

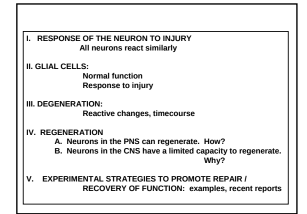
Symptoms of spinal cord injury:

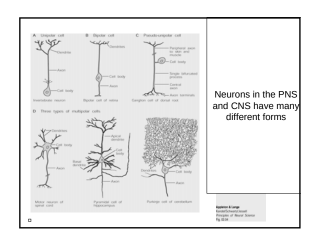
involuntary muscle spasms
/oss of voluntary movement
sensation, balance
control of breathing
autonomic functions (blood pressure)
bladder, sexual, bowel control

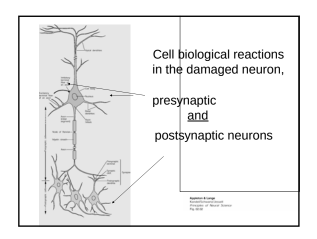
All due to destruction of long ascending or descending spinal pathways

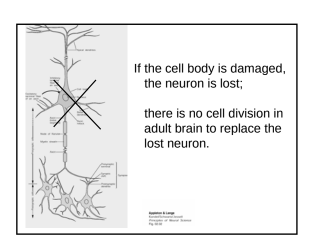
TO REPAIR THESE PATHWAYS,
AXONS must REGROW

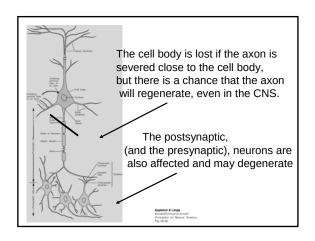
SYNAPTIC CIRCUITS must be RE-ESTABLISHED









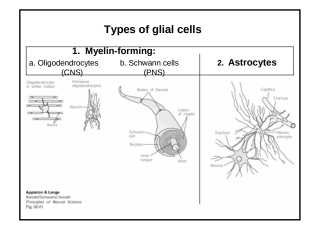


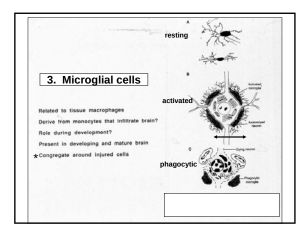
I. RESPONSE OF THE NEURON TO INJURY (summary)

- A. All neurons despite different morphologies - react similarly
- A. Principles
 - -If cell body damaged, the neuron dies, and is not replaced by cell division in mature brain.
 - -If the axon is damaged or severed at a distance from the soma, there is a good chance of regeneration, primarily in the PNS.
 - -CNS neurons have the capacity to regenerate.

I. RESPONSE OF THE NEURON TO INJURY

- II. GLOSSARY OF GLIAL CELLS: Normal function, response to injury
- III. DEGENERATION: Signs, Timecourse
- IV. REGENERATION
 - A. Neurons in the PNS can regenerate their axons. How?
 - Neurons in the CNS have a limited capacity to regenerate axons. Why?
- V. EXPERIMENTAL STRATEGIES TO PROMOTE REPAIR AND RECOVERY OF FUNCTION: Principles, examples





Myelin forming cells: (myelin important for conduction). oligodendroglia in CNS

Schwann cells in PNS.

oligodendrocytes (CNS) are inhibitory to axon regrowth in adult CNS regeneration;

Schwann cells (PNS) are supportive, as a growth surface and releaser of growth factors.

Astroglia -

development: supports axon growth and cell migration; mature: important for ion flux, synaptic function, blood-brain barrier

injury: accumulate in scar, release excess matrix; inhibit axon growth?

Microglia (resting) and macrophages (active) -

cells of immune system, similar to monocytes.

injury: help or hinder?

