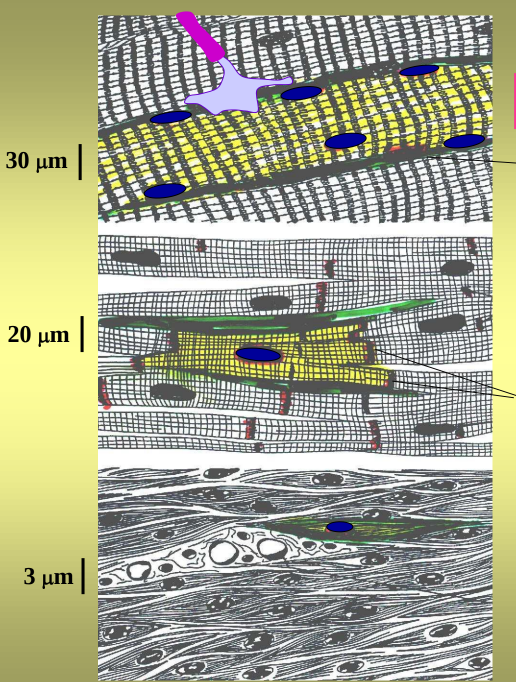
SKELETAL, CARDIAC, AND SMOOTH MUSCLES

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

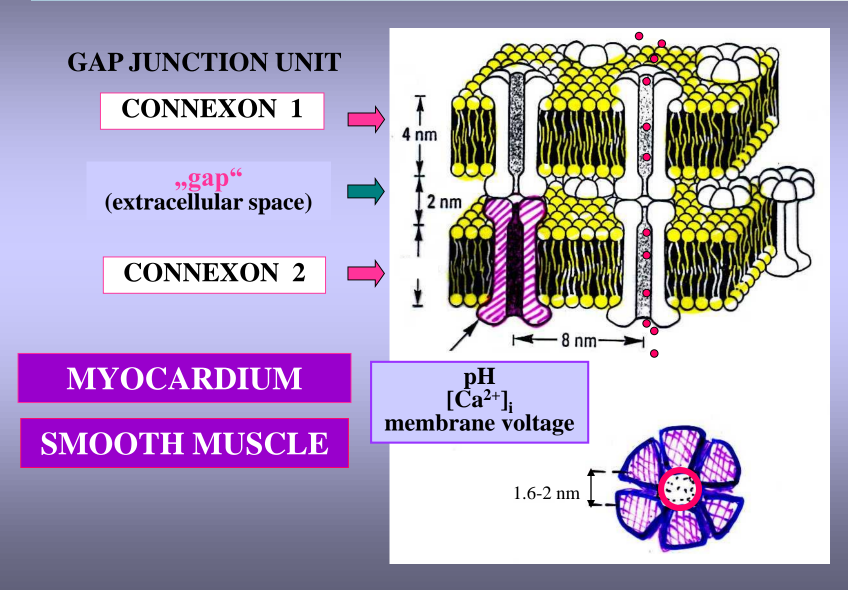
intercalated discs

SMOOTH MUSCLE

(vascular system, airways, gastrointestinal and urogenital systems)

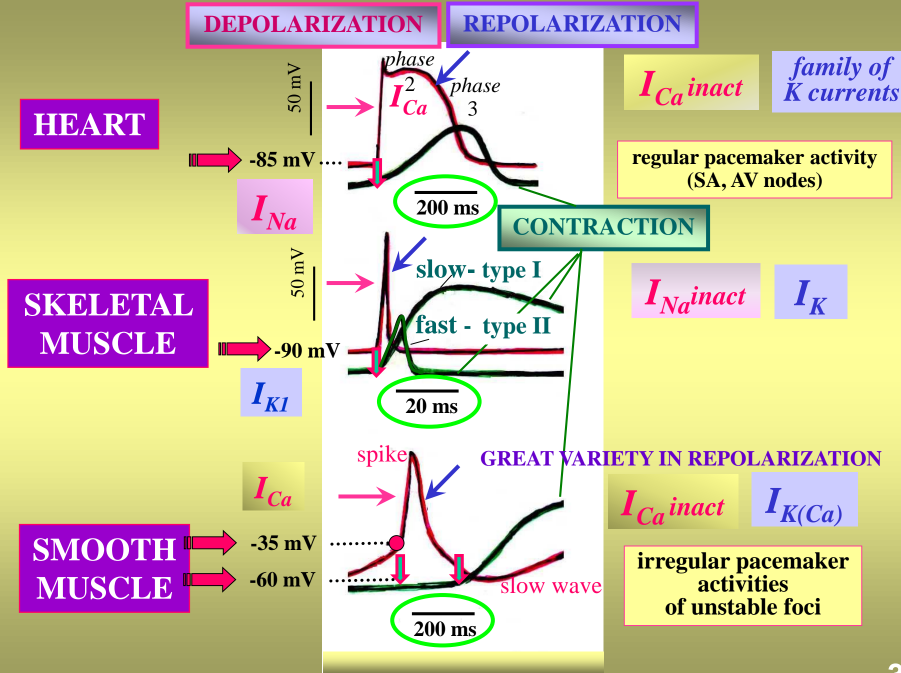
ELECTRICAL CONNECTIONS "GAP JUNCTIONS"

