endings, however, are most stimulated at the onset of stretch, whereas the secondary endings respond in a more tonic (sustained) fashion as stretch is maintained. Sudden, rapid stretching of a muscle activates both types of sensory endings, and is thus a more powerful stimulus for the muscle spindles

than a slower, more gradual stretching that has less of an effect on the primary sensory endings. Since the activation of the sensory endings in muscle spindles produces a reflex contraction, the force of this reflex contraction is greater in response to rapid stretch than to gradual stretch.

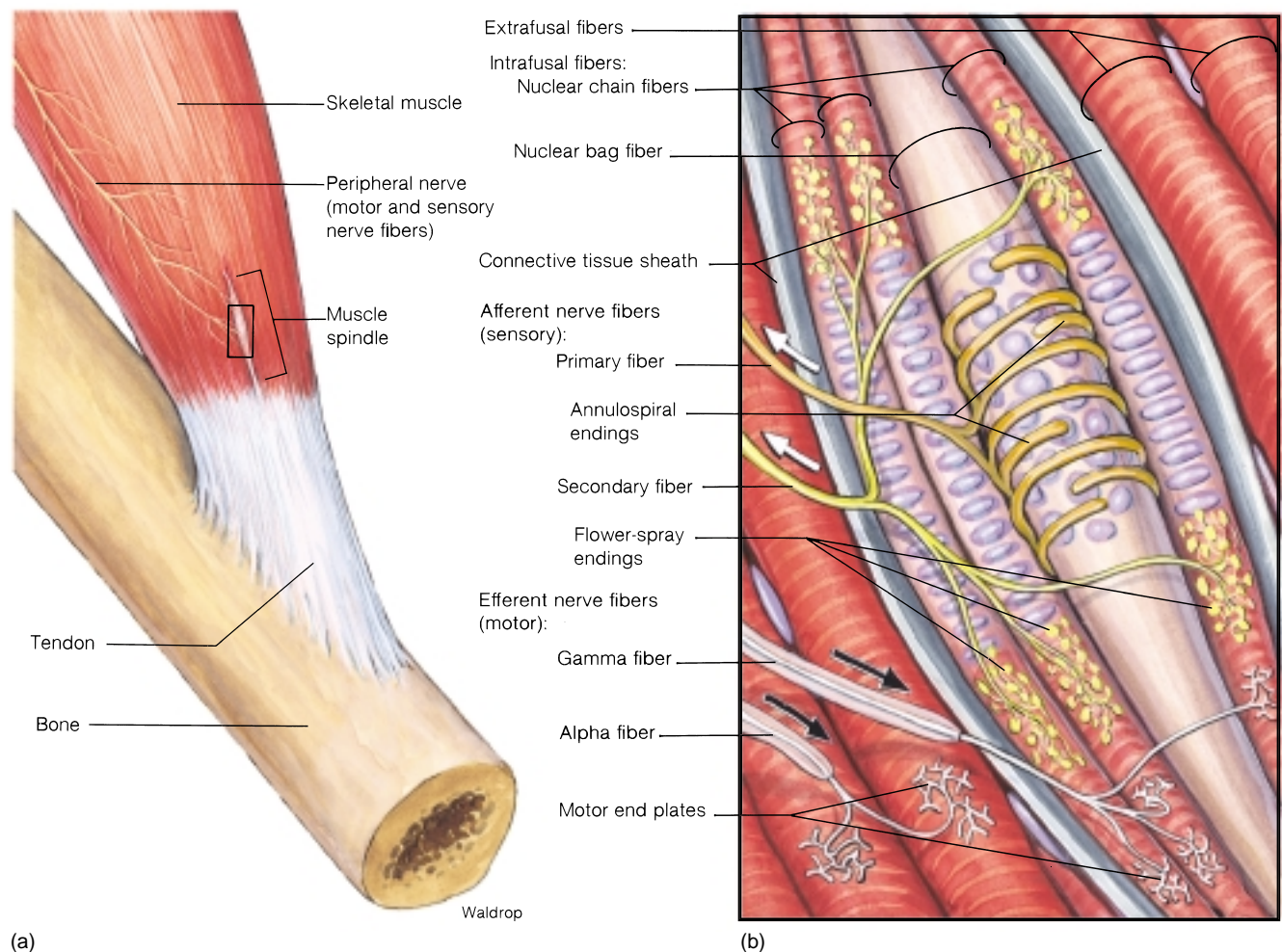


Rapid stretching of skeletal muscles produces very forceful muscle contractions as a result of the activation of primary and secondary endings in the muscle spindles and the monosynaptic stretch reflex. This can result in painful muscle spasms, as may occur, for example, when muscles are forcefully pulled in the process of setting broken bones. Painful muscle spasms may be avoided in physical exercise by stretching slowly and thereby stimulating mainly the secondary endings in the muscle spindles. A slower rate of stretch also allows time for the inhibitory Golgi tendon organ reflex to occur and promote muscle relaxation.

## Alpha and Gamma Motoneurons

In the spinal cord, two types of lower motor neurons innervate skeletal muscles. The motor neurons that innervate the extrafusal muscle fibers are called **alpha motoneurons**; those that innervate the intrafusal fibers are called **gamma motoneurons** (fig. 12.24). The alpha motoneurons are faster conducting (60–90 meters per second) than the thinner gamma motoneurons (10–40 meters per second). Since only the extrafusal muscle fibers are sufficiently strong and numerous to cause a muscle to shorten, only stimulation by the alpha motoneurons can cause muscle contraction that results in skeletal movements.

The intrafusal fibers of the muscle spindle are stimulated to contract by gamma motoneurons, which represent one-third of all efferent fibers in spinal nerves. However, because the intrafusal fibers are too few in number and their contraction too weak to cause a muscle to shorten, stimulation by gamma motoneurons results only in isometric contraction of the spindles. Since myofibrils are present in the poles but absent in the central regions of intrafusal fibers, the more distensible central region of the intrafusal fiber is pulled toward the ends in response to



■ **Figure 12.24** The location and structure of a muscle spindle. (a) A muscle spindle within a skeletal muscle. (b) The structure and innervation of a muscle spindle.

stimulation by gamma motoneurons. As a result, the spindle is tightened. This effect of gamma motoneurons, which is sometimes termed *active stretch* of the spindles, serves to increase the sensitivity of the spindles when the entire muscle is passively stretched by external forces. The activation of gamma motoneurons thus enhances the stretch reflex and is an important factor in the voluntary control of skeletal movements.

## Coactivation of Alpha and Gamma Motoneurons

Most of the fibers in the descending motor tracts synapse with interneurons in the spinal cord; only about 10% of the descending fibers synapse directly with the lower motor neurons. It is likely that very rapid movements are produced by direct synapses with the lower motor neurons, whereas most other movements are produced indirectly via synapses with spinal interneurons, which in turn stimulate the motor neurons.

*Upper motor neurons*—neurons in the brain that contribute fibers to descending motor tracts—usually stimulate alpha and gamma motoneurons simultaneously. Such stimulation is known as **coactivation**. Stimulation of alpha motoneurons results in muscle contraction and shortening; stimulation of gamma motoneurons stimulates contraction of the intrafusal fibers, and thus “takes out the slack” that would otherwise be present in the spindles as the muscles shorten. In this way, the spindles remain under tension and provide information about the length of the muscle even while the muscle is shortening.

Under normal conditions, the activity of gamma motoneurons is maintained at the level needed to keep the muscle spindles under proper tension while the muscles are relaxed. Undue relaxation of the muscles is prevented by stretch and activation of the spindles, which in turn elicits a reflex contraction (described in the next section). This mechanism produces a normal resting muscle length and state of tension, or **muscle tone**.

## Skeletal Muscle Reflexes

Although skeletal muscles are often called voluntary muscles because they are controlled by descending motor pathways that are under conscious control, they often contract in an unconscious, reflex fashion in response to particular stimuli. In the simplest type of reflex, a skeletal muscle contracts in response to the stimulus of muscle stretch. More complex reflexes involve inhibition of antagonistic muscles and regulation of a number of muscles on both sides of the body.

### The Monosynaptic Stretch Reflex

Reflex contraction of skeletal muscles occurs in response to sensory input and does not depend on the activation of upper motor neurons. The **reflex arc**, which describes the nerve impulse pathway from sensory to motor endings in such reflexes, involves only a few synapses within the CNS. The simplest of all reflexes—the *muscle stretch reflex*—consists of only one synapse within the CNS. The sensory neuron directly synapses with the motor neuron, without involving spinal cord interneurons. The stretch reflex is thus a **monosynaptic reflex** in terms of the individual reflex arcs (although, of course, many sensory neurons are activated at the same time, leading to the activation of many motor neurons). Resting skeletal muscles are maintained at an optimal length, as previously described under the heading “Length-Tension Relationship,” by stretch reflexes.

The stretch reflex is present in all muscles, but it is most dramatic in the extensor muscles of the limbs. The **knee-jerk reflex**—the most commonly evoked stretch reflex—is initiated by striking the patellar ligament with a rubber mallet. This stretches the entire body of the muscle, and thus passively stretches the spindles within the muscle so that sensory nerves with primary (annulospiral) endings in the spindles are activated. Axons of these sensory neurons synapse within the ventral gray matter of the spinal cord with *alpha motoneurons*. These large, fast-conducting motor nerve fibers stimulate the extrafusal fibers of the extensor muscle, resulting in isotonic contraction and the knee jerk. This is an example of negative feedback—stretching of the muscles (and spindles) stimulates shortening of the muscles (and spindles). These events are summarized in table 12.8 and illustrated in figure 12.25.