....not well-understood

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III. DEGENERATION: Signs, Timecourse

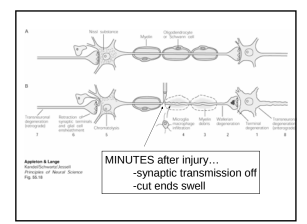
IV. REGENERATION

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REACTIONS TO INJURY WITHIN THE NEURON:

- Immediately
 1. Synaptic transmission off

 2. Cut ends pull apart and seal up, and swell, due to axonal transport in both directions



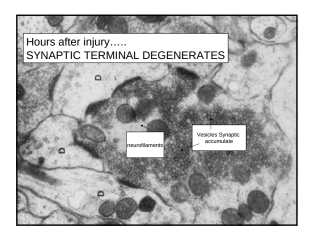
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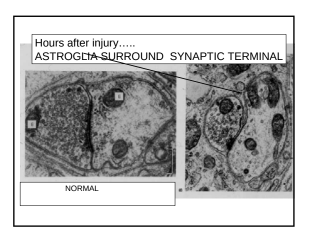
mmediately -

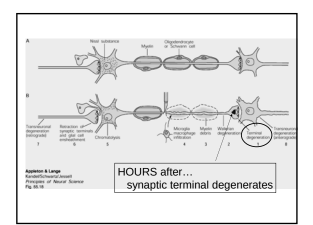
- Synaptic transmission off
- Cut ends pull apart and seal up, and swell,
 due to axonal transport in both directions

Hours -

- 3. Synaptic terminal degenerates accumulation of neurofilaments, vesicles.
- Astroglia surround terminal normally:
 after axotomy, astroglia interpose between terminal and target
 and cause terminal to be pulled away from postsynaptic cell.







REACTIONS TO INJURY WITHIN THE NEURON:

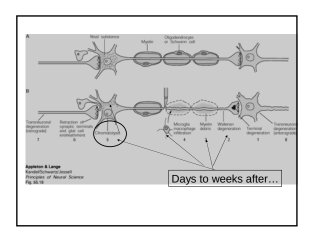
Immediately -

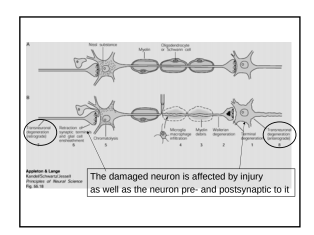
- Synaptic transmission off
 Cut ends pull apart and seal up, and swell, due to axonal transport in both directions

- Hours later 3. Synaptic terminal degenerates accumulation of NF, vesicles.
 4. Astroglia suround terminal normally;

after axotomy, interpose between terminal and target and cause terminal to be pulled away from postsynaptic

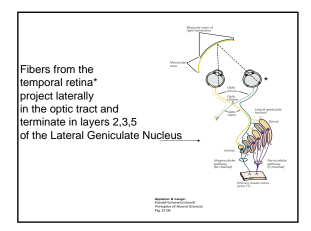
- days weeks 5. Myelin breaks up and leaves debris (myelin hard to break down).
- 6. Axon undergoes Wallerian degeneration
- 7. Chromatolysis cell body swells; nissl and nucleus eccentric.
 - **If axon cut in PNS or CNS, changes are the same.
- **The damaged neuron is affected by injury, as well as the pre- and postsynaptic neurons to it

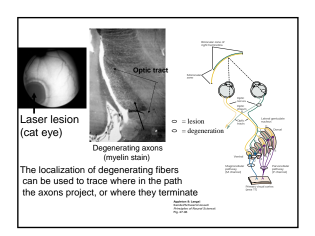




Severing the axon causes degenerative changes in the injured neuron AND in the cells that have synaptic connections with the injured neuron.

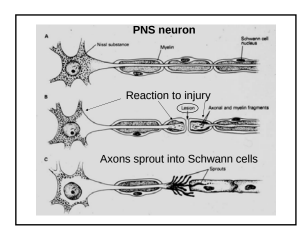
Classically, degeneration of fibers and their targets has been used to trace neuronal circuits experimentally, and still is used to understand pathology post-mortem