BASIC STRUCTURAL ELEMENTS OF FUNCTIONAL SYNCYTIUM



SKELETAL, CARDIAC, AND SMOOTH MUSCLES

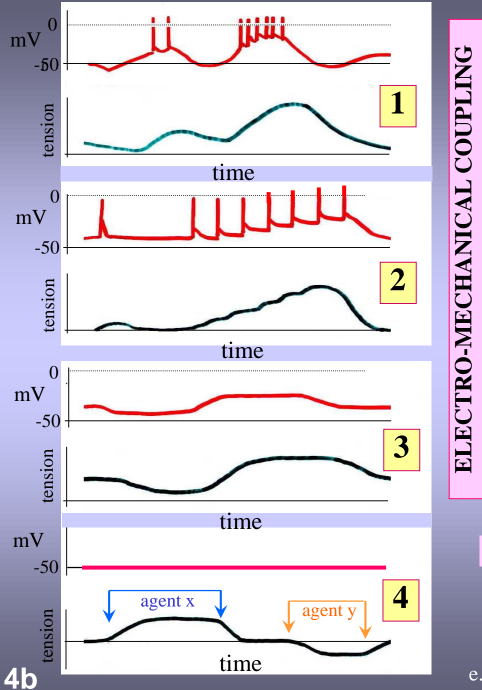
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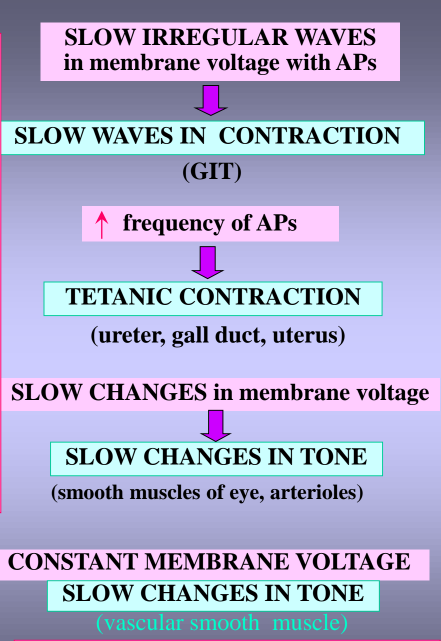


SMOOTH MUSCLE CELL

TRIGGERING AND MODULATION OF MECHANICAL RESPONSES

GREAT VARIETY IN ELECTRO-MECHANICAL RELATIONS





NEUROHUMORAL STIMULATION

e.g. via LIGAND-RECEPTOR activation pathways

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, ...)

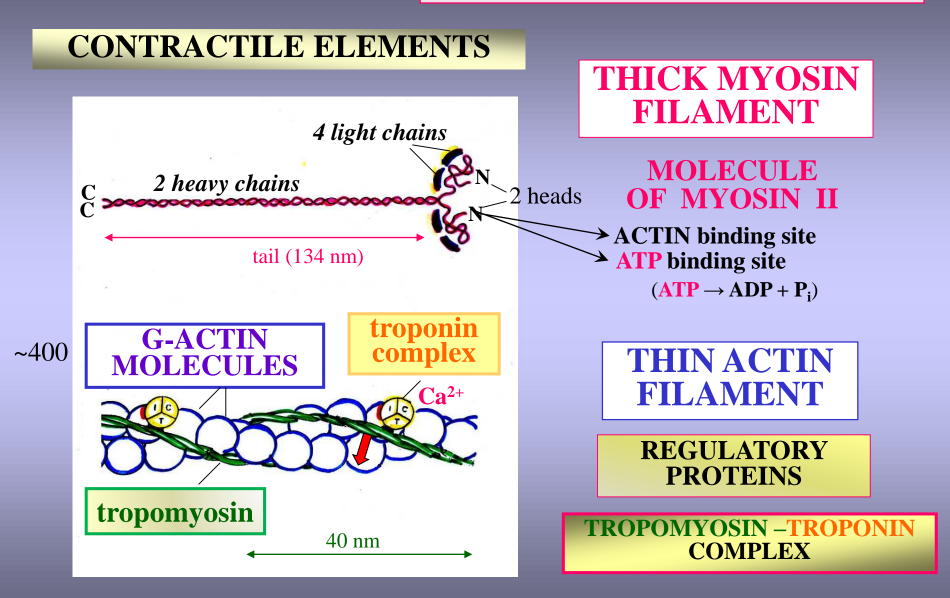
LOCAL TISSUE FACTORS (NO, adenosine, P_{CO2}, P_{O2}, pH, ...)