Damage to spinal nerves, or to the cell bodies of lower motor neurons (by poliovirus, for example), produces a **flaccid paralysis**, characterized by reduced muscle tone, depressed stretch reflexes, and atrophy. Damage to upper motor neurons or descending motor tracts at first produces spinal shock in which there is a flaccid paralysis. This is followed in a few weeks by **spastic paralysis**, characterized by increased muscle tone, exaggerated stretch reflexes, and other signs of hyperactive lower motor neurons.

The appearance of spastic paralysis suggests that upper motor neurons normally exert an inhibitory effect on lower alpha and gamma motor neurons. When this inhibition is removed, the gamma motoneurons become hyperactive and the spindles thus become overly sensitive to stretch. This can be demonstrated dramatically by forcefully dorsiflecting the patient's foot (pushing it up) and then releasing it. Forced extension stretches the antagonistic flexor muscles, which contract and produce the opposite movement (plantar flexion). Alternative activation of antagonistic stretch reflexes produces a flapping motion known as *clonus*.

### Golgi Tendon Organs

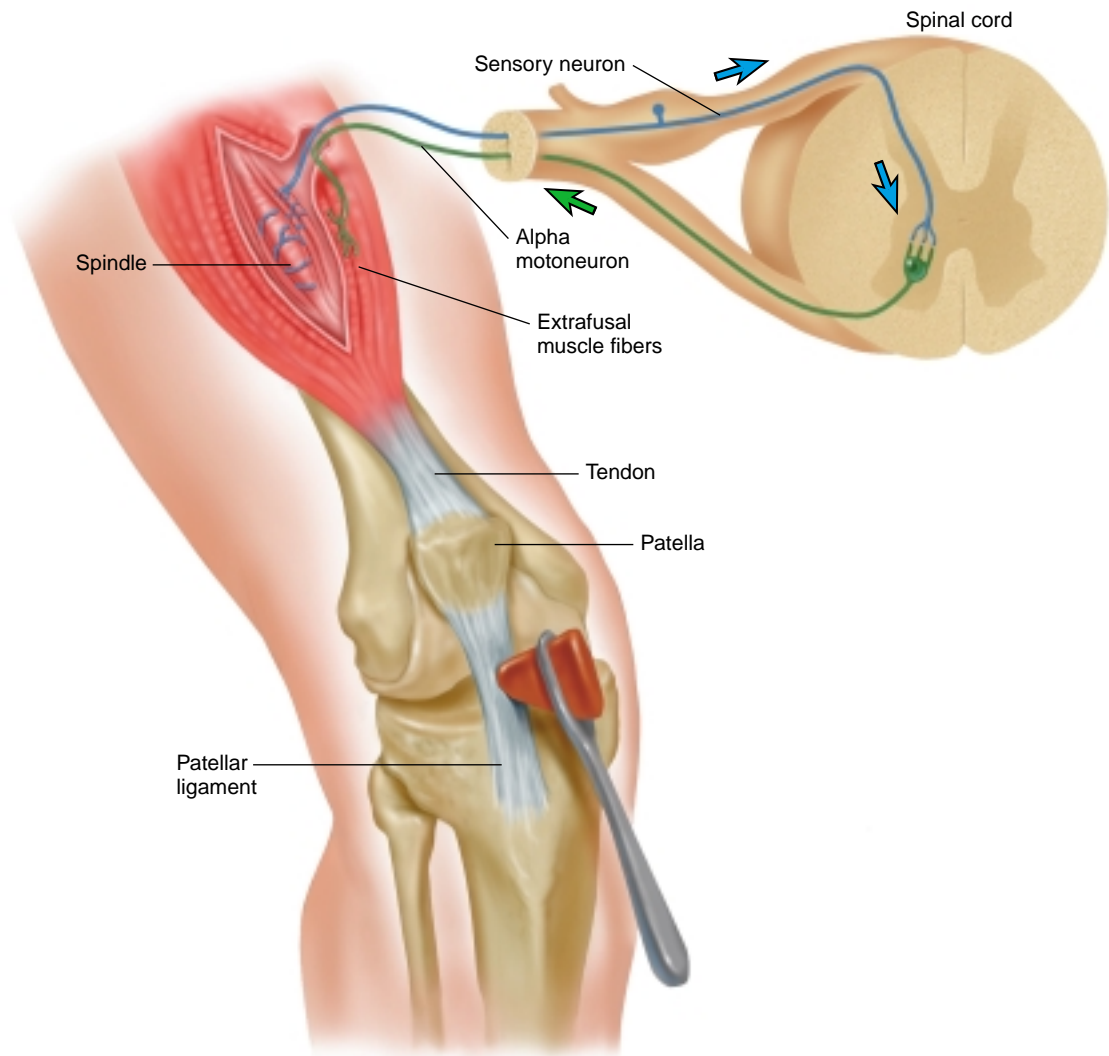
The **Golgi tendon organs** continuously monitor tension in the tendons produced by muscle contraction or passive stretching of a muscle. Sensory neurons from these receptors synapse with interneurons in the spinal cord; these interneurons, in turn, have *inhibitory synapses* (via IPSPs and postsynaptic inhibition—chapter 7) with motor neurons that innervate the muscle (fig. 12.26). This inhibitory Golgi tendon organ reflex is called a **disynaptic reflex** (because two synapses are crossed in the CNS), and it helps to prevent excessive muscle contractions or excessive passive muscle stretching. Indeed, if a muscle is stretched extensively, it will actually relax as a result of the inhibitory effects produced by the Golgi tendon organs.

### Reciprocal Innervation and the Crossed-Extensor Reflex

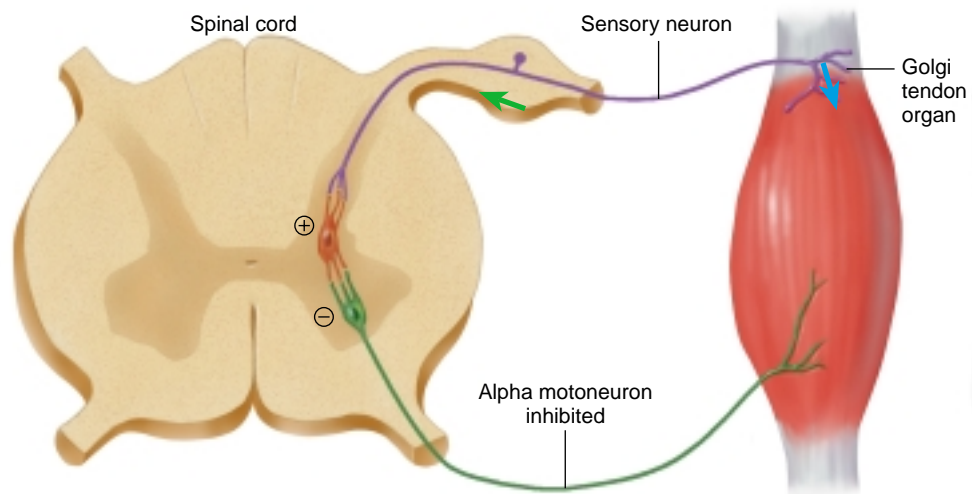
In the knee-jerk and other stretch reflexes, the sensory neuron that stimulates the motor neuron of a muscle also stimulates

**Table 12.8 Summary of Events in a Monosynaptic Stretch Reflex**

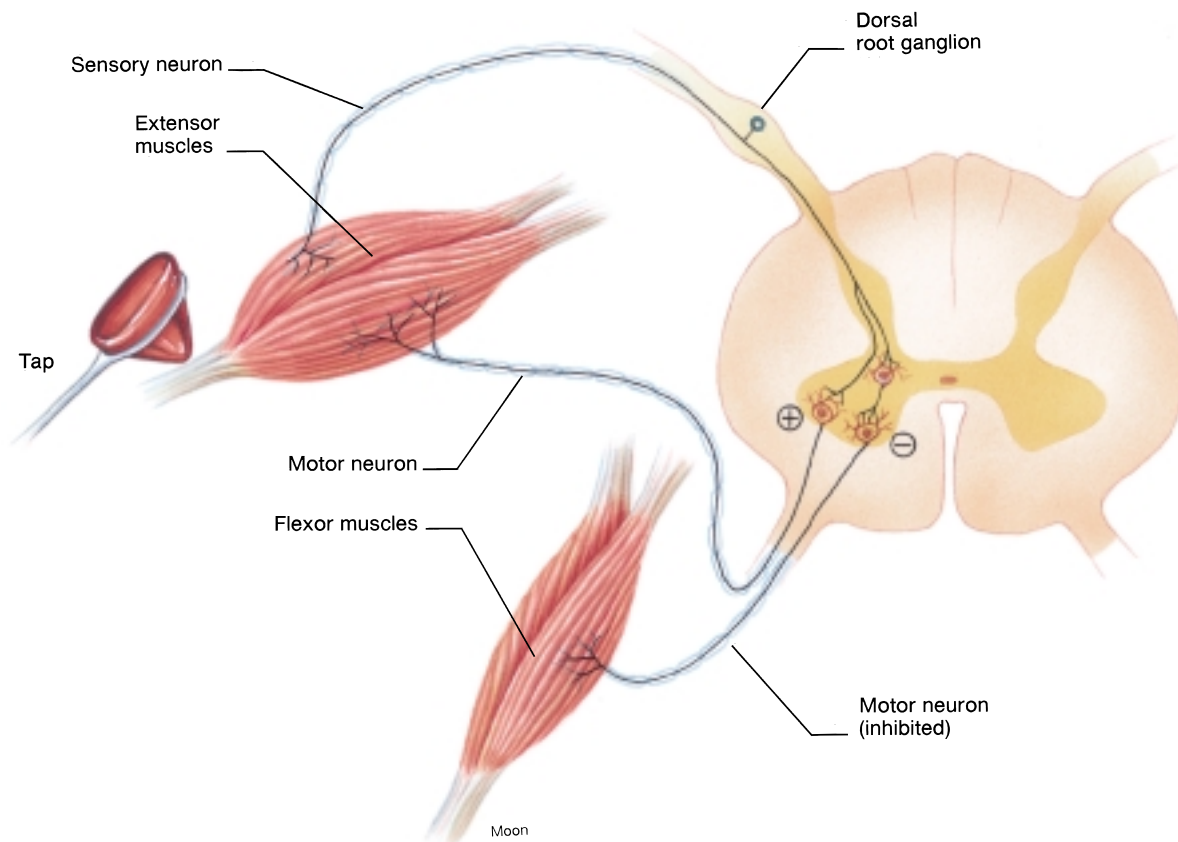
1. Passive stretch of a muscle (produced by tapping its tendon) stretches the spindle (intrafusal) fibers.
2. Stretching of a spindle distorts its central (bag or chain) region, which stimulates dendritic endings of sensory nerves.
3. Action potentials are conducted by afferent (sensory) nerve fibers into the spinal cord on the dorsal roots of spinal nerves.
4. Axons of sensory neurons synapse with dendrites and cell bodies of somatic motor neurons located in the ventral horn gray matter of the spinal cord.
5. Efferent nerve impulses in the axons of somatic motor neurons (which form the ventral roots of spinal nerves) are conducted to the ordinary (extrafusal) muscle fibers. These neurons are alpha motoneurons.
6. Release of acetylcholine from the endings of alpha motoneurons stimulates the contraction of the extrafusal fibers, and thus of the whole muscle.
7. Contraction of the muscle relieves the stretch of its spindles, thus decreasing electrical activity in the spindle afferent nerve fibers.



