SKELETAL, CARDIAC, AND SMOOTH MUSCLES

SKELETAL, CARDIAC, AND SMOOTH MUSCLES

- Structural characteristics
 - Electrical and mechanical activities
 - Molecular mechanisms of contraction
 - Biophysical properties of muscle as a whole
 - Mechanisms of gradation/modulation of contraction
 - Overview of characteristic properties of skeletal, cardiac, and smooth muscles



SKELETAL MUSCLE

sarcolemma

CARDIAC MUSCLE

intercalated discs

SMOOTH MUSCLE

(vascular system, airways, gastrointestinal and urogenital systems)

ELECTRICAL CONNECTIONS "GAP JUNCTIONS"

BASIC STRUCTURAL ELEMENTS OF FUNCTIONAL SYNCYTIUM



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SMOOTH MUSCLE CELL

TRIGGERING AND MODULATION OF MECHANICAL RESPONSES

GREAT VARIETY IN ELECTRO-MECHANICAL RELATIONS



SMOOTH MUSCLE CELL

MECHANICAL RESPONSES can be triggered/modulated

- by different patterns of ELECTRICAL ACTIVITY ELECTRO-MECHANICAL COUPLING ELECTRICAL STIMULATION
- by different NEUROHUMORAL STIMULATION
 NEUROTRANSMITTERS (acetylcholine, noradrenaline, ...)
 NEURAL STIMULATION

HORMONES (e.g. progesterone, oxytocin, angiotensin II, ...)

LOCAL TISSUE FACTORS (NO, adenosine, P_{CO2}, P_{O2}, pH, ...)

HUMORAL STIMULATION

 by STRETCH of the smooth muscle cell (STRETCH-ACTIVATED CHANNELS)
 MECHANICAL STIMULATION

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CROSS STRIATED MUSCLES



CROSS STRIATED MUSCLES





CROSS-STRIATED MUSCLE

ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION



CROSS-STRIATED MUSCLE

ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION

MOLECULAR LEVEL



CROSS-STRIATED MUSCLE

ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION



CROSS-STRIATED MUSCLE





Animated model of interaction of <u>myosin head</u> with <u>actin filament (</u>,, paddling ")

Myosin – MOLECULAR MOTOR

It consumes chemical energy released from *hydrolysis of ATP* and converts it into the motion (*mechanical work*)

troponin – tropomyosin complex





CONTRACTION VARIANTS OF SMOOTH MUSCLE CELL



SMOOTH MUSCLE

1

PHASIC variant of CONTRACTION - mode of cycling



Adapted from Berne and Levi (2004)

SMOOTH MUSCLE



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ISOMETRIC AND ISOTONIC CONTRACTION



TENSION-LENGTH RELATIONSHIP

SKELETAL MUSC	TOTAL tension TOTAL tension ACTIVE tension ACTIVE tension PASSIVE tension increase in muscle length (in cm)	n ion sion <i>vivo</i>
PASSIVE tension	tension of <i>unstimulated muscle</i> at gradual s	stretching
(ELASTIC COMPONENTS)		
TOTAL tension	ISOMETRIC CONTRACTIONS of <i>stime</i>	<u>ilated</u>
<i>muscle</i> at gradually increased <i>initial (resting) length</i>		
ACTIVE tension	<i>difference</i> between TOTAL and PASSIVE tension curves at any length (<i>tension actually generated by contractile elements</i>)	

ACTIVE TENSION of **cross striated muscles** as a **function** of **INITIAL LENGTH of SARCOMERE**



SMOOTH MUSCLE

CHARACTERISTIC FEATURES

• GREAT EXTENSIBILITY

(e.g. myocytes of <u>urinary bladder</u> can lengthen up to 200%, myocytes of <u>uterus</u> even up to 1000% at the end of pregnancy in relation to their original state)

PLASTICITY

No direct relation between the **LENGTH** and **TENSION** in smooth muscle cells. Stretch-induced *increased tension* almost *immediately spontaneously decreases*.

Analogous relation is valid between **VOLUME** and **PRESSURE** in **hollow organs** (*stomach, intestines, urinary bladder, ...*).

PLASTICITY OF SMOOTH MUSCLE



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SKELETAL MUSCLE

MAIN FACTORS IN GRADATION OF CONTRACTION

- ↑ *frequency* of discharges in *motor neuron* ⇒ FREQUENCY
 SUMMATION of contractions in skeletal muscle fibre
 (TETANIC CONTRACTION)
- *number* of activated MOTOR UNITS by increasing voluntary effort ⇒ SPATIAL SUMMATION (multiple fibre summation) RECRUITMENT OF MOTOR UNITS



SKELETAL MUSCLE

GRADATION of CONTRACTION by ↑ FREQUENCY of STIMULATIONSINGLE MUSCLE FIBRE

RANGE OF SUMMATION

physiological behaviour of skeletal myocyte



1 Hz = 1 impulse/sec

CARDIAC MUSCLE

MAIN FACTORS IN GRADATION OF CONTRACTION

- ↑ DIASTOLIC FILLING of ventricles in vivo (,,preload")

 ↑ contraction of ventricles proportionate to the stretching
 of cardiomyocytes at the end of diastole

 FRANK-STARLING'S LAW
- **FREQUENCY of electrical activity** of cardiac cells *via* modulation of pacemaker activity of SA node by sympathetic nerves positive FREQUENCY EFFECT
- **LIGAND-RECEPTOR ACTIVATION CASCADES** leading to $\uparrow [Ca^{2+}]_i$ (noradrenalin, adrenalin, thyroxine, ...)



