ZOO 332H1S Lecture 5,6 Jan. 2003 (AJE)

Channels, resting and action potentials:

<u>Experimental</u>
<u>evidence</u>



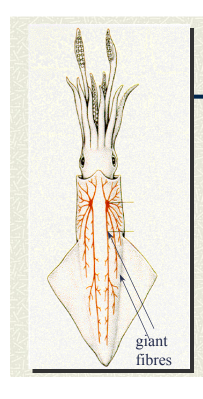


A. F. Huxley, I

Hodgkin (1949) and Huxley (1974)

Major advances occur because:

- **♯** Discover a good preparation
- **♯** Find or develop appropriate technology
- **■** Ask the right questions
- **■** Work hard and think clearly



Loligo, the squid

"It is arguable that the introduction of the squid giant nerve fibre by J.Z. Young in 1936 did more for axonology than any other single advance during the last forty years. Indeed a distinguished neurophysiologist remarked recently at a congress dinner (not, I thought, with the utmost tact), 'It's the squid that really ought to be given the Nobel Prize.'"

A.L. Hodgkin, 1973

3

Giant Nerve Cells of the Squid (see next slide for description) Giant axon Brain-Presynaptic (2nd level) Stellate nerve 1st-level Smaller axons neuron Stellate -2nd-level ganglion neuron 3rd-level neuron Postsynaptic Cross (3rd level) section Stellate nerve with 1 mm 1 mm giant axon Squid giant axon = 800 µm diameter Mammalian axon = 2 μm diameter

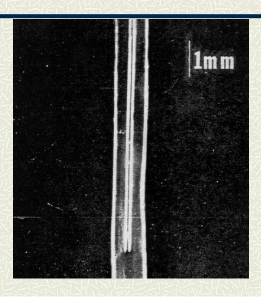
Cont...Giant Nerve Cells of the Squid

After Purves et al (2001):

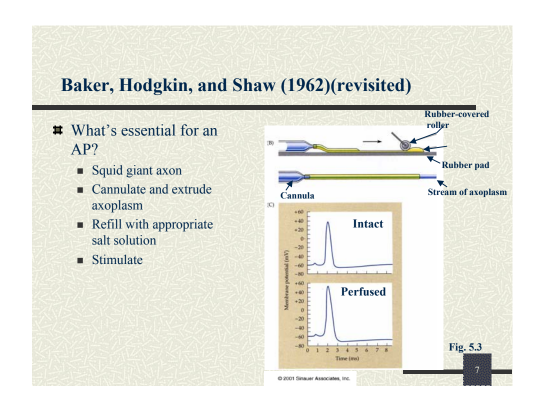
Diagram of squid, showing location of its giant nerve cells. Different colours indicate the neuronal components of the escape circuitry. The first- and second-level neurons originate in the brain, while the third-level neurons are in the stellate ganglion and innervate muscle cells of the mantle. (B) Giant synapses within the stellate ganglion. The second-level neuron forms a series of fingerlike processes, each of which makes an extraordinarily large synapse with a single third-level neuron. (C) Structure of a giant axon of a third-level neuron lying within its nerve. The difference in the diameters of a squid giant axon and a mammalian axon are shown below. However, note that some mammalian motor neurons are as large as $20~\mu m$ in diameter.

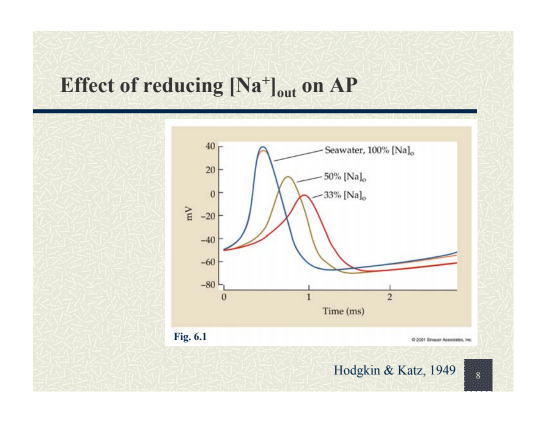
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Using "microelectrodes"



Hodgkin, Huxley, and others pioneered recording intra-cellularly. They used glass capillaries inserted longitudinally into the squid giant axon.