■ **Figure 12.25** The knee-jerk reflex. This is an example of a monosynaptic stretch reflex.



■ **Figure 12.26** The action of the Golgi tendon organ. An increase in muscle tension stimulates the activity of sensory nerve endings in the Golgi tendon organ. This sensory input stimulates an interneuron, which in turn inhibits the activity of a motor neuron innervating that muscle. This is therefore a disynaptic reflex.



■ **Figure 12.27** A diagram of reciprocal innervation. Afferent impulses from muscle spindles stimulates alpha motoneurons to the agonists muscle (the extensor) directly, but (via an inhibitory interneuron) they inhibit activity in the alpha motoneuron to the antagonist muscle.

interneurons within the spinal cord via collateral branches. These interneurons inhibit the motor neurons of antagonist muscles via inhibitory postsynaptic potentials (IPSPs). This dual stimulatory and inhibitory activity is called **reciprocal innervation** (fig. 12.27).

When a limb is flexed, for example, the antagonistic extensor muscles are passively stretched. Extension of a limb similarly stretches the antagonistic flexor muscles. If the monosynaptic stretch reflexes were not inhibited, reflex contraction of the antagonistic muscles would always interfere with the intended movement. Fortunately, whenever the “intended,” or agonist muscles, are stimulated to contract, the alpha and gamma motoneurons that stimulate the antagonist muscles are inhibited.

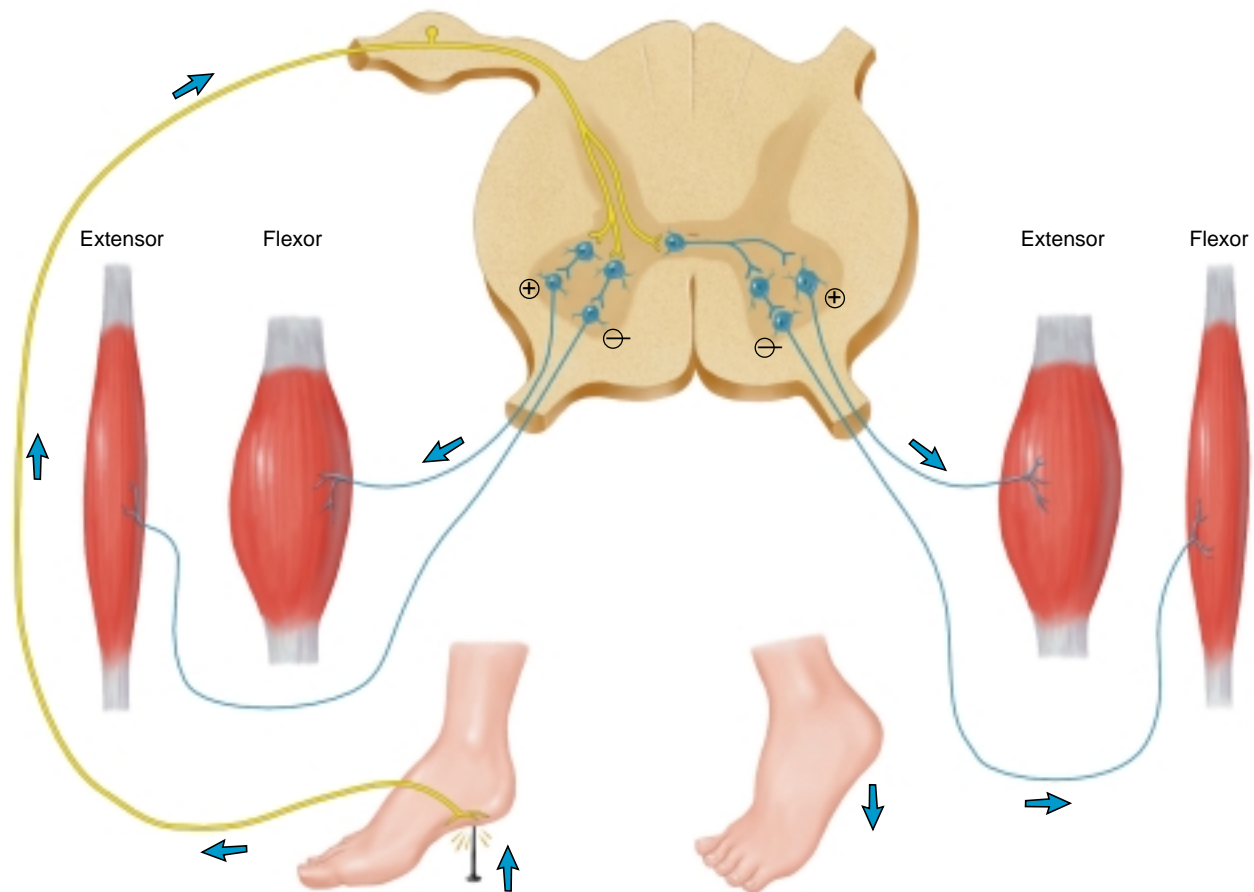
The stretch reflex, with its reciprocal innervations, involves the muscles of one limb only and is controlled by only one segment of the spinal cord. More complex reflexes involve muscles controlled by numerous spinal cord segments and affect muscles on the contralateral side of the cord. Such reflexes involve **double reciprocal innervation** of muscles.

Double reciprocal innervation is illustrated by the **crossed-extensor reflex**. If you step on a tack with your right foot, for example, this foot is withdrawn by contraction of the

flexors and relaxation of the extensors of your right leg. The contralateral left leg, by contrast, extends to help support your body during this withdrawal reflex. The extensors of your left leg contract while its flexors relax. These events are illustrated in figure 12.28.

## Upper Motor Neuron Control of Skeletal Muscles

As previously described, upper motor neurons are neurons in the brain that influence the control of skeletal muscle by lower motor neurons (alpha and gamma motoneurons). Neurons in the precentral gyrus of the cerebral cortex contribute axons that cross to the contralateral sides in the pyramids of the medulla oblongata; these tracts are thus called **pyramidal tracts** (chapter 8). The pyramidal tracts include the *lateral* and *ventral corticospinal tracts*. Neurons in other areas of the brain produce the **extrapyramidal tracts**. The major extrapyramidal tract is the *reticulospinal tract*, which originates in the reticular formation of the medulla oblongata and pons. Brain areas that influence the activity of extrapyramidal tracts are believed to produce the inhibition of lower motor neurons described in the preceding section.



■ **Figure 12.28** The crossed-extensor reflex. This complex reflex demonstrates double reciprocal innervation.

### Cerebellum

The **cerebellum**, like the cerebrum, receives sensory input from muscle spindles and Golgi tendon organs. It also receives fibers from areas of the cerebral cortex devoted to vision, hearing, and equilibrium.

There are no descending tracts from the cerebellum. The cerebellum can influence motor activity only indirectly, through its output to the vestibular nuclei, red nucleus, and basal nuclei. These structures, in turn, affect lower motor neurons via the vestibulospinal tract, rubrospinal tract, and reticulospinal tract. It is interesting that all output from the cerebellum is inhibitory; these inhibitory effects aid motor coordination by eliminating inappropriate neural activity. Damage to the cerebellum interferes with the ability to coordinate movements with spatial judgment. Under- or overreaching for an object may occur, followed by *intention tremor*, in which the limb moves back and forth in a pendulum-like motion.

### Basal Nuclei

The **basal nuclei**, sometimes called the **basal ganglia**, include the *caudate nucleus*, *putamen*, and *globus pallidus* (chapter 8). Often included in this group are other nuclei of the *thalamus*, *subthalamus*, *substantia nigra*, and *red nucleus*. Acting directly

via the rubrospinal tract and indirectly via synapses in the reticular formation and thalamus, the basal nuclei have profound effects on the activity of lower motor neurons.

In particular, through their synapses in the reticular formation, the basal nuclei exert an inhibitory influence on the activity of lower motor neurons. Damage to the basal nuclei thus results in increased muscle tone, as previously described. People with such damage display *akinesia*, lack of desire to use the affected limb, and *chorea*, sudden and uncontrolled random movements (table 12.9).



