	Outline		
	• anatomical structure		
Mechanics of Skeletal Muscle	• physiological basis of muscle contraction (muscle nerve interaction, sliding filament theory of force development)		
Ozkaya and Nordin Chapter 9, pages 213-214	 effect on muscle force of stimulation frequency, muscle fibre type, muscle length, velocity of shortening/ lengthening, muscle geometry (PCSA, angle of pennation) 		
	• quick-release experiments		
	• Hill's active state model of muscle contraction		
KIN 2012007-1Stephen Robinovitch, Ph.D.1	2		
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Muscle contraction may occur with or without a change in muscle length

- muscle "contraction" refers to force development in activated muscle, rather than decrease in length
- contractions may be:
 - *isometric*: constant muscle length
 - *isokinetic*: constant muscle velocity
 - isotonic: constant muscle load
 - <u>concentric</u>: shortening
 - eccentric: lengthening
- strains are greatest during concentric contractions, where muscle may shorten by 50-70 percent

5

Eccentric and concentric contractions

- <u>Concentric</u> (energy generating, positive work) contractions tend to increase joint angular velocity, and increase the total energy of the system
- Eccentric (energy absorbing, negative work) *contractions* tend to decrease (or brake) joint angular velocity, and reduce the total energy of the system

Concentric Eccentric (braking) (accelerating) contraction contraction of m_2



of m₂

6

Z-disc structure

Calcium is needed for muscle contraction

- the sarcoplasmic reticulum (SR) is a membrane that surrounds myofibrils
- at the onset of an action potential, the SR releases calcium
- calcium then binds to troponin, triggering muscle contraction
- the SR re-sequesters calcium at the end of the action potential, thereby inducing muscle relaxation



Muscle force results from interaction between contractile proteins

- sliding filament model proposes that muscle force arises from cyclic binding between thick and thin filaments of the sarcomere
- thin filaments contain actin. troponin C, and tropomyosin
- thick filaments contain myosin
- in the absence of calcium. tropomyosin prevents myosin from attaching to actin

Muscle force results from interaction between contractile proteins (cont)

- during the action potential, calcium binds to troponin C, inducing a conformal change in tropomyosin
- simultaneously, adenosine triphosphate (ATP) is hydrolyzed by ATPase in the myosin head, providing the metabolic energy required for cross-bridge attachment
- at the end of the action potential, calcium re-uptake causes a reconfiguration of tropomyosin to a position which prevents cross-bridge attachment



9

Factors affecting muscle force development (a partial list)

- muscle geometry (e.g., physiological crosssectional area (PCSA), angle of pennation)
- number of activated motor neurons, frequency of discharge
- muscle fibre type
- muscle length
- velocity of shortening/ lengthening

10

gross muscle geometry affects muscle force

- peak muscle force increases linearly with physiologic cross-sectional area (PCSA)
- *pennate muscles*: have fibers that run oblique to long axis of the muscle
- pennate muscles have larger PCSA (and muscle force), but smaller length and shortening velocity, than parallel fiber muscles



Muscle-nerve interaction

- *motor unit*: a single motor nerve axon and all the muscle fibers it contacts
- a motor nerve enters muscle and splits into numerous axons; each axon contacts 10-2000 muscle fibers
- each muscle fiber is innervated by only one motor nerve axon, and contracts in response to an action potential in that axon



muscle	# muscle fibers	# motor units	av. fibers per motor u.
<u>platysma</u>	27,100	1,100	25
Brachioradialis	130,000	330	410
Tibialis anterior	250,000	450	600
gastrocnemius	1,120,000	580	2,000

stimulation frequency affects muscle force: twitch and tetanus

• muscle force can be modulated by varying: (1) the number of recruited motor neurons, and (2) the frequency of discharge (i.e., stimulation rate) in motor neurons