Effects of increasing conductance's $(g_{Na+}$ and $g_{K+})$

- (A) Positive feedback of Na⁺ entry on depolarization
- (B) K⁺ efflux leads to repolarization

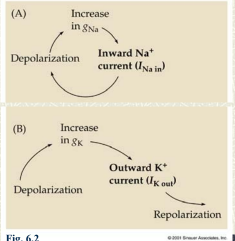
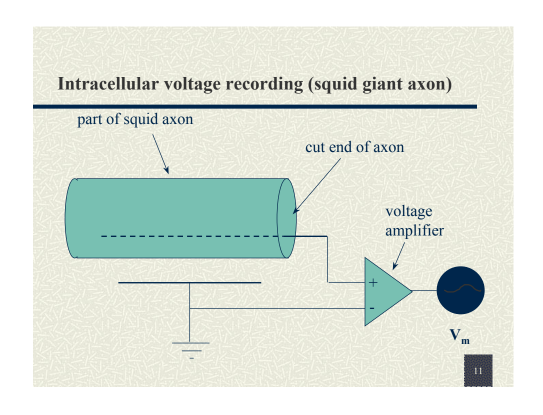
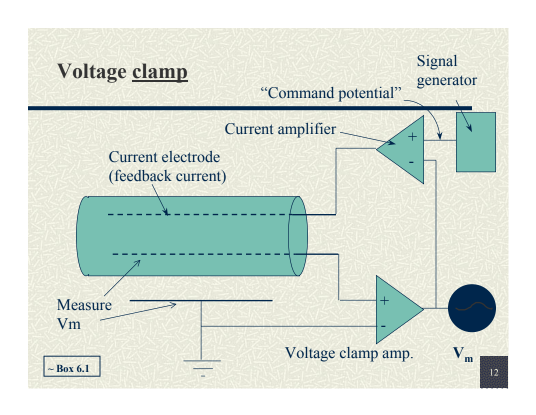


Fig. 6.2

To figure out ionic currents and changes in conductance

- \blacksquare Problem: during AP, V_m , I_m , and G_m are all changing
- Solution: if we hold one variable steady, measure a second, we can calculate the third
- ♯ The "voltage clamp" was developed to do this by Cole, Hodgkin, Huxley, Katz and others around 1949





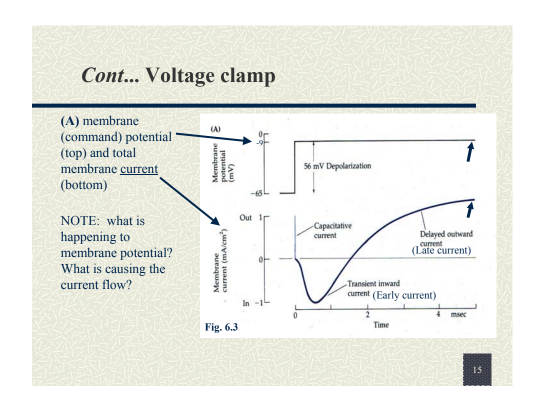
How voltage clamp works...

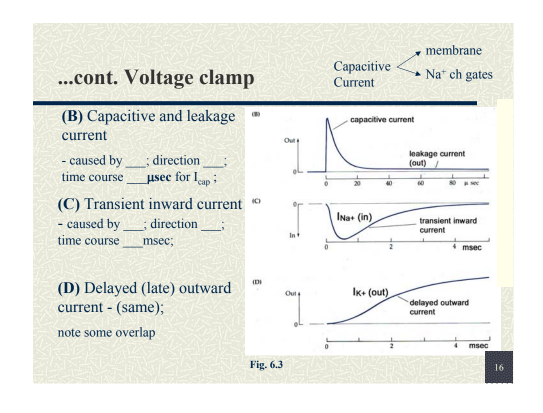
- **■** Voltage electrode records V_m
- **♯** Signal generator outputs command potential, *i.e.*, desired voltage
- Current amplifier sends out current if command potential different from V_m
- \sharp This current causes V_m to move to the command potential