**Parkinson's disease** (or *paralysis agitans*) is a disorder of the basal nuclei involving degeneration of fibers from the substantia nigra. These fibers, which use dopamine as a neurotransmitter, are required to antagonize the effects of other fibers that use acetylcholine (ACh) as a transmitter. The relative deficiency of dopamine compared to ACh is believed to produce the symptoms of Parkinson's disease, including resting tremor. This "shaking" of the limbs tends to disappear during voluntary movements and then reappear when the limb is again at rest.

## Table 12.9 Symptoms of Upper Motor Neuron Damage

**Babinski's reflex**—Extension of the great toe when the sole of the foot is stroked along the lateral border

**Spastic paralysis**—High muscle tone and hyperactive stretch reflexes; flexion of arms and extension of legs

**Hemiplegia**—Paralysis of upper and lower limbs on one side—commonly produced by damage to motor tracts as they pass through internal capsule (such as by cerebrovascular accident—stroke)

**Paraplegia**—Paralysis of the lower limbs on both sides as a result of lower spinal cord damage

**Quadriplegia**—Paralysis of upper and lower limbs on both sides as a result of damage to the upper region of the spinal cord or brain

**Chorea**—Random uncontrolled contractions of different muscle groups (as in Saint Vitus' dance) as a result of damage to basal nuclei

**Resting tremor**—Shaking of limbs at rest; disappears during voluntary movements; produced by damage to basal nuclei

**Intention tremor**—Oscillations of the arm following voluntary reaching movements; produced by damage to cerebellum

## Test Yourself Before You Continue

1. Draw a muscle spindle surrounded by a few extrafusal fibers. Indicate the location of primary and secondary sensory endings and explain how these endings respond to muscle stretch.
2. Describe all of the events that occur from the time the patellar tendon is struck with a mallet to the time the leg kicks.
3. Explain how a Golgi tendon organ is stimulated and describe the disynaptic reflex that occurs.
4. Explain the significance of reciprocal innervation and double reciprocal innervation in muscle reflexes.
5. Describe the functions of gamma motoneurons and explain why they are stimulated at the same time as alpha motoneurons during voluntary muscle contractions.
6. Explain how a person with spinal cord damage might develop clonus.

## Cardiac and Smooth Muscles

Cardiac muscle, like skeletal muscle, is striated and contains sarcomeres that shorten by sliding of thin and thick filaments. But while skeletal muscle requires nervous stimulation to contract, cardiac muscle can produce impulses and contract spontaneously. Smooth muscles lack sarcomeres, but they do contain actin and myosin that produce contractions in response to a unique regulatory mechanism.

Unlike skeletal muscles, which are voluntary effectors regulated by somatic motor neurons, cardiac and smooth muscles are involuntary effectors regulated by autonomic motor neurons. Although there are important differences between skeletal mus-

cle and cardiac and smooth muscle, there are also significant similarities. All types of muscle are believed to contract by means of sliding of thin filaments over thick filaments. The sliding of the filaments is produced by the action of myosin cross bridges in all types of muscles, and excitation-contraction coupling in all types of muscles involves  $\text{Ca}^{2+}$ .

## Cardiac Muscle

Like skeletal muscle cells, cardiac (heart) muscle cells, or **myocardial cells**, are striated; they contain actin and myosin filaments arranged in the form of sarcomeres, and they contract by means of the sliding filament mechanism. The long, fibrous skeletal muscle cells, however, are structurally and functionally separated from each other, whereas the myocardial cells are short, branched, and interconnected. Each myocardial cell is tubular in structure and joined to adjacent myocardial cells by electrical synapses, or **gap junctions** (see chapter 7, fig. 7.19).

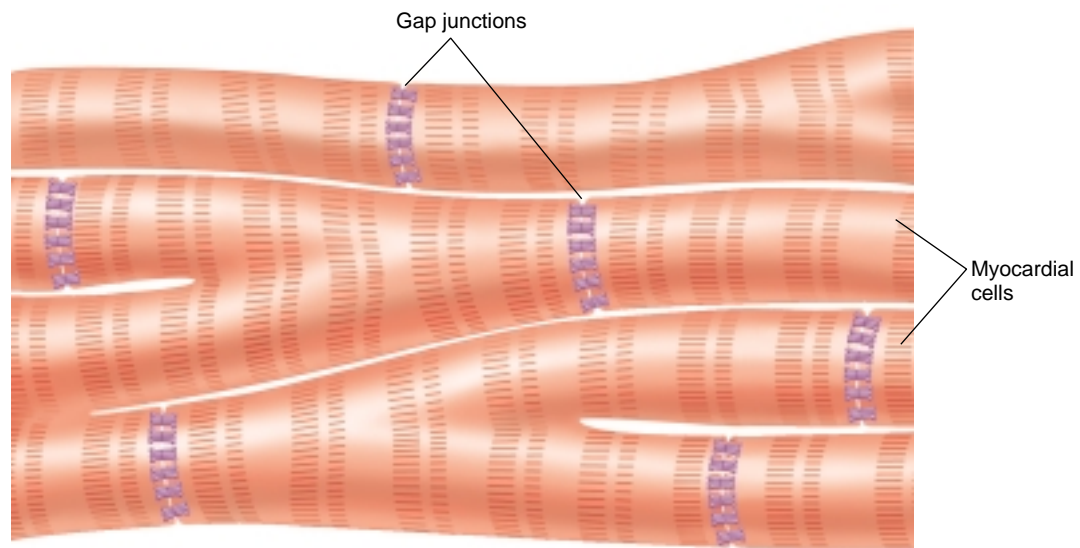
The gap junctions are concentrated at the ends of each myocardial cell (fig. 12.29), which permits electrical impulses to be conducted primarily along the long axis from cell to cell. Gap junctions in cardiac muscle have an affinity for stain that makes them appear as dark lines between adjacent cells when viewed in the light microscope. These dark-staining lines are known as *intercalated discs* (fig. 12.30).

Electrical impulses that originate at any point in a mass of myocardial cells, called a **myocardium**, can spread to all cells in the mass that are joined by gap junctions. Because all cells in a myocardium are electrically joined, a myocardium behaves as a single functional unit. Thus, unlike skeletal muscles that produce contractions that are graded depending on the number of cells stimulated, a myocardium contracts to its full extent each time because all of its cells contribute to the contraction. The ability of the myocardial cells to contract, however, can be increased by the hormone epinephrine and by stretching of the heart chambers. The heart contains two distinct myocardia (atria and ventricles), as will be described in chapter 13.

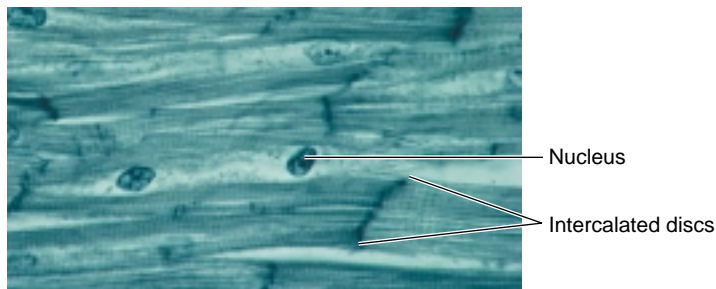
Unlike skeletal muscles, which require external stimulation by somatic motor nerves before they can produce action potentials and contract, cardiac muscle is able to produce action potentials automatically. Cardiac action potentials normally originate in a specialized group of cells called the *pacemaker*. However, the rate of this spontaneous depolarization, and thus the rate of the heartbeat, are regulated by autonomic innervation. Regulation of the cardiac rate is described more fully in chapter 14.

## Smooth Muscle

Smooth (visceral) muscles are arranged in circular layers in the walls of blood vessels and bronchioles (small air passages in the lungs). Both circular and longitudinal smooth muscle layers occur in the tubular digestive tract, the ureters (which transport urine), the ductus deferentia (which transport sperm cells), and the uterine tubes (which transport ova). The alternate contraction of circular and longitudinal smooth muscle layers in the intestine produces **peristaltic waves**, which propel the contents of these tubes in one direction.



■ **Figure 12.29** Myocardial cells are interconnected by gap junctions. The gap junctions are fluid-filled channels through the plasma membrane of adjacent cells that permit the conduction of impulses from one cell to the next. The gap junctions are concentrated at the ends of each myocardial cell, and each gap junction is composed of connexin proteins (also see chapter 7, fig. 7.19).



■ **Figure 12.30** Cardiac muscle. Notice that the cells are short, branched, and striated and that they are interconnected by intercalated discs.

Although smooth muscle cells do not contain sarcomeres (which produce striations in skeletal and cardiac muscle), they do contain a great deal of actin and some myosin, which produces a ratio of thin to thick filaments of about 16 to 1 (in striated muscles the ratio is 2 to 1). Unlike striated muscles, in which the thin filaments are relatively short (extending from a Z disc into a sarcomere), the thin filaments of smooth muscle cells are quite long. They attach either to regions of the plasma membrane of the smooth muscle cell or to cytoplasmic protein structures called **dense bodies**, which are analogous to the Z discs of striated muscle (fig. 12.31*b*).

In smooth muscle, the myosin proteins of the thick filaments are stacked vertically so that their long axis is perpendicular to the long axis of the thick filament (fig. 12.31*c*). In this way, the myosin heads can form cross bridges with actin all along the length of the thick filaments. This is different from the horizontal arrangement of myosin proteins in the thick filaments

of striated muscles (see fig. 12.10), which is required to cause the shortening of sarcomeres.