SMOOTH MUSCLE

MAIN FACTORS IN GRADATION OF CONTRACTION / TONUS

■ **DEPOLARIZATION of the smooth muscle membrane** with or without triggering of action potentials via opening of the *voltage dependent calcium channels* $\Rightarrow \uparrow [Ca^{2+}]_i$

FACTORS <u>in</u>dependent on membrane depolarization

- *Ligand-receptor activation cascades* leading to $\uparrow [Ca^{2+}]_i$ (e.g. *via activation* of PLC $\Rightarrow \uparrow IP_3$ releasing Ca²⁺ from SR)
- Stretching of the smooth muscle cell ⇒ opening of the stretch-activated channels ⇒ ↑ [Ca²⁺]_i

 \uparrow Ca²⁺-calmodulin complex

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SKELETAL MUSCLE

MAIN CHARACTERISTIC FEATURES

- *Multinucleated* long cylindrical cells (max. length up to 20 cm)
- *Rich* sarcoplasmic reticulum
- *Regular arrangement* of thick and thin filaments in sarcomeres (*cross striation*)
- Activity strongly dependent on *motor nerve supply* (excitation transmitted via *motor end-plate*)
- Without intercellular connections (no gap junctions between muscle cells)
- Motor neurons branch to innervate more muscle cells (*motor unit* defined as one motor neuron with 5-1000 myocytes)
- Summation of contractions (tetanus) is a physiological property of muscle fibre
- Activity under *voluntary control*



motor unit

MAIN TYPES OF SKELETAL MUSCLE FIBRES

TYPE ISLOW - RED

e.g. muscles of the back, soleus m.

- *Slow* (posture-maintaining) *contractions*
- Motor units contain slowly conducting motor neurons

High OXIDATIVE CAPACITY and <u>high</u> resistance to fatigue

TYPE II

FAST (RED /WHITE)

e.g. extraocular muscles, muscles of the hand

- *Short* twitches for fine skilled movements
- Motor units with rapidly conducting motor neurons

TYPE IIa (FAST-RED) and TYPE IIb (FAST-WHITE)

Proportion of OXIDATIVE and GLYCOLYTIC metabolism determines the resistance to fatigue

Sport activities cause gradual transformation from IIb into IIa

CARDIAC MUSCLE

MAIN CHARACTERISTIC FEATURES

- Branched and interconnected cells with one nucleus in the centre (length ~100 μm)
- Well (moderately) developed sarcoplasmic reticulum
- *Regular arrangement* of thick and thin filaments in sarcomeres (*cross striation*)
- Excitations (contractions) are independent on nerve supply (*specialized pacemaker cells*)
- Functional syncytium (electrical connections *gap junctions*)
- *Receptors* for *neurotransmitters* (released from neuron endings) and *hormones* (brought by circulation); activity is *modulated* by *local mediators*
- *Long refractory period prevents* cells from tetanic contraction (which would be life threatening)
- Activity is **not** under *voluntary* control

SMOOTH MUSCLE

MAIN CHARACTERISTIC FEATURES

- Thin *spindle-shaped* cells of various length (20-200 μm) with *one nucleus* in the centre
- *Irregular arrangement* of thick and thin filaments; no cross striation
- Poorly developped sarcoplasmic reticulum, TT system is missing
- Contractions of *visceral muscles* can be triggered independently on nerve supply (*slow irregular unstable pacemaker activity*); functional syncytium (*gap junctions*)
- Slow *phasic*, often *tonic*, even *tetanic* contractions
- Numerous *receptors* for *neurotransmitters* (released from neuron endings) and *hormones* (brought by circulation). Activity is greatly modulated by *local mediators* (local tissue factors)
- Activity can be triggered by stretch (*stretch activated channels*)
- Great extensibility and plasticity
- Activity without voluntary control

TYPES OF SMOOTH MUSCLE

VISCERAL "SINGLE UNIT"

e.g. stomach, intestine, uterus, ureter, ...

- Functional syncytium (gap junctions)
- Excitation and contraction can be evoked *in the absence of nerve supply* (*slow irregular pacemakers in multiple foci* shifting from place to place, *gap junctions*)
- Contraction evoked by **stretching** (*stretch-activated channels*)

MULTIUNITstimulated by neurons

e.g. arterioles, m. ciliaris, muscle of iris, ...

- Myocytes need the stimulation by autonomic "motor" neurons releasing *acetylcholine / norepinephrine, ...* (AUTONOMIC "MOTOR UNITS")
- Cells are **not** interconnected by **gap junctions**, **APs** are **not** triggered
- Synapses "*en passant*" in the course of the neuron endings
- Contractions are *finely graded* and *localized*