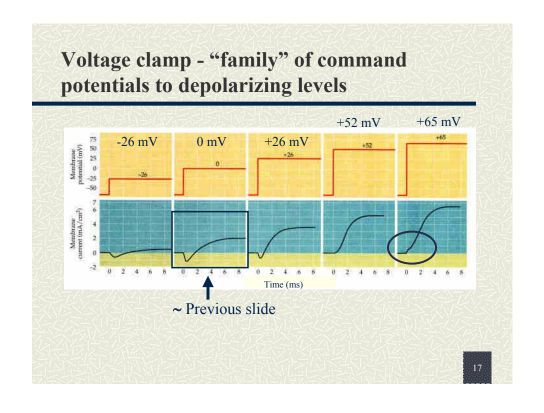
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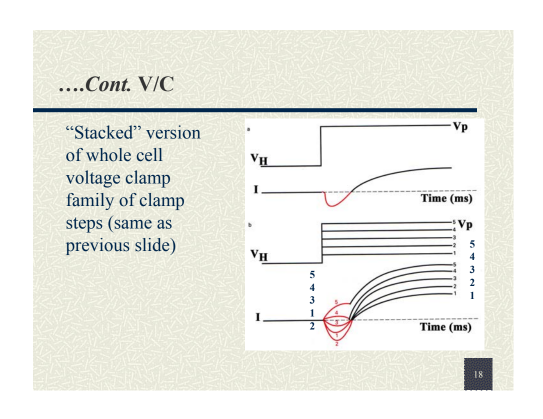
... Voltage clamp continued

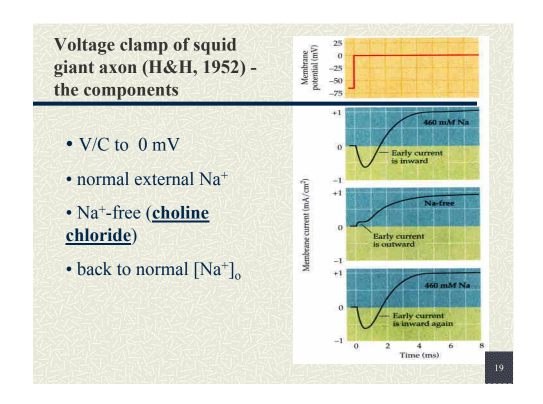
- \blacksquare The size of the current needed to "clamp" V_m , *i.e.*, to counteract ΔV_m , depends on membrane conductance (Ohm's Law)
- Therefore the size of the current gives a measure of the *actual current flowing* at this voltage, and of the *underlying conductance changes*, and their *time course*

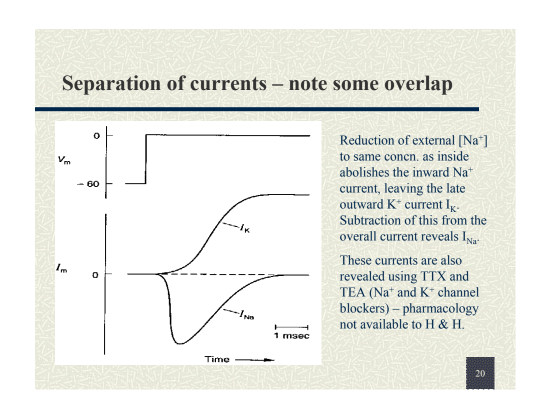












Discovery of neuroactive drugs, examples...

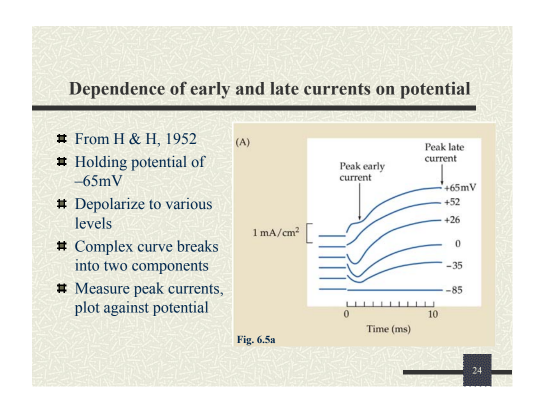
- # *Tetrodotoxin* (TTX) from puffer fish selectively blocks voltage-gated Na⁺ channels (other sources available as well)
- **♯** *Tetraethylammonium* (TEA) ions block voltagegated K⁺ channels

21

cont...Discovery of neuroactive drugs has led to powerful tools

- # powerful tools with which to investigate channels
 - **■** Kinetics
 - **■** Operation
 - Pharmacology
 - Etc.