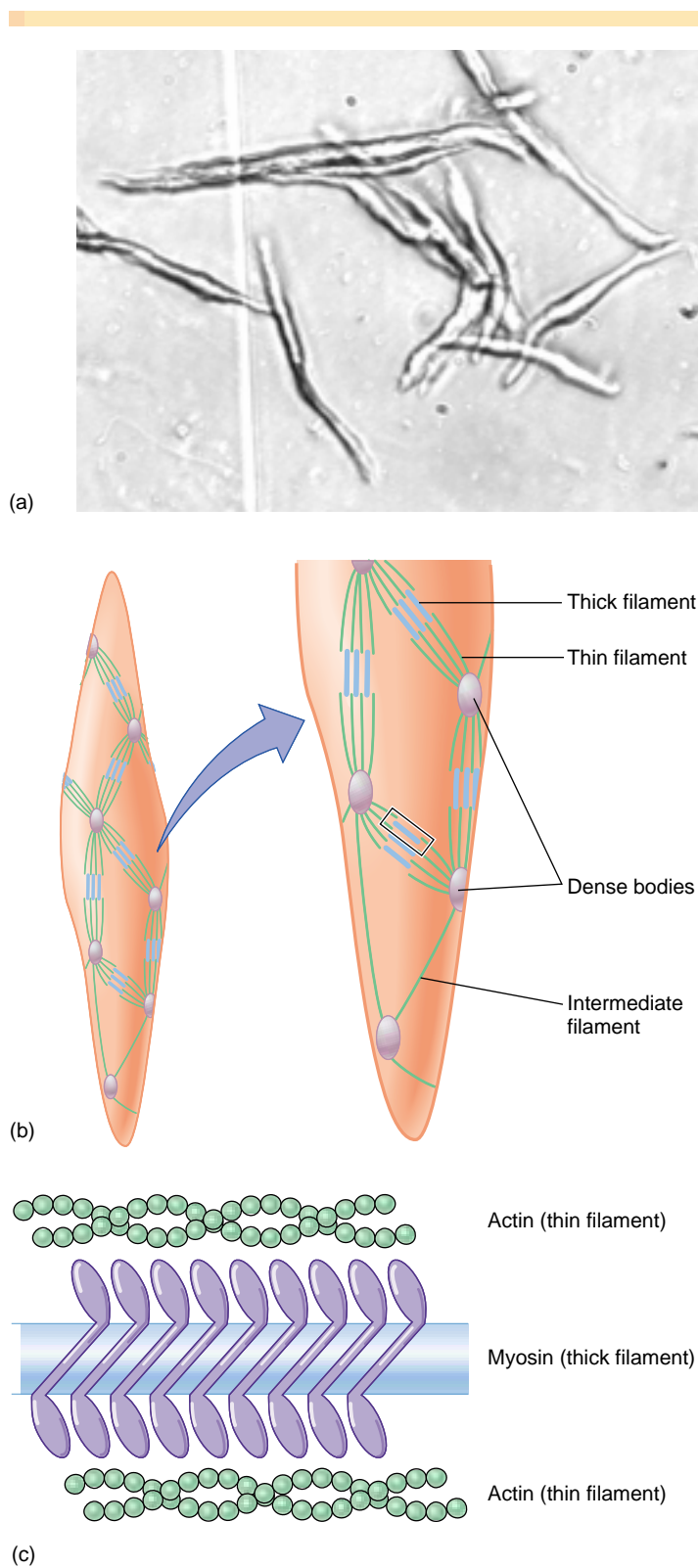
The arrangement of the contractile apparatus in smooth muscle cells, and the fact that it is not organized into sarcomeres, is required for proper smooth muscle function. Smooth muscles must be able to contract even when greatly stretched—in the urinary bladder, for example, the smooth muscle cells may be stretched up to two and a half times their resting length. The smooth muscle cells of the uterus may be stretched up to eight times their original length by the end of pregnancy. Striated muscles, because of their structure, lose their ability to contract when the sarcomeres are stretched to the point where actin and myosin no longer overlap.

### **Excitation-Contraction Coupling in Smooth Muscles**

As in striated muscles, the contraction of smooth muscles is triggered by a sharp rise in the  $\text{Ca}^{2+}$  concentration within the cytoplasm of the muscle cells. However, the sarcoplasmic reticulum of smooth muscles is less developed than that of skeletal muscles, and  $\text{Ca}^{2+}$  released from this organelle may account for only the initial phase of smooth muscle contraction. Extracellular  $\text{Ca}^{2+}$  diffusing into the smooth muscle cell through its plasma membrane is responsible for sustained contractions. This  $\text{Ca}^{2+}$  enters primarily through voltage-regulated  $\text{Ca}^{2+}$  channels in the plasma membrane. The opening of these channels is graded by the amount of depolarization; the greater the depolarization, the more  $\text{Ca}^{2+}$  will enter the cell and the stronger will be the smooth muscle contraction.

The events that follow the entry of  $\text{Ca}^{2+}$  into the cytoplasm are somewhat different in smooth muscles than in striated muscles. In striated muscles,  $\text{Ca}^{2+}$  combines with troponin. Troponin, however, is not present in smooth muscle cells. In smooth muscles,  $\text{Ca}^{2+}$  combines with a protein in the cytoplasm called **calmodulin**, which is structurally similar to troponin. Calmodulin





**Figure 12.31** Smooth muscle and its contractile apparatus. (a) A photomicrograph of freshly isolated vascular smooth muscle cells (400 $\times$ ). (b) Arrangement of thick and thin filaments in smooth muscles. Note that dense bodies are also interconnected by intermediate fibers. (c) The myosin proteins are staked in a different arrangement in smooth muscles than in striated muscles.

was previously discussed in relation to the function of  $\text{Ca}^{2+}$  as a second messenger in hormone action (chapter 11). The calmodulin- $\text{Ca}^{2+}$  complex thus formed combines with and activates **myosin light-chain kinase (MLCK)**, an enzyme that catalyzes the phosphorylation (addition of phosphate groups) of *myosin light chains*, a component of the myosin cross bridges. In smooth muscle (unlike striated muscle), the phosphorylation of myosin cross bridges is the regulatory event that permits them to bind to actin and thereby produce a contraction (fig. 12.32).

Unlike the situation in striated muscle cells, which produce all-or-none action potentials, smooth muscle cells can produce graded depolarizations and contractions without producing action potentials. Indeed, only these graded depolarizations are conducted from cell to cell in many smooth muscles. The greater the depolarization of a smooth muscle cell, the more  $\text{Ca}^{2+}$  will enter, and the more MLCK enzymes will be activated. With more MLCK enzymes activated, more cross bridges will become phosphorylated and able to bind to actin. In this way, a stronger depolarization of the smooth muscle cell leads to a stronger contraction.

Relaxation of the smooth muscle follows the closing of the  $\text{Ca}^{2+}$  channels and lowering of the cytoplasmic  $\text{Ca}^{2+}$  concentrations. Under these conditions, calmodulin dissociates from the myosin light-chain kinase, thereby inactivating this enzyme. The phosphate groups that were added to the myosin are then removed by a different enzyme, a *myosin phosphatase* (fig. 12.32). Dephosphorylation inhibits the cross bridge from binding to actin and producing another power stroke.

In addition to being graded, the contractions of smooth muscle cells are slow and sustained. The slowness of contraction is related to the fact that myosin ATPase in smooth muscle is slower in its action (splitting ATP for the cross-bridge cycle) than it is in striated muscle. The sustained nature of smooth muscle contraction is explained by the theory that cross bridges in smooth muscles can enter a *latch state*.

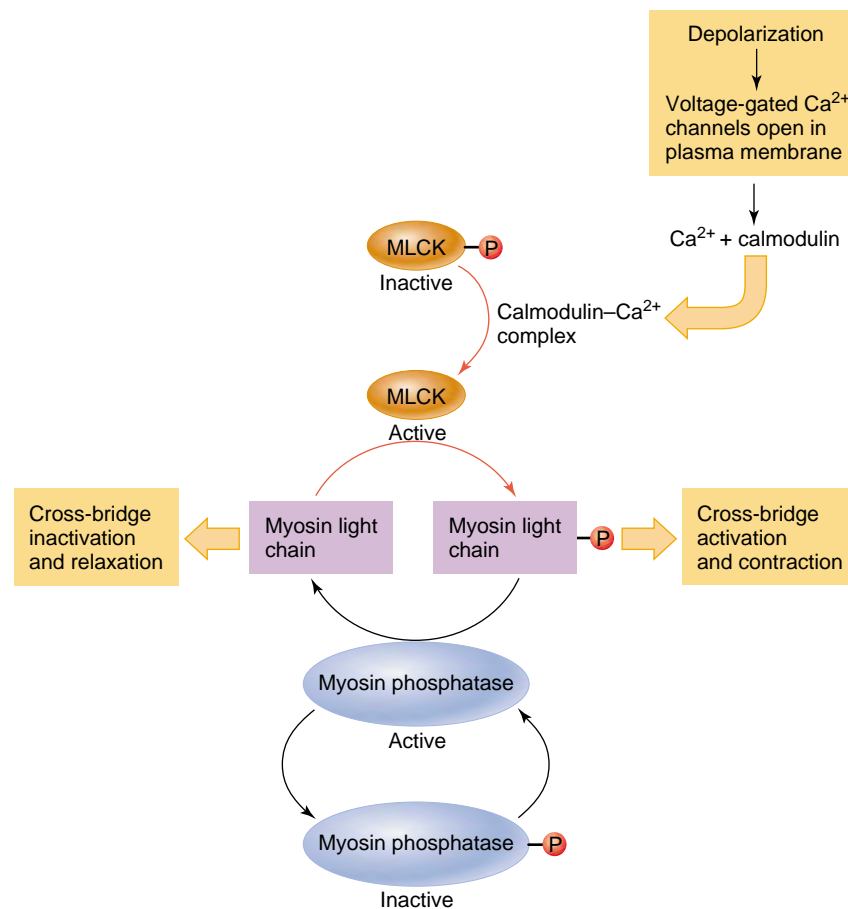
The latch state allows smooth muscle to maintain its contraction in a very energy-efficient manner, hydrolyzing less ATP than would otherwise be required. This ability is obviously important for smooth muscles, given that they encircle the walls of hollow organs and must sustain contractions for long periods of time. The mechanisms by which the latch state is produced, however, are complex and poorly understood.