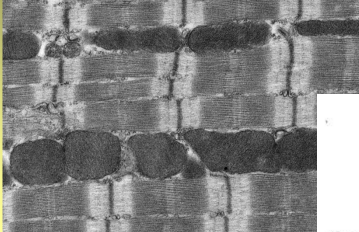
HUMORAL STIMULATION

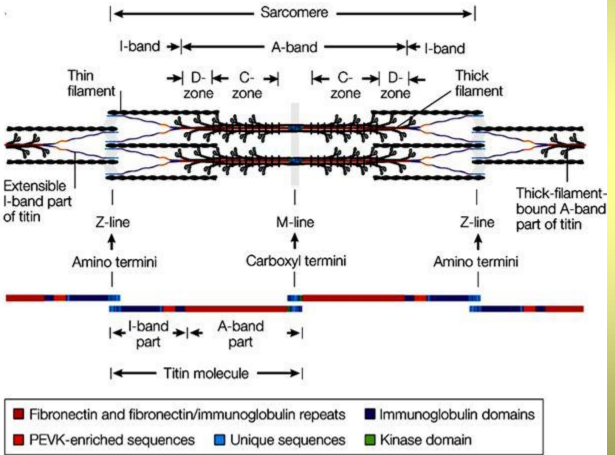
 by STRETCH of the smooth muscle cell (STRETCH-ACTIVATED CHANNELS)
 MECHANICAL STIMULATION

SKELETAL, CARDIAC, AND SMOOTH MUSCLES

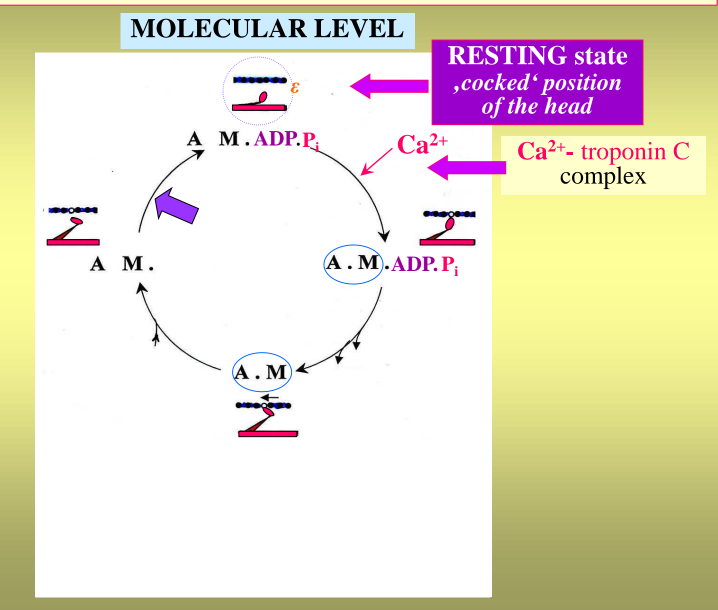
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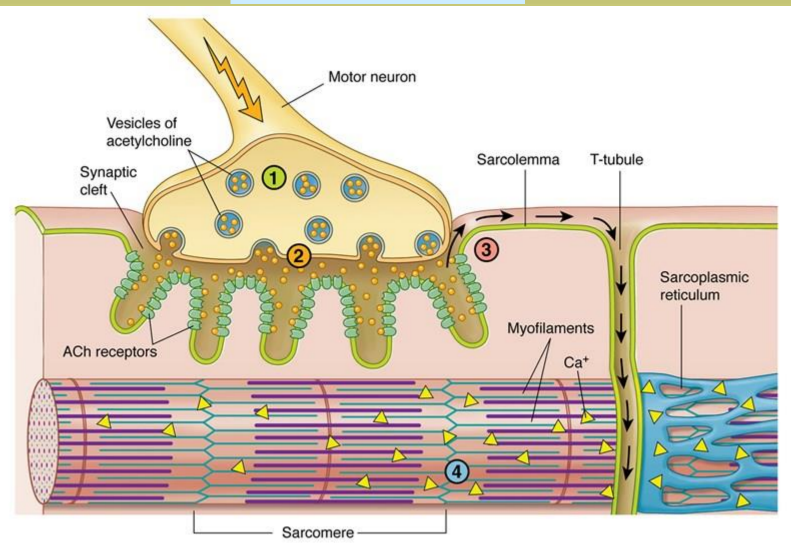


ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION

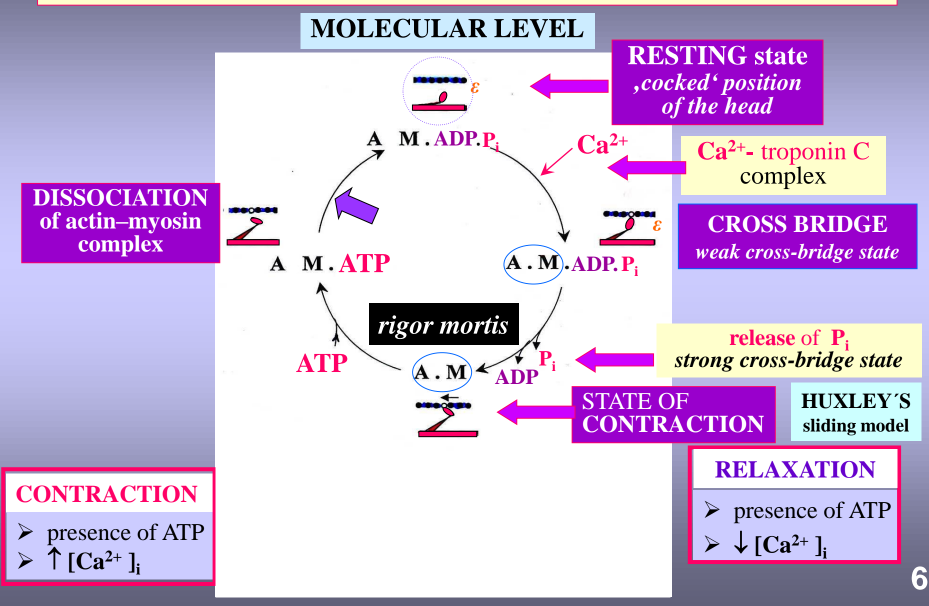


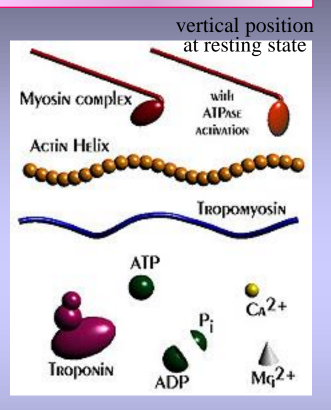
ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION

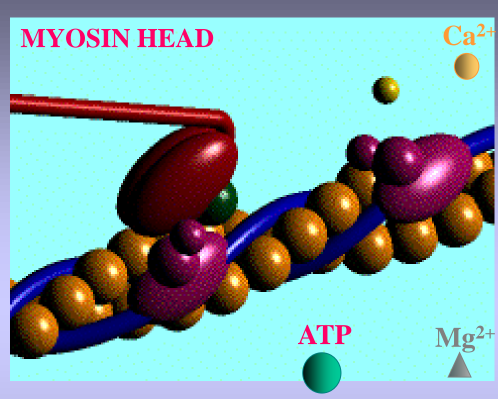
MOLECULAR LEVEL



ONE ELEMENTARY CYCLE OF CONTRACTION AND RELAXATION







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

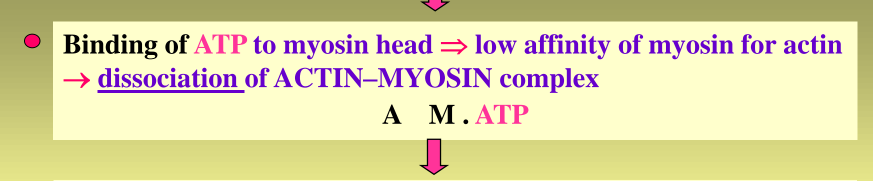
Myosin – MOLECULAR MOTOR

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

troponin – tropomyosin complex

MOLECULAR MECHANISM OF CONTRACTION

- Binding of Ca²⁺ to TROPONIN C ⇒ shift of troponin-tropomyosin complex → <u>actin binding sites</u> for myosin heads <u>are uncovered</u>
- Formation of CROSS BRIDGES between actin and myosin (weak cross-bridge state)
 A.M^e. ADP. P_i
- Release of P_i (strong cross-bridge state) ⇒ conformational changes in myosin head-neck junction → tilt of the myosin head (power stroke)
 → sliding of thin on thick filaments ⇒ SHORTENING OF SARCOMERE
- ADP is released → actomyosin complex is left in a rigid 'attached' state A.M



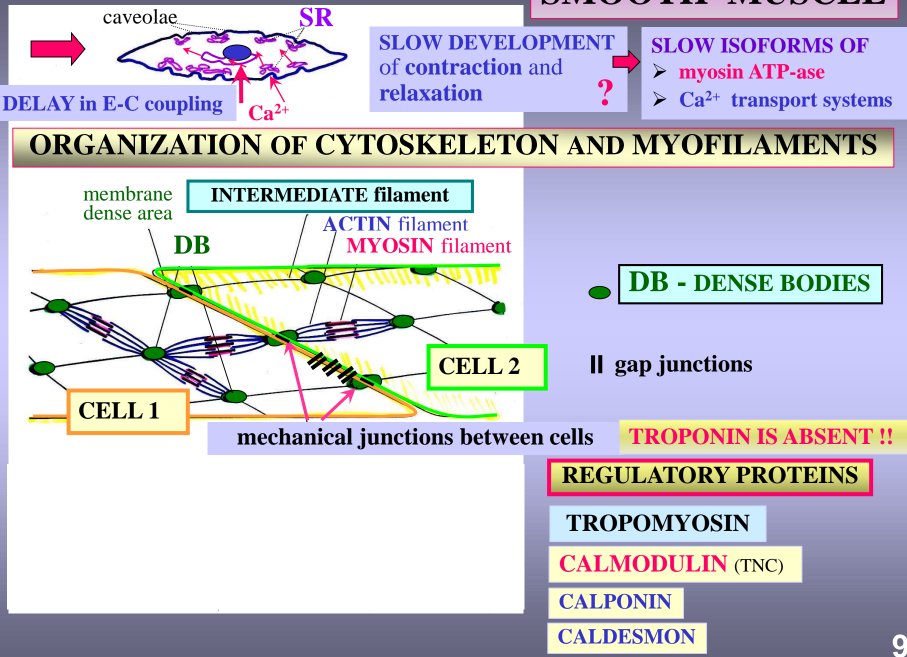
ATP-ase activity of myosin head ⇒ partial hydrolysis of ATP, the gained energy is used for re-cocking of the myosin head (analogy of the stretched spiral spring).