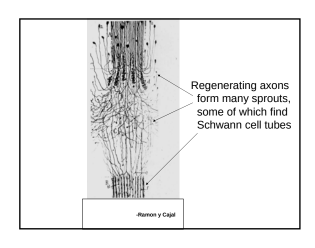


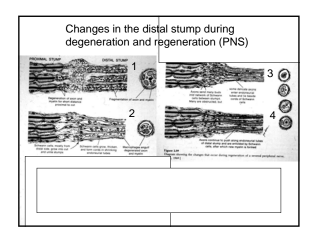


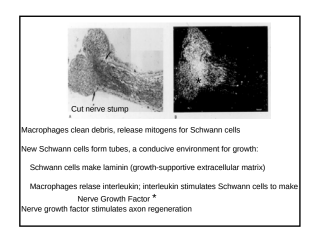
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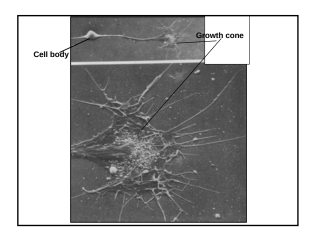
- II. GLOSSARY OF GLIAL CELLS: Normal function, response to injury
- III. DEGENERATION: Signs, Timecourse, applications of "reading" trans-synaptic degeneration
- IV. REGENERATION
 - A. Neurons in the PNS can regenerate their axons. How?
 - B. Neurons in the CNS have a limited capacity to regenerate axons. Why?
- V. EXPERIMENTAL STRATEGIES TO PROMOTE REPAIR AND RECOVERY OF FUNCTION: Principles, examples

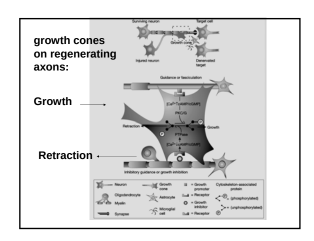












- IV. Neurons in the PNS can regenerate their axons. HOW? (summary)
- a. After degeneration of distal axon and myelin, macrophages clean up debris.
- b. Macrophages release mitogens that induce Schwann cells to divide
- c. The myelin-forming Schwann cells repopulate the nerve sheaths;
- d. Schwann cells make laminin
- e. Macrophages make interleukin, which induces Schwann cells to make Nerve Growth Factor.
- e. Axons sprout, and some sprouts enter new Schwann cell tubes
- f. Axonal growth cones successfully grow

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B. Neurons in the mature CNS have a limited capacity to regenerate axons. WHY?

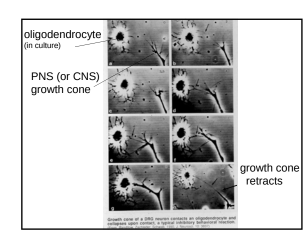
CNS axons can regrow, but...

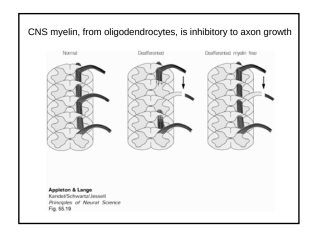
Growth is impeded by negative elements in the environment -myelin proteins (NOGO, MAG, Omgp) increase

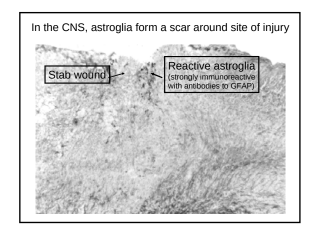
- inhibitory proteoglycans increase

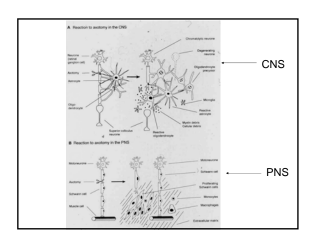
Intracellular growth factors such as GAP-43 (important for intracellular signaling/growth cone advance) are low

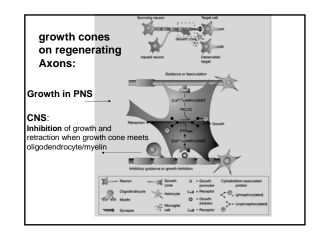
- -growth factors have different distributions compared to young brain
- -normally growth-supporting extracellular matrix (laminin) is sparse











B. Neurons in the CNS have a limited capacity to regenerate axons. WHY? (Summary)

CNS axons can regrow, but...