The three muscle types—skeletal, cardiac, and smooth—are compared in table 12.10.



Drugs such as *nifedipine* and related newer compounds are **calcium channel blockers**. These drugs block  $\text{Ca}^{2+}$  channels in the membrane of smooth muscle cells within the walls of blood vessels, causing the muscles

to relax and the vessels to dilate. This effect, called *vasodilation*, may be helpful in treating some cases of hypertension (high blood pressure). Calcium-channel-blocking drugs are also used when spasm of the coronary arteries (*vasospasm*) produces *angina pectoris*, which is pain caused by insufficient blood flow to the heart.



■ **Figure 12.32** Excitation-contraction coupling in smooth muscle. When Ca<sup>2+</sup> passes through voltage-gated channels in the plasma membrane it enters the cytoplasm and binds to calmodulin. The calmodulin-Ca<sup>2+</sup> complex then activates myosin light-chain kinase (MLCK) by removing a phosphate group. The activated MLCK, in turn, phosphorylates the myosin light chains, thereby activating the cross bridges to cause contraction. Contraction is ended when myosin phosphatase becomes activated. Upon its activation, myosin phosphatase removes the phosphates from the myosin light chains and thereby inactivates the cross bridges.

**Table 12.10** Comparison of Skeletal, Cardiac, and Smooth Muscle

Skeletal Muscle	Cardiac Muscle	Smooth Muscle
Striated; actin and myosin arranged in sarcomeres	Striated; actin and myosin arranged in sarcomeres	Not striated; more actin than myosin; actin inserts into dense bodies and cell membrane
Well-developed sarcoplasmic reticulum and transverse tubules	Moderately developed sarcoplasmic reticulum and transverse tubules	Poorly developed sarcoplasmic reticulum; no transverse tubules
Contains troponin in the thin filaments	Contains troponin in the thin filaments	Contains calmodulin, a protein that, when bound to Ca <sup>2+</sup> , activates the enzyme myosin light-chain kinase
Ca <sup>2+</sup> released into cytoplasm from sarcoplasmic reticulum	Ca <sup>2+</sup> enters cytoplasm from sarcoplasmic reticulum and extracellular fluid	Ca <sup>2+</sup> enters cytoplasm from extracellular fluid, sarcoplasmic reticulum, and perhaps mitochondria
Cannot contract without nerve stimulation; denervation results in muscle atrophy	Can contract without nerve stimulation; action potentials originate in pacemaker cells of heart	Maintains tone in absence of nerve stimulation; visceral smooth muscle produces pacemaker potentials; denervation results in hypersensitivity to stimulation
Muscle fibers stimulated independently; no gap junctions	Gap junctions present as intercalated discs	Gap junctions generally present

## Clinical Investigation Clues

Remember that Maria was taking a calcium channel-blocking drug to treat her hypertension.

How do such drugs help to lower blood pressure?

Is it likely that this drug contributed to Maria's skeletal muscle pain and fatigue?

Could it raise her blood  $\text{Ca}^{2+}$  levels?

If not, what could raise her blood  $\text{Ca}^{2+}$ ?

### Single-Unit and Multiunit Smooth Muscles

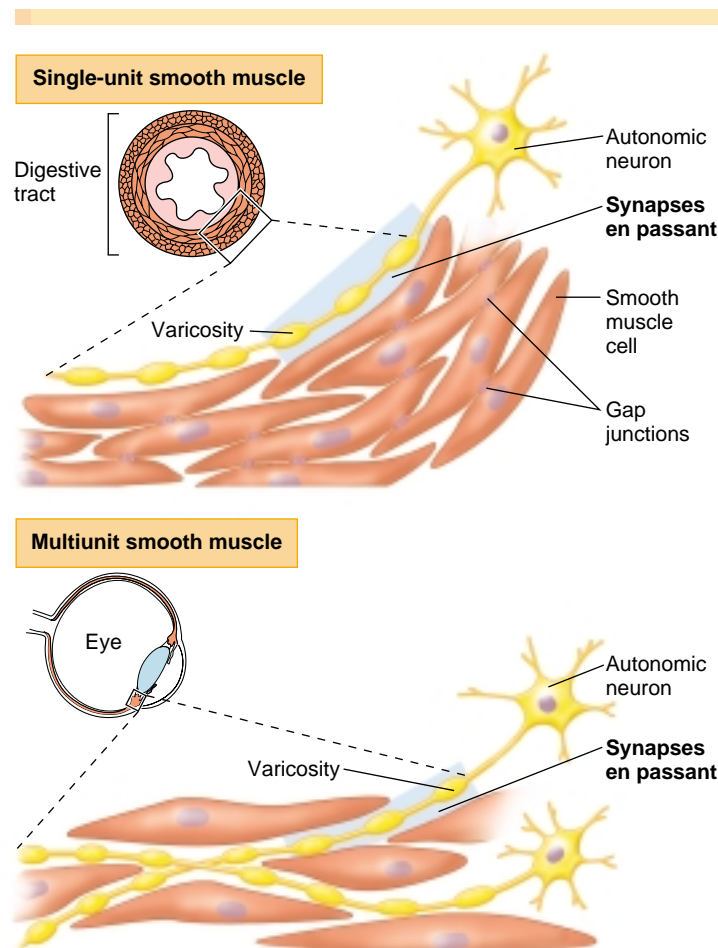
Smooth muscles are often grouped into two functional categories: **single-unit** and **multiunit** (fig. 12.33). Single-unit smooth muscles have numerous gap junctions (electrical synapses) between adjacent cells that weld them together electrically; they thus behave as a single unit, much like cardiac muscle. Most smooth muscles—including those in the digestive tract and uterus—are single-unit.

Only some cells of single-unit smooth muscles receive autonomic innervation, but the ACh released by the axon can diffuse to other smooth muscle cells. Binding of ACh to its muscarinic receptors causes depolarization by closing  $\text{K}^+$  channels, as described in chapter 7. Such stimulation, however, only modifies the automatic behavior of single-unit smooth muscles. Single-unit smooth muscles display *pacemaker* activity, in which certain cells stimulate others in the mass. This is similar to the situation in cardiac muscle. Single-unit smooth muscles also display intrinsic, or *myogenic*, electrical activity and contraction in response to stretch. For example, the stretch induced by an increase in the volume of a ureter or a section of the digestive tract can stimulate myogenic contraction. Such contraction does not require stimulation by autonomic nerves.

Contraction of multiunit smooth muscles, by contrast, requires nerve stimulation. Multiunit smooth muscles have few, if any, gap junctions. The cells must thus be stimulated individually by nerve fibers. Examples of multiunit smooth muscles are the arrector pili muscles in the skin and the ciliary muscles attached to the lens of the eye.

### Autonomic Innervation of Smooth Muscles

The neural control of skeletal muscles differs significantly from that of smooth muscles. A skeletal muscle fiber has only one junction with a somatic nerve fiber, and the receptors for the neurotransmitter are located only at the neuromuscular junction. By contrast, the entire surface of smooth muscle cells contains neurotransmitter receptor proteins. Neurotransmitter molecules are released along a stretch of an autonomic nerve fiber that is located some distance from the smooth muscle cells. The regions of the autonomic fiber that release transmitters appear as bulges, or *varicosities*, and the neurotransmitters released from these varicosities stimulate a number of smooth muscle cells. Since there are numerous varicosities along a stretch of an autonomic nerve ending, they form synapses “in passing”—or *synapses en passant*—with the smooth muscle cells (fig. 12.33).



**Figure 12.33** Single-unit and multiunit smooth muscle. In single-unit smooth muscle, the individual smooth muscle cells are electrically joined by gap junctions, so that depolarizations can spread from one cell to the next. In multiunit smooth muscle, each smooth muscle cell must be stimulated by an axon. The axons of autonomic neurons have varicosities, which release neurotransmitters, and which form *synapses en passant* with the smooth muscle cells.

## Test Yourself Before You Continue

1. Explain how cardiac muscle differs from skeletal muscle in its structure and regulation of contraction.
2. Contrast the structure of a smooth muscle cell with that of a skeletal muscle fiber and discuss the advantages of each type of structure.
3. Describe the events by which depolarization of a smooth muscle cell results in contraction and explain why smooth muscle contractions are slow and sustained.
4. Distinguish between single-unit and multiunit smooth muscles.