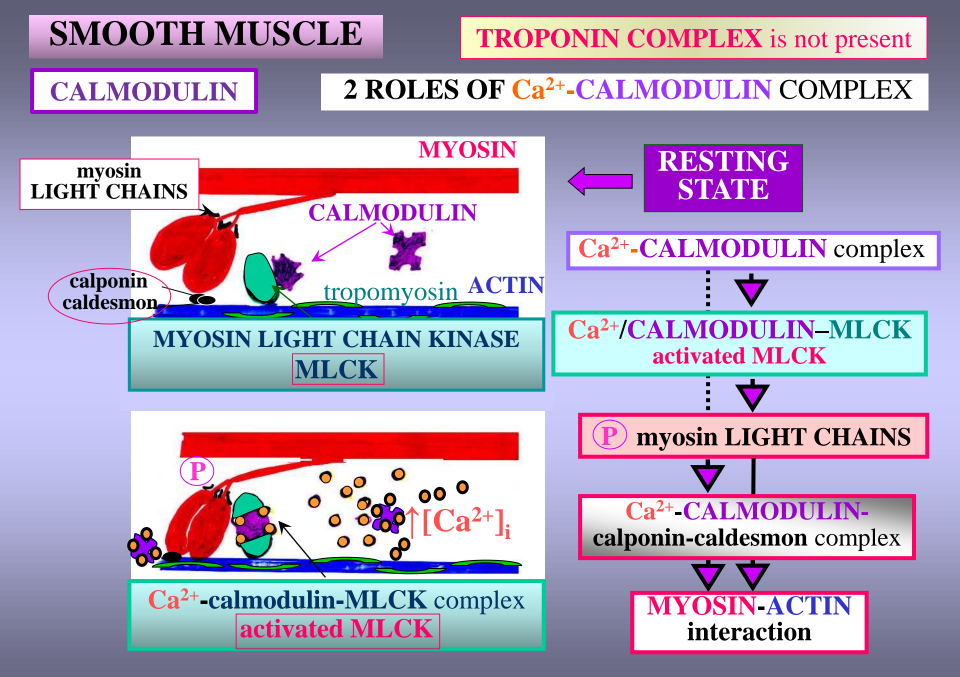
A M^{ε} . ADP. P_{i}



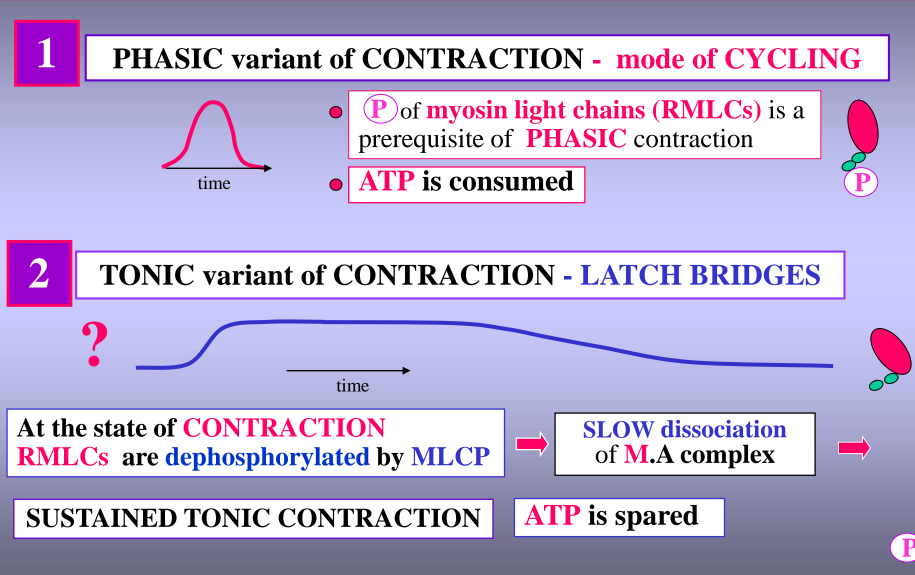
• **RELAXATION of the muscle cell** results from the **presence of ATP** and \downarrow [Ca²⁺]_i (Ca²⁺ is pumped back into SR and pumped out of the cell)