Pharmacological separation of membrane currents Out 10 ■ Voltage clamp to 0 mV **(A)** is normal bath **=** 300 nM TTX (B) € Out 10 TTX **■** Low mM TEA **■** Time-course of individual conductance changes during an A.P. Out 10 TEA can be calculated from v/c records Time (ms) Fig. 6.4



cont...Dependence of early and late currents on potential

- Plot peak current vs. potential to which membrane is stepped
- ➡ Peak late current, <-65mV is linear (passive resistor); depolarization involves vactivated K⁺ conductance, additional current thru membrane
- Early current complex: with depolarization dealing with decreasing driving force on Na⁺, increasing conductance of vactivated Na⁺ channels, (inactivation)
- **■** Negative slope conductance

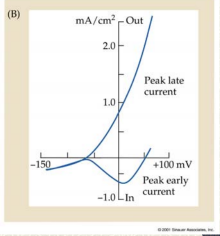


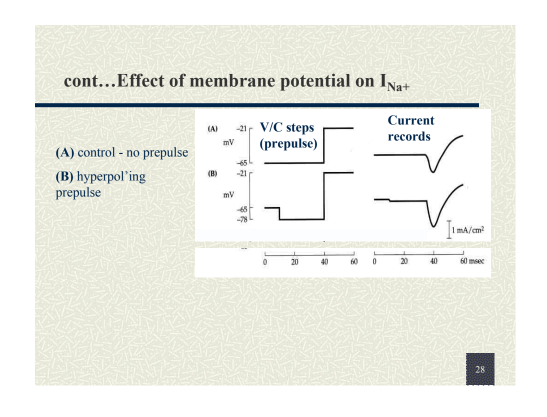
Fig. 6.5b

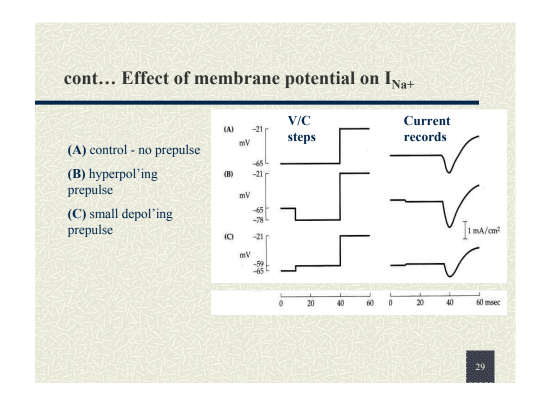
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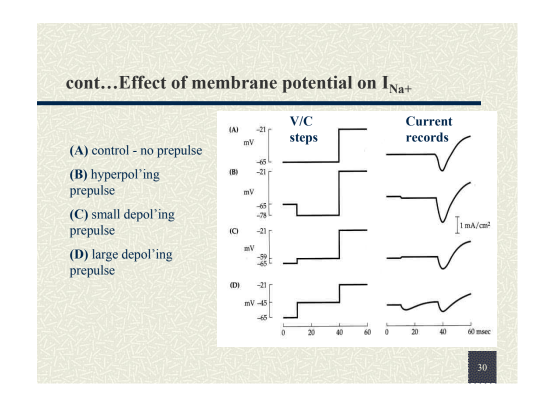
Complex interaction between channels and membrane potential

- **♯** Operate independently
- **■** Dependence on voltage
- ? Does the starting point of V_m influence subsequent inward current recorded?