# INTERACTIONS

## HPer Links of the Muscular System with Other Body Systems

### Integumentary System

- The skin helps to protect all organs of the body from invasion by pathogens . . . (p. 000)
- The smooth muscles of cutaneous blood vessels are needed for the regulation of cutaneous blood flow . . . . . (p. 000)
- The arrector pili muscles in the skin produce goose bumps . . . . . (p. 000)

### Skeletal System

- Bones store calcium, which is needed for the control of muscle contraction . . . . (p. 000)
- The skeleton provides attachment sites for muscles . . . . . (p. 000)
- Joints of the skeleton provide levers for movement . . . . . (p. 000)
- Muscle contractions maintain the health and strength of bone . . . . . (p. 000)

### Nervous System

- Somatic motor neurons stimulate contraction of skeletal muscles . . . (p. 000)
- Autonomic neurons stimulate smooth muscle contraction or relaxation . . (p. 000)
- Autonomic nerves increase cardiac output during exercise . . . . . (p. 000)
- Sensory neurons from muscles monitor muscle length and tension . . . . . (p. 000)

### Endocrine System

- Sex hormones promote muscle development and maintenance . . . . . (p. 000)
- Parathyroid hormone and other hormones regulate blood calcium and phosphate concentrations . . . . . (p. 000)

- Epinephrine and norepinephrine influence contractions of cardiac muscle and smooth muscles . . . . . (p. 000)
- Insulin promotes glucose entry into skeletal muscles . . . . . (p. 000)
- Adipose tissue secretes hormones that regulate the sensitivity of muscles to insulin . . . . . (p. 000)

### Circulatory System

- Blood transports O<sub>2</sub> and nutrients to muscles and removes CO<sub>2</sub> and lactic acid . . . . . (p. 000)
- Contractions of skeletal muscles serve as a pump to assist blood movement within veins . . . . . (p. 000)
- Cardiac muscle enables the heart to function as a pump . . . . . (p. 000)
- Smooth muscle enables blood vessels to constrict and dilate . . . . . (p. 000)

### Respiratory System

- The lungs provide oxygen for muscle metabolism and eliminate carbon dioxide . . . . . (p. 000)
- Respiratory muscles enable ventilation of the lungs . . . . . (p. 000)

### Urinary System

- The kidneys eliminate creatinine and other metabolic wastes from muscle . . . . . (p. 000)
- The kidneys help to regulate the blood calcium and phosphate concentrations . . . . . (p. 000)

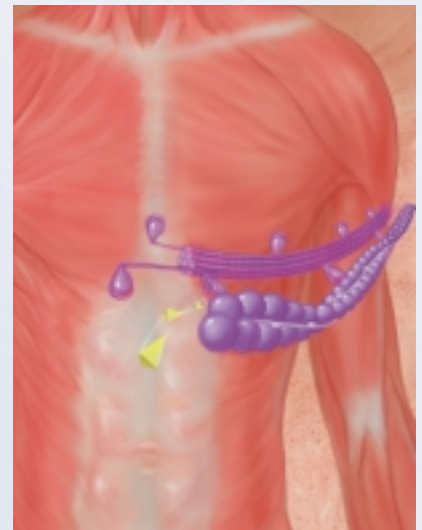
- Muscles of the urinary tract are needed for the control of urination . . . . . (p. 000)

### Digestive System

- The GI tract provides nutrients for all body organs, including muscles . . . . . (p. 000)
- Smooth muscle contractions push digestion products along the GI tract . . . . . (p. 000)
- Muscular sphincters of the GI tract help to regulate the passage of food . . . . . (p. 000)

### Reproductive System

- Testicular androgen promotes growth of skeletal muscle . . . . . (p. 000)
- Muscle contractions contribute to orgasm in both sexes . . . . . (p. 000)
- Uterine muscle contractions are required for vaginal delivery of a fetus . . . . . (p. 000)



## Summary

### Skeletal Muscles 326

- I. Skeletal muscles are attached to bones by tendons.
  - A. Skeletal muscles are composed of separate cells, or fibers, that are attached in parallel to the tendons.
  - B. Individual muscle fibers are covered by the endomysium; bundles of fibers, called fascicles, are covered by the perimysium; and the entire muscle is covered by the epimysium.
  - C. Skeletal muscle fibers are striated.
    1. The dark striations are called A bands, and the light regions are called I bands.
    2. Z lines are located in the middle of each I band.
- II. The contraction of muscle fibers *in vivo* is stimulated by somatic motor neurons.
  - A. Each somatic motor axon branches to innervate numerous muscle fibers.
  - B. The motor neuron and the muscle fibers it innervates are called a motor unit.
    1. When a muscle is composed of a relatively large number of motor units (such as in the hand), there is fine control of muscle contraction.
    2. The large muscles of the leg have relatively few motor units, which are correspondingly large in size.
    3. Sustained contractions are produced by the asynchronous stimulation of different motor units.

### Mechanisms of Contraction 331

- I. Skeletal muscle cells, or fibers, contain structures called myofibrils.
  - A. Each myofibril is striated with dark (A) and light (I) bands. In the middle of each I band are Z lines.
  - B. The A bands contain thick filaments, composed primarily of myosin.
    1. The edges of each A band also contain thin filaments, which overlap the thick filaments.
2. The central regions of the A bands contain only thick filaments—these regions are the H bands.
- C. The I bands contain only thin filaments, composed primarily of actin.
- D. Thin filaments are composed of globular actin subunits known as G-actin. A protein known as tropomyosin is also located at intervals in the thin filaments. Another protein—troponin—is attached to the tropomyosin.
- II. Myosin cross bridges extend out from the thick filaments to the thin filaments.
  - A. At rest, the cross bridges are not attached to actin.
    1. The cross-bridge heads function as ATPase enzymes.
    2. ATP is split into ADP and  $P_i$ , activating the cross bridge.
  - B. When the activated cross bridges attach to actin, they release  $P_i$  and undergo a power stroke.
  - C. At the end of a power stroke, the cross bridge releases the ADP and binds to a new ATP.
    1. This allows the cross bridge to detach from actin and repeat the cycle.
    2. Rigor mortis is caused by the inability of cross bridges to detach from actin because of a lack of ATP.
- III. The activity of the cross bridges causes the thin filaments to slide toward the centers of the sarcomeres.
  - A. The filaments slide—they do not shorten—during muscle contraction.
  - B. The lengths of the H and I bands decrease, whereas the A bands stay the same length during contraction.
- IV. When a muscle is at rest, the  $Ca^{2+}$  concentration of the sarcoplasm is very low and cross bridges are prevented from attaching to actin.
  - A. The  $Ca^{2+}$  is actively transported into the sarcoplasmic reticulum.
  - B. The sarcoplasmic reticulum is a modified endoplasmic reticulum that surrounds the myofibrils.
- V. Action potentials are conducted by transverse tubules into the muscle fiber.
  - A. Transverse tubules are invaginations of the cell membrane that almost touch the sarcoplasmic reticulum.
  - B. Action potentials in the transverse tubules stimulate the release of  $Ca^{2+}$  from the sarcoplasmic reticulum.
- VI. When action potentials cease,  $Ca^{2+}$  is removed from the sarcoplasm and stored in the sarcoplasmic reticulum.

### Contractions of Skeletal Muscles 339

- I. Muscles *in vitro* can exhibit twitch, summation, and tetanus.
  - A. The rapid contraction and relaxation of muscle fibers is called a twitch.
  - B. A whole muscle also produces a twitch in response to a single electrical pulse *in vitro*.
    1. The stronger the electric shock, the stronger the muscle twitch—whole muscles can produce graded contractions.
    2. The graded contraction of whole muscles is due to different numbers of fibers participating in the contraction.
  - C. The summation of fiber twitches can occur so rapidly that the muscle produces a smooth, sustained contraction known as tetanus.
  - D. When a muscle exerts tension without shortening, the contraction is termed isometric; when shortening does occur, the contraction is isotonic.
- II. The series-elastic component refers to the elastic composition of the muscle and its associated structures, which must be stretched tight before the tension exerted by the muscle can cause movement.
- III. The strength of a muscle contraction is dependant upon its resting length.
  - A. If the muscle is too short or too long prior to stimulation, the filaments in the sarcomeres will not have an optimum amount of overlap.

- B. At its normal resting length in vivo, a muscle is at its optimum length for contraction.

### Energy Requirements of Skeletal Muscles 342

- I. Aerobic cell respiration is ultimately required for the production of ATP needed for cross-bridge activity.
  - A. Resting muscles and muscles performing light exercise obtain most of their energy from fatty acids.
  - B. During moderate exercise, just below the lactate threshold, energy is obtained about equally from fatty acids and glucose.
  - C. Glucose, from the muscle's stored glycogen and from blood plasma, becomes an increasingly important energy source during heavy exercise.
  - D. New ATP can be quickly produced from the combination of ADP with phosphate derived from phosphocreatine.
  - E. Muscle fibers are of three types.
    - 1. Slow-twitch red fibers are adapted for aerobic respiration and are resistant to fatigue.
    - 2. Fast-twitch white fibers are adapted for anaerobic respiration.
    - 3. Intermediate fibers are fast-twitch but adapted for aerobic respiration.
- II. Muscle fatigue may be caused by a number of mechanisms.
  - A. Fatigue during sustained maximal contraction may be produced by the accumulation of extracellular  $K^+$  as a result of high levels of nerve activity.
  - B. Fatigue during moderate exercise is primarily a result of anaerobic respiration by fast-twitch fibers.
    - 1. The production of lactic acid lowers the intracellular pH, which inhibits glycolysis and decreases ATP concentrations.
    - 2. Decreased ATP inhibits excitation-contraction coupling, possibly due to a cellular loss of  $Ca^{2+}$ .
- III. Physical training affects the characteristics of the muscle fibers.
  - A. Endurance training increases the aerobic capacity of muscle fibers and their use of fatty acids for energy, so that their reliance on glycogen and anaerobic respiration—and thus their susceptibility to fatigue—is reduced.
  - B. Resistance training causes hypertrophy of muscle fibers because of an increase in the size and number of myofibrils.