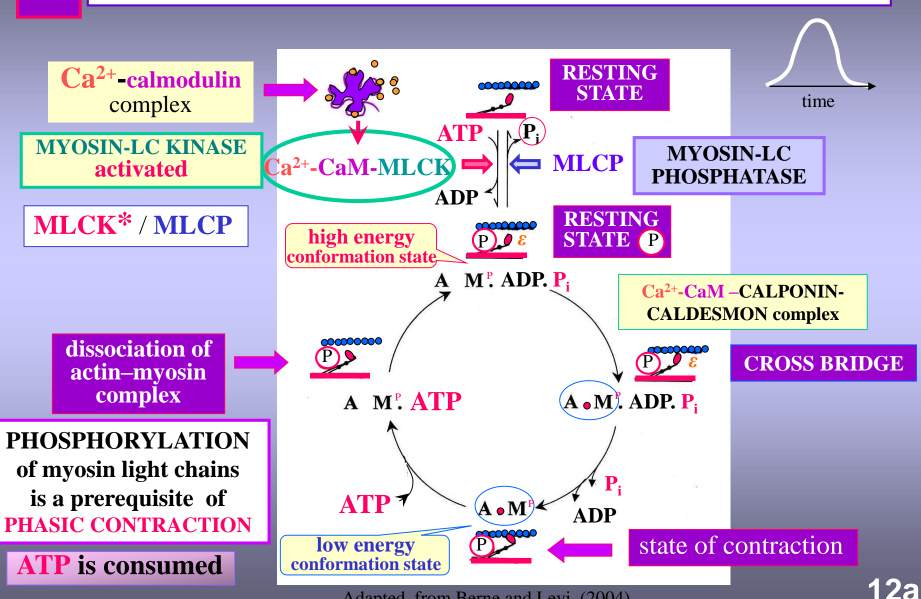
CONTRACTION VARIANTS OF SMOOTH MUSCLE CELL