Feature of v-activated Na⁺ channels, inactivation, first elucidated by H & H. (A) control - no prepulse (A) control - no prepulse Fig. 6.6a







Peak Na+ current vs. prepulse potential amount of activation in the "system"

- I_{Na no step} constant
- If current increases after a step then $I_{\text{Na}^+\,\text{step}}/~I_{\text{Na}^+\,\,\text{no}}$ $_{\text{step}}$ is > 1.0
- If current **decreases** after a step then I_{Na^+step}/I_{Na^+no} step is < 1.0
- intersection of RP and no change (1.0) shows amount of activation normally in system at
- h represents amount of activation

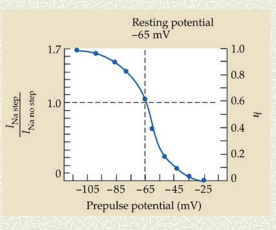


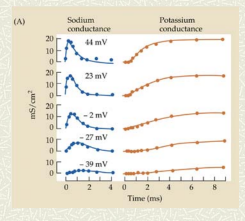
Fig. 6.6e

A further note on gNa+ and gK+

- **■** Examples of data for Na⁺ and K⁺ conductances
- **♯** Voltage steps to indicated potentials
- \blacksquare Peak g_{Na^+} and g_{K^+} both increase with depol'n

$$g_{Na^{+}} = \frac{I_{Na^{+}}}{(V_{m} - E_{Na})} \;\; ; \label{eq:gNa+}$$

And similarly for g_{K+}



cont...A further note on gNa+ and gK+

- both increase progressively as neuron is depol'd
- □ note the similar voltage dependence for each "g"
- ♯ both sigmoidal functions of membrane potential
- **#** quite small at -ve potentials
- **♯** maximal at +ve potentials
- exquisitely dependent on Vm at intermediate potentials
- deduction about mechanism of g-change senses V across membrane

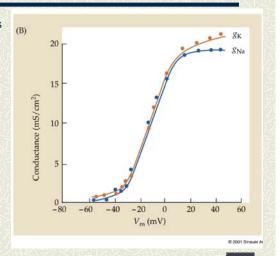


Fig. 6.7

33

Threshold and refractory period

Threshold

- Generally about 15mV depolarization required to reach (several factors influence exact value)
- Point at which gNa⁺ > outward gK⁺, initiates +ve feedback cycle of Na⁺ entry

♯ Refractory Period (absolute and relative)

- Na⁺ inactivation
- high g_{K+}

Hodgkin, Huxley, and Katz (1952)

Hodgkin & Huxley (and Katz) reconstructed the conductance changes underlying the AP from voltage clamp data.

- Apply change in potential (ΔV) across the membrane (V/C);
- Measure current flow (I);
- Calculate conductance (G) (remember: V=IR (R=1/G))

Bottom lines: point by point reconstruction; data generated experimentally essentially same as during an AP. Mathematical model accurate, AP is a predictable phenomenon.

Also, What does this tell you about the voltage clamp technique?

25

H & H finally compared their calculated APs with the observed APs

Reconstruction of the AP (mathematically)

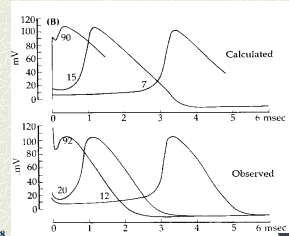
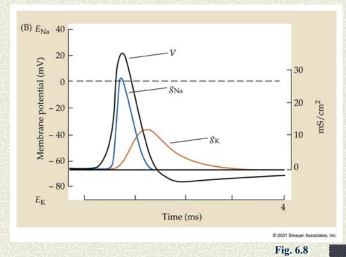