Growth is impeded by negative elements in the environment -extracelluar matrix (laminin) is sparse; inhibitory proteoglycans increas -growth factors have different distributions compared to young brain

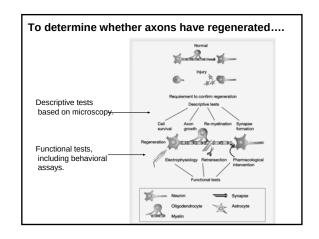
Intracellular growth elements such as GAP-43 (important for intracellular signaling/growth cone advance) are low

*Glial cells inhibit growth

Oligodendrocytes (CNS myelin) are the most inhibitory Astrocytes accumulate in the scar around injury site Macrophages also accumulate; role of microglia unclear

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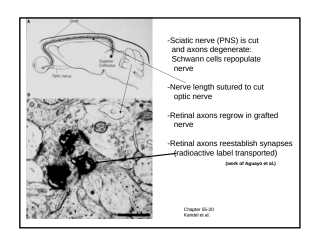


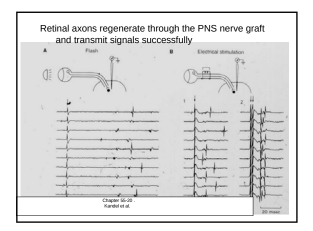
Therapeutic Strategies:

- 1. Implant
 - lengths of peripheral nerve (a natural "bridge")

Or

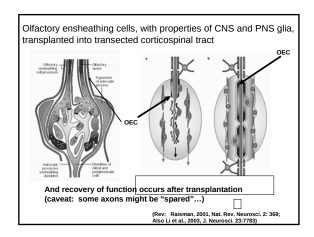
- artificial plastic tubes lined with supportive glia

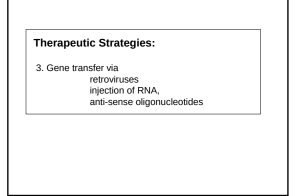


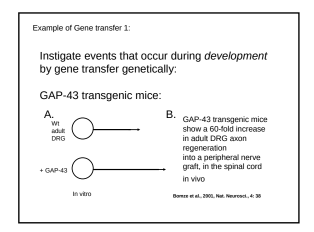


Therapeutic Strategies:

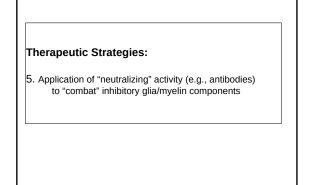
- 2. Transplant/ implant into or near site of injury:
 -fetal tissue (containing immature neurons and glia)
 or stem cells, with potential of becoming either
 - -cell lines or normal cells transfected with a gene for e.g., neurotrophins (positive) antibodies (against inhibitory myelin)
 - -"good" glia: olfactory ensheathing glia*

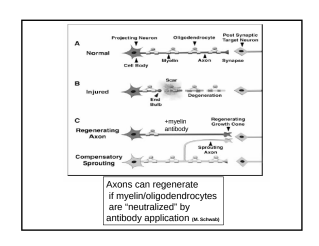












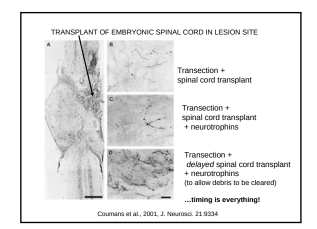
COMBINATION OF APPROACHES:

#2. Cellular Transplants

Transplant embryonic spinal cord

Plus....

#4. Delivery of growth factors



Embryonic spinal cord transplants plus neurotrophins lead to functional recovery after spinal cord transection

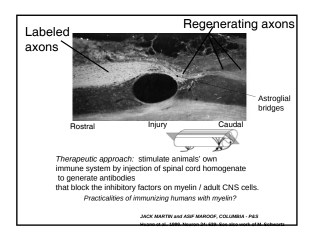
A

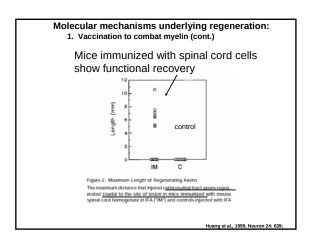
Transection only:
No weight support spinal cord tp + neurotrophins