SMOOTH MUSCLE

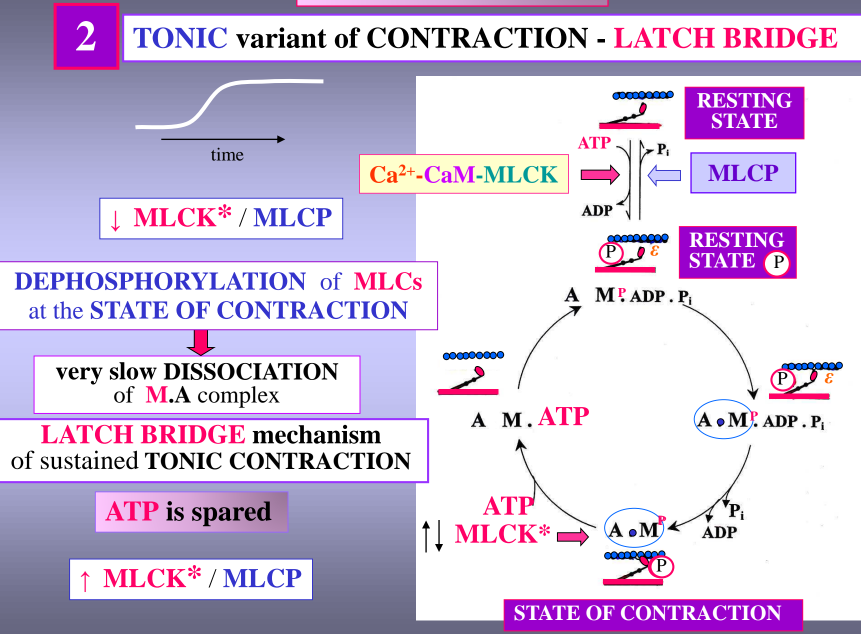
1

PHASIC variant of CONTRACTION - mode of cycling

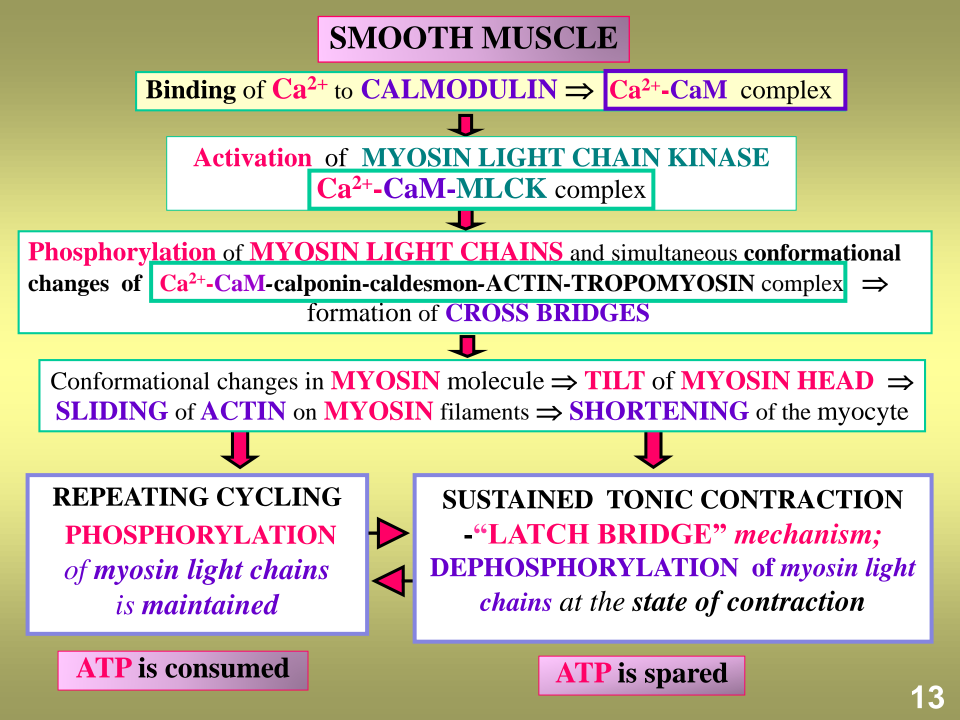


Adapted from Berne and Levi (2004)

SMOOTH MUSCLE



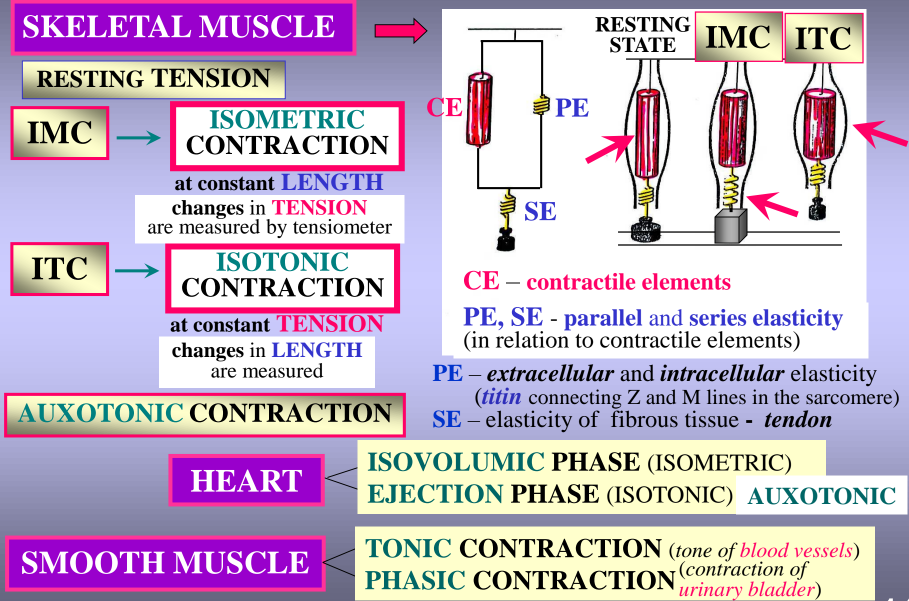
Adapted from Berne and Levi (2004)



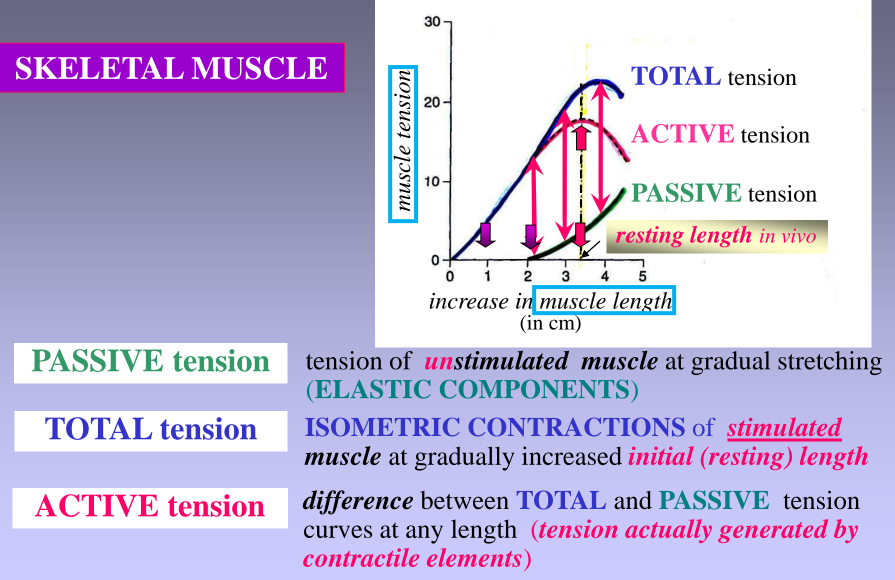
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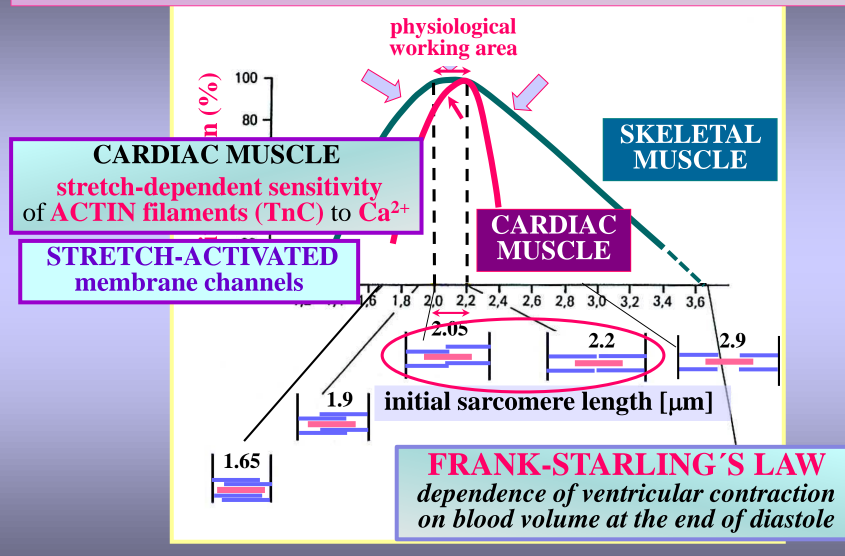
ISOMETRIC AND ISOTONIC CONTRACTION