### Neural Control of Skeletal Muscles 347

- I. The somatic motor neurons that innervate the muscles are called lower motor neurons.
  - A. Alpha motoneurons innervate the ordinary, or extrafusal, muscle fibers. These are the fibers that produce muscle shortening during contraction.
  - B. Gamma motoneurons innervate the intrafusal fibers of the muscle spindles.
- II. Muscle spindles function as length detectors in muscles.
  - A. Spindles consist of several intrafusal fibers wrapped together. The spindles are in parallel with the extrafusal fibers.
  - B. Stretching of the muscle stretches the spindles, which excites sensory endings in the spindle apparatus.
    - 1. Impulses in the sensory neurons travel into the spinal cord in the dorsal roots of spinal nerves.
    - 2. The sensory neuron synapses directly with an alpha motoneuron within the spinal cord, which produces a monosynaptic reflex.
    - 3. The alpha motoneuron stimulates the extrafusal muscle fibers to contract, thus relieving the stretch. This is called the stretch reflex.
  - C. The activity of gamma motoneurons tightens the spindles, thus making them more sensitive to stretch and better able to monitor the length of the muscle, even during muscle shortening.
- III. The Golgi tendon organs monitor the tension that the muscle exerts on its tendons.
  - A. As the tension increases, sensory neurons from Golgi tendon organs inhibit the activity of alpha motoneurons.
- B. This is a disynaptic reflex because the sensory neurons synapse with interneurons, which in turn make inhibitory synapses with motoneurons.
- IV. A crossed-extensor reflex occurs when a foot steps on a tack.
  - A. Sensory input from the injured foot causes stimulation of flexor muscles and inhibition of the antagonistic extensor muscles.
  - B. The sensory input also crosses the spinal cord to cause stimulation of extensor and inhibition of flexor muscles in the contralateral leg.
- V. Most of the fibers of descending tracts synapse with spinal interneurons, which in turn synapse with the lower motor neurons.
  - A. Alpha and gamma motoneurons are usually stimulated at the same time, or coactivated.
  - B. The stimulation of gamma motoneurons keeps the muscle spindles under tension and sensitive to stretch.
  - C. Upper motor neurons, primarily in the basal nuclei, also exert inhibitory effects on gamma motoneurons.
- VI. Neurons in the brain that affect the lower motor neurons are called upper motor neurons.
  - A. The fibers of neurons in the precentral gyrus, or motor cortex, descend to the lower motor neurons as the lateral and ventral corticospinal tracts.
    - 1. Most of these fibers cross to the contralateral side in the brain stem, forming structures called the pyramids; therefore, this system is called the pyramidal system.
    - 2. The left side of the brain thus controls the musculature on the right side, and vice versa.
  - B. Other descending motor tracts are part of the extrapyramidal system.
    - 1. The neurons of the extrapyramidal system make numerous synapses in different areas of the brain, including the midbrain, brain stem, basal nuclei, and cerebellum.

2. Damage to the cerebellum produces intention tremor, and degeneration of dopaminergic neurons in the basal nuclei produces Parkinson's disease.

### Cardiac and Smooth Muscles 354

- I. Cardiac muscle is striated and contains sarcomeres.
  - A. In contrast to skeletal muscles, which require neural stimulation to contract, action potentials in the heart originate in myocardial cells; stimulation by neurons is not required.
  - B. Also unlike the situation in skeletal muscles, action potentials can cross from one myocardial cell to another.
- II. The thin and thick filaments in smooth muscles are not organized into sarcomeres.
  - A. The thin filaments extend from the plasma membrane and from dense bodies in the cytoplasm.
  - B. The myosin proteins are stacked perpendicular to the long axis of the thick filaments, so they can bind to actin all along the length of the thick filament.
  - C. Depolarizations are graded and conducted from one smooth muscle cell to the next.
    1. The depolarizations stimulate the entry of  $\text{Ca}^{2+}$ , which binds to calmodulin; this complex then activates myosin light-chain kinase, which phosphorylates the myosin heads.
  2. Phosphorylation of the myosin heads is needed for them to be able to bind to actin and produce contractions.
  - D. Smooth muscles are classified as single-unit, if they are interconnected by gap junctions, and as multiunit if they are not so connected.
  - E. Autonomic neurons have varicosities that release neurotransmitter all along their length of contact with the smooth muscle cells, making synapses en passant.

## Review Activities

### Test Your Knowledge of Terms and Facts

1. A graded whole muscle contraction is produced in vivo primarily by variations in
  - a. the strength of the fiber's contraction.
  - b. the number of fibers that are contracting.
  - c. both of the above.
  - d. neither of the above.
2. The series-elastic component of muscle contraction is responsible for
  - a. increased muscle shortening to successive twitches.
  - b. a time delay between contraction and shortening.
  - c. the lengthening of muscle after contraction has ceased.
  - d. all of the above.
3. Which of the following muscles have motor units with the highest innervation ratio?
  - a. leg muscles
  - b. arm muscles
  - c. muscles that move the fingers
  - d. muscles of the trunk
4. The stimulation of gamma motoneurons produces
  - a. isotonic contraction of intrafusal fibers.
  - b. isometric contraction of intrafusal fibers.
  - c. either isotonic or isometric contraction of intrafusal fibers.
  - d. contraction of extrafusal fibers.
5. In a single reflex arc involved in the knee-jerk reflex, how many synapses are activated within the spinal cord?
  - a. thousands
  - b. hundreds
  - c. dozens
  - d. two
  - e. one
6. Spastic paralysis may occur when there is damage to
  - a. the lower motor neurons.
  - b. the upper motor neurons.
  - c. either the lower or the upper motor neurons.
7. When a skeletal muscle shortens during contraction, which of the following statements is *false*?
  - a. The A bands shorten.
  - b. The H bands shorten.
  - c. The I bands shorten.
  - d. The sarcomeres shorten.
8. Electrical excitation of a muscle fiber *most directly* causes
  - a. movement of tropomyosin.
  - b. attachment of the cross bridges to actin.
  - c. release of  $\text{Ca}^{2+}$  from the sarcoplasmic reticulum.
  - d. splitting of ATP.
9. The energy for muscle contraction is *most directly* obtained from
  - a. phosphocreatine.
  - b. ATP.
  - c. anaerobic respiration.
  - d. aerobic respiration.
10. Which of the following statements about cross bridges is *false*?
  - a. They are composed of myosin.
  - b. They bind to ATP after they detach from actin.
  - c. They contain an ATPase.
  - d. They split ATP before they attach to actin.
11. When a muscle is stimulated to contract,  $\text{Ca}^{2+}$  binds to
  - a. myosin.
  - b. tropomyosin.
  - c. actin.
  - d. troponin.
12. Which of the following statements about muscle fatigue is *false*?
  - a. It may result when ATP is no longer available for the cross-bridge cycle.
  - b. It may be caused by a loss of muscle cell  $\text{Ca}^{2+}$ .
  - c. It may be caused by the accumulation of extracellular  $\text{K}^+$ .
  - d. It may be a result of lactic acid production.

13. Which of the following types of muscle cells are *not* capable of spontaneous depolarization?
- single-unit smooth muscle
  - multiunit smooth muscle
  - cardiac muscle
  - skeletal muscle
- e. both *b* and *d*  
f. both *a* and *c*
14. Which of the following muscle types is striated and contains gap junctions?
- single-unit smooth muscle
  - multiunit smooth muscle
  - cardiac muscle
  - skeletal muscle
15. In an isotonic muscle contraction,
- the length of the muscle remains constant.
  - the muscle tension remains constant.
  - both muscle length and tension are changed.
  - movement of bones does not occur.

### Test Your Understanding of Concepts and Principles

- Using the concept of motor units, explain how skeletal muscles *in vivo* produce graded and sustained contractions.
- Describe how an isometric contraction can be converted into an isotonic contraction using the concepts of motor unit recruitment and the series-elastic component of muscles.
- Trace the sequence of events in which the cross bridges attach to the thin filaments when a muscle is stimulated by a nerve. Why don't the cross bridges attach to the thin filaments when a muscle is relaxed?
- Using the sliding filament theory of contraction, explain why the contraction strength of a muscle is maximal at a particular muscle length.
- Explain why muscle tone is first decreased and then increased when descending motor tracts are damaged. How is muscle tone maintained?
- Explain the role of ATP in muscle contraction and muscle relaxation.
- Why are all the muscle fibers of a given motor unit of the same type? Why are smaller motor units and slow-twitch muscle fibers used more frequently than larger motor units and fast-twitch fibers?
- What changes occur in muscle metabolism as the intensity of exercise is increased? Describe the changes that occur as a result of endurance training and explain how these changes raise the level of exercise that can be performed before the onset of muscle fatigue.
- Compare the mechanism of excitation-coupling in striated muscle with that in smooth muscle.
- Compare cardiac muscle, single-unit smooth muscle, and multiunit smooth muscle with respect to the regulation of their contraction.

### Test Your Ability to Analyze and Apply Your Knowledge

- Your friend eats huge helpings of pasta for two days prior to a marathon, claiming such “carbo loading” is of benefit in the race. Is he right? What are some other things he can do to improve his performance?
- Compare muscular dystrophy and amyotrophic lateral sclerosis (ALS) in terms of their causes and their effects on muscles.
- Why is it important to have a large amount of stored high-energy phosphates in the form of creatine phosphate for the function of muscles during exercise? What might happen to a muscle in your body if it ever ran out of ATP?
- How is electrical excitation of a skeletal muscle fiber coupled to muscle contraction? Speculate on why the exact mechanism of this coupling has been difficult to determine.
- How would a rise in the extracellular  $\text{Ca}^{2+}$  concentration affect the beating of a heart? Explain the mechanisms involved. Lowering the blood  $\text{Ca}^{2+}$  concentration can cause muscle spasms. What might be responsible for this effect?

## Related Websites

Check out [www.mhhe.com/fox](http://www.mhhe.com/fox) for links to sites containing resources related to the Muscles. These links are monitored to ensure the URLs are updated as needed. Examples of the sites you will find include:

Muscular Dystrophy Association  
Mayo Clinic Health Oasis (ALS, calcium channel blockers)