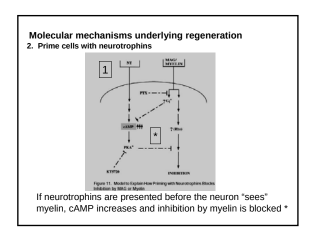
Figure 7. State accretion

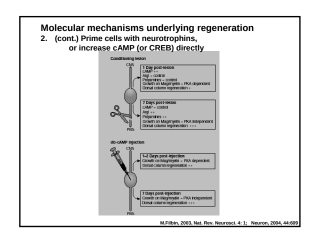
Molecular mechanisms underlying regeneration:

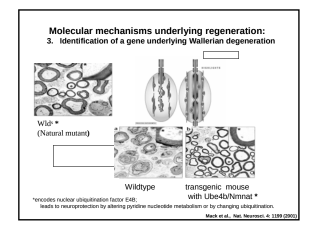
- 1. Vaccination to combat myelin
- 2. Prime cells with neurotrophins
- 3. Identification of a gene underlying Wallerian degeneration
- 4. Increase (good) microglia in eye by stabbing lens
- 5. Signals that travel from injury site back to nucleus
- Molecules that increase, decrease during inflammation, degeneration, regeneration
- 7. Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

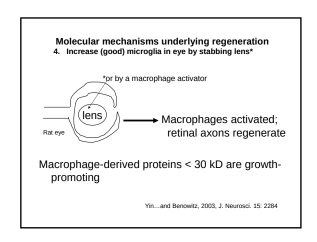


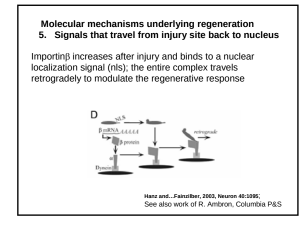


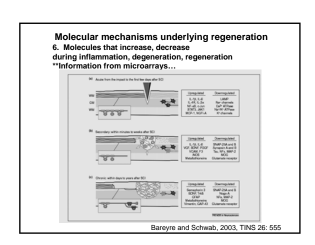


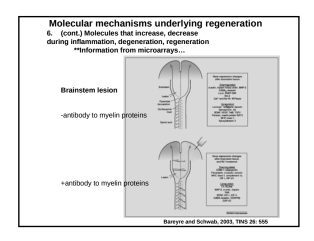


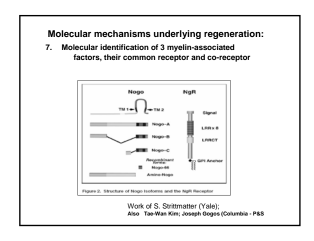


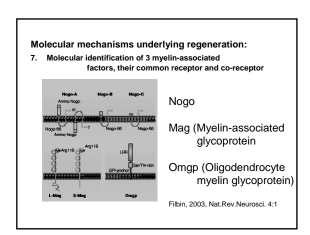


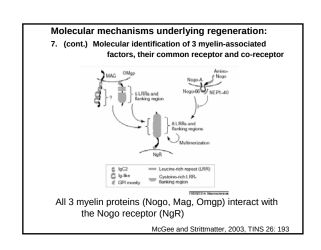


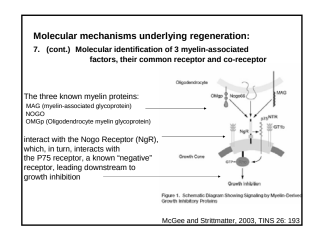


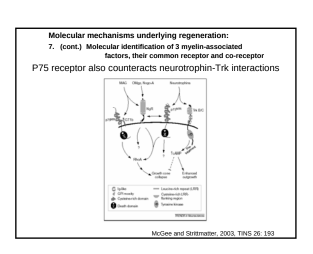












The bottom line...what treatments work in humans with spinal cord injury??

The case of Christopher Reeves...

Mice, cats, rats and humans that have been completely spinalized can regain greater locomotor performance if they are trained to perform that task, by robotics...

Edgerton and Roy, 2002, Curr Op Neurobiol 12:658

(Measures of recovery: Curt, Schwab, Deitz, 2004 Spinal Cord: 42:1)

