



*Instructions concerning a dislocation of a vertebra in the neck. "If you examine a man with a neck injury... and find he is without sensation in both arms and both legs, and unable to move them, and he is incontinent of urine... it is due to the laceration of the spinal cord caused by dislocation of a cervical vertebra. This is a condition which cannot be treated." Edwin Smith Surgical Papyrus, Case 27, Thebes, c. 1550 B.C. Taken from (Woodard, J. H. ed.) The Edwin Smith Surgical Papyrus © The University of Chicago Press, 1930.*

**GLIAL CELLS AND NERVE INJURY**

Carol Mason 1/27/04

**Symptoms of spinal cord injury:**

- involuntary muscle spasms
- loss 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**

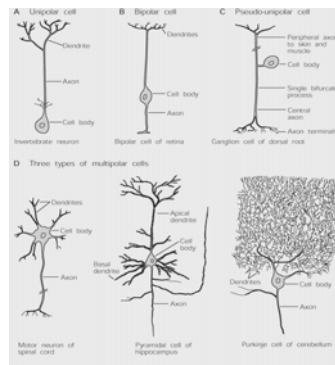
**I. RESPONSE OF THE NEURON TO INJURY**  
All neurons react similarly

**II. GLIAL CELLS:**  
Normal function  
Response to injury

**III. DEGENERATION:**  
Reactive changes, timecourse

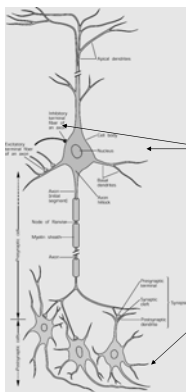
**IV. REGENERATION**  
A. Neurons in the PNS can regenerate. How?  
B. Neurons in the CNS have a limited capacity to regenerate. Why?

**V. EXPERIMENTAL STRATEGIES TO PROMOTE REPAIR / RECOVERY OF FUNCTION:** examples, recent reports



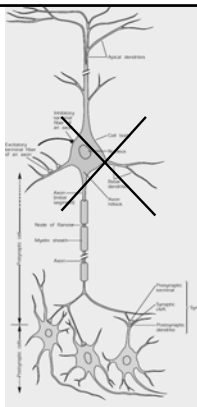
Neurons in the PNS and CNS have many different forms

Appelton & Lange  
Kandel/Chosson/Jessell  
Principles of Neural Science  
Fig. 12.14



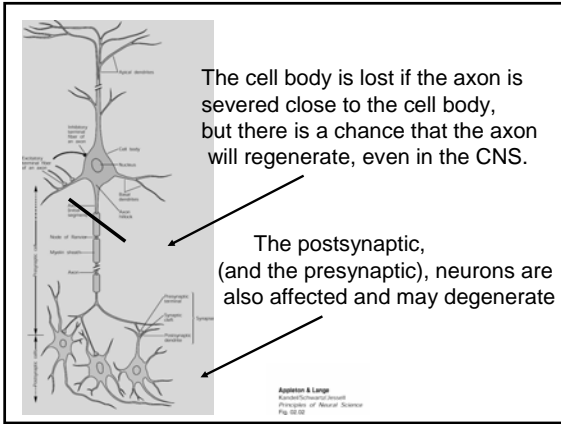
Cell biological reactions in the damaged neuron, presynaptic and postsynaptic neurons

Appelton & Lange  
Kandel/Chosson/Jessell  
Principles of Neural Science  
Fig. 12.12



If the cell body is damaged, the neuron is lost; there is no cell division in adult brain to replace the lost neuron.

Appelton & Lange  
Kandel/Chosson/Jessell  
Principles of Neural Science  
Fig. 12.12



**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

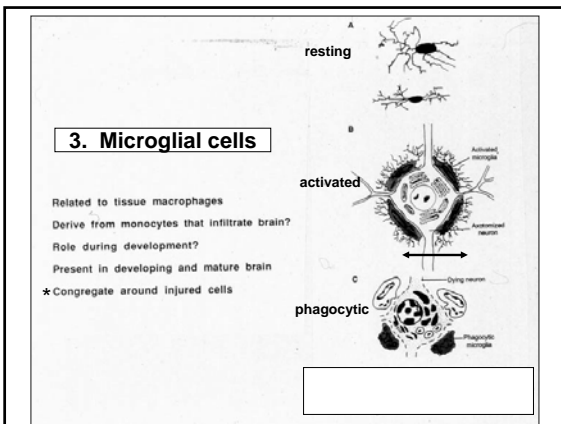
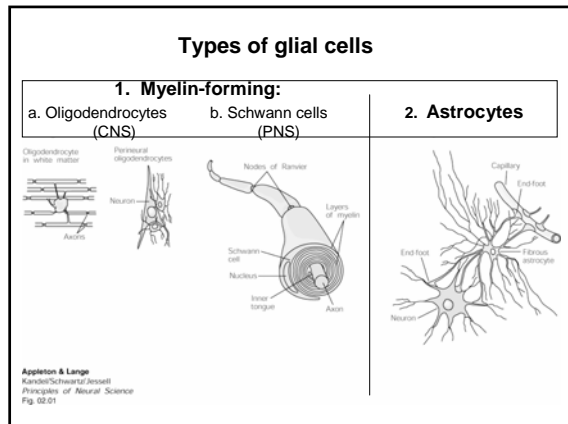