- a single action potential (S_1) produces a *twitch* contraction, a quick rise and slow fall in force
- a tetanus occurs when a new action potential (S_2) arrives before the previous twitch has dissipated, and there is force summation
- at stimulation frequencies >30/s, there are no twitch transients (fused tetanus)



active force development in the sarcomere depends on actin-myosin overlap

- (A): no overlap between actin and myosin, zero developed tension
- between (A) and (B): tension increases linearly as overlap increases
- between (B) and (C): maximum overlap & maximum tension
- left of (C): interference between actin filaments reduces ability of crossbridges to develop tension
- •left of (D): myosin filaments collide with Z-lines and fold, and force declines rapidly



muscle fiber type affects the speed and strength of muscle force

Fibre type

SO (slow,

oxidative

• fast glycolytic (FG, type IIB, white meat) fibers: twitch contraction times less than 55 ms. can generate 2-3 times more force than slow fibers, but highly fatigable, ATP generation through conversion of glucose to lactic acid (glycolysis)

• slow oxidative (SO, type I, red meat): fatigue FOG (fast, oxidative resistant, ATP generation through oxidative glycolytic) phosphorylation of blood glucose and free fatty acids

• fast oxidative glycolytic (FOG, type IIA): intermediate degree of fatigue resistance, ATP FG (fast, glycolytic) generation through oxidative phosphorylation and glycolysis



Twitch



muscle length affects force development in whole

• the tension developed in a whole muscle is the sum of active force due to muscle contraction and passive force due the passive stiffness of tendon and muscle

• the passive force is negligible for lengths less that the normal resting length (l_0)

• the active force follows the tension-length behaviour of the sarcomere, and scales with muscle activation



509

25 %

LENGTH 16

muscle velocity affects force development in whole muscle

- force (*T*) is greater during lengthening than shortening contractions
- the greater the shortening velocity (v), the smaller the force (explains why we cannot lift heavy objects quickly)
- in the shortening regime, mechanical power output is maximum when T and v are around one-third their maximum values



17

Muscle force-velocity behaviour is described by the Hill Equation

An empirical relation that describes the force-velocity behaviour of muscle during shortening is the Hill Equation. The equation can be written as : $(T + a)(v + b) = (T_o + a)b)$ where T_o is the isometric (zero velocity) tension, and v_{max} is the maximum (zero tension) velocity = $\frac{bT_o}{a}$. The instantaneous power is given by $P = T \cdot v$.



Greatest force is developed when lengthening near resting length



Hill's active state model of muscle contraction

• Hill assumed:

(1) for a given length, muscle always develops the same peak force $T_0(x_1,t)$;

(2) if the muscle is shortening, some force is dissipated in overcoming inherent viscous resistance

- *B*: muscle damping constant, which must be a nonlinear function of shortening velocity and temperature
- *K_{SE}*: stiffness of the series elastic component; represents force-deflection properties of tendon
- *K*_{*PE*}: stiffness of the parallel elastic component; represents force-deflection properties of sarcolemma, epimysium, perimysium, and endomysium



19

Quick-release experiments

- hold muscle length fixed with the catch
- stimulate muscle to produce peak (isometric) force T₀
- instantly release catch
- at the instant of release, muscle force changes to a value (T) that depends on weight in pan
- in this example, T < T₀ so the muscle shortens, rather than lengthens



21

Quick-release experiments (cont)

- there is an instant change (Δx_2) in total muscle length following release
- this occurs in the tendon, which is relatively elastic and in series with the muscle (*K*_{SE})
- this is followed by a more gradual change (Δx_i) in total muscle length
- as *T* increases, there is a decrease in *v* (slope of dashed line), reflecting that muscle cannot shorten quickly under high loads
- combinations of *T* and *v* reflect the force-velocity properties of a given muscle



Review Questions

- What is a motor unit?
- What is the role of calcium, ATP, troponin, tropomyosin, actin, and myosin in muscle contraction?
- What structure(s) contribute to the passive force-length behaviour of muscle?
- What characteristics of muscle might make it easier for us to quickly stop than to quickly start a movement?
- Why is there an optimal muscle length, above or below which there is a decrease in the force developed by activated muscle?
- During shortening, what combination of force and velocity is approximately maximizes muscle power?
- What are quick release experiments, and what data do they provide?