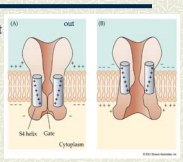
Fig. 6.8



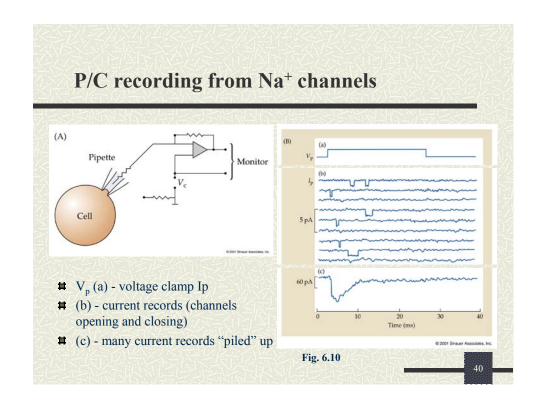


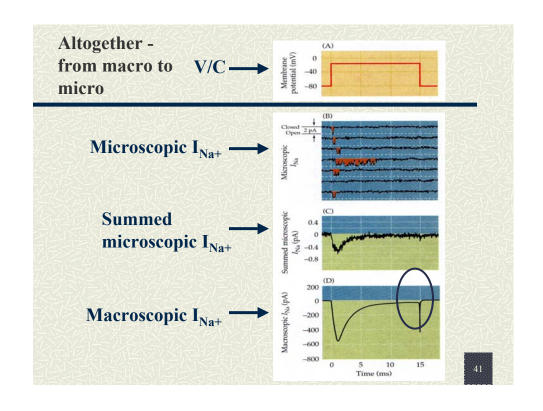
Hodgkin and Huxley's <u>prediction</u> re. Na⁺ channel "gating currents" (1952)

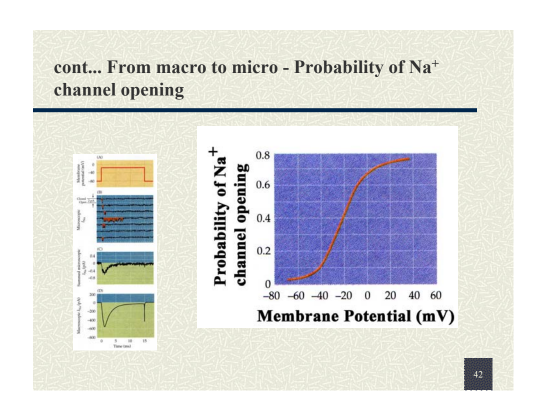
- suggested Na⁺ channel **activation** >> **movement** of **charged particles** ("structure") within the membrane
- appear as very small current in response to voltage step (**asymmetrical**), *i.e.*, doesn't occur when hyperpolarize
- block Na⁺ channels with TTX and cancel symmetrical capacitive current due to voltage step; remainder = gating current (Armstrong and Bezanilla, 1974)



cont. Na+ channels - gating currents a Depolarize (A) • (C) method of separating gating current from capacitative current 1 msec · for depol'g and hyperpol'g V/C steps, I_{cap} are equal and opposite b Hyperpolarize **(B)** electronically subtract out I_{cap} and I_{gating} remains \bullet (A) I_{gating} and I_{Na} (when reduced [Na] $_{o})$ • (B) I_{gating} alone (TTX)







More on Na+ currents - what's in the "system"?

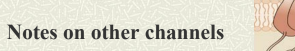
Remember...

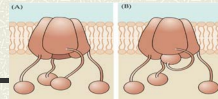
- probabilistic events Na⁺ channel opening and closing
- dependence on voltage
- Na⁺ channel activation and inactivation appear to be "separate" events

12

What experimental evidence suggested:

- inactivation was a distinct phenomenon, separable from the activation process (by H & H, 1952!)
- **pronase** experiments supported this (pronase what? where? Not effective externally)(1973, 1977)
- further support veratridine (lily) and batrachotoxin (BTX)(frog skin), both eliminate inactivation; but also alter V-dependence of activation (Na⁺ channels open at rest)(1980)(i.e., not perfectly specific)
- antibody directed at cytoplasmic domain of Na⁺ channel greatly prolonged single channel Na⁺ currents (1989)
- site-directed mutagenesis alter or delete specific a.a. in cytoplasmic domain (3 specific aa's in loop that acts to "plug" the channel), inactivation not occur (1997)





- **■** Delayed rectifier (*a.k.a.* voltage gated K⁺ channel) does not inactivate
- # Many species of K⁺ channels, different properties; for eg.
 - A-channels: inactivate with maintained depolarization; ball and chain model similar to Na⁺ channels (NMWF 107, 109); inactivate rapidly and usually inactivated at rest (require hyperpolarization to activate)
 - M-channels: <u>m</u>uscarinic AChR; 2nd messenger involved; activate gK⁺ (note mistake in text pg 109)
 - S-channels: open at rest; 5-HT via 2nd messenger, closes K⁺ channel
 - Ca²⁺-activated K⁺ channels

45

Role of calcium in excitation

- **♯** Voltage-gated Ca²⁺ channels
- **■** Distribution
- **Importance**
 - excitability (threshold complex interaction with local charges)
 - release of neurotransmitter
 - contraction of muscle fibres
 - etc.