TENSION-LENGTH RELATIONSHIP



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



SMOOTH MUSCLE

CHARACTERISTIC FEATURES

• GREAT EXTENSIBILITY

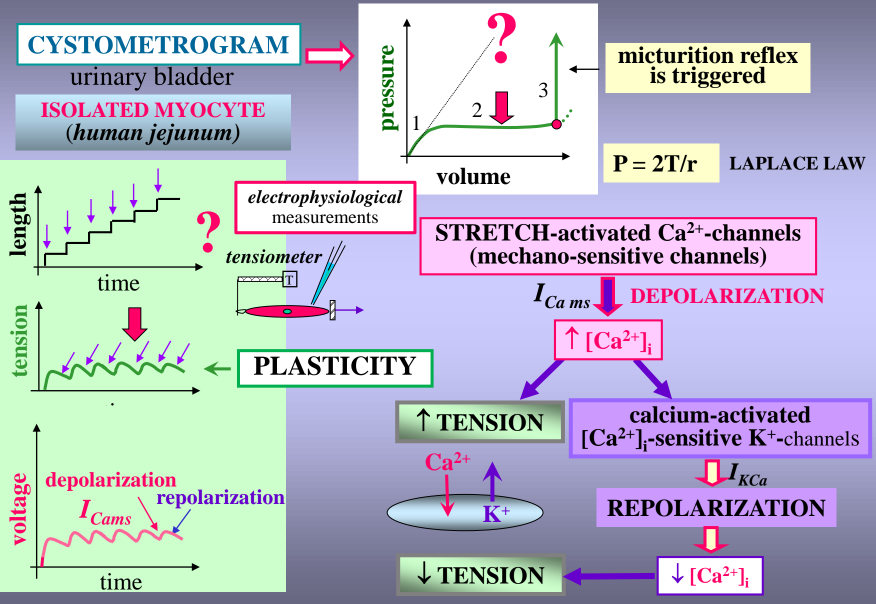
(e.g. **myocytes** of **<u>urinary bladder</u>** can lengthen up to **200%**, **myocytes** of **<u>uterus</u>** even up to **1000%** at the end of pregnancy <u>in relation to their original state</u>)

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



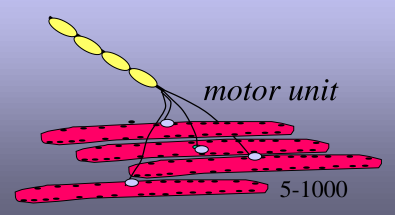
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SKELETAL MUSCLE

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

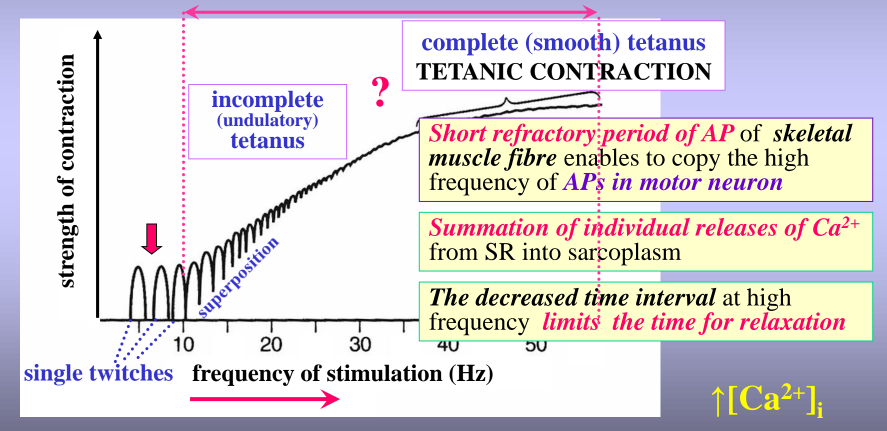


SKELETAL MUSCLE

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

SLOW - RED TYPE I

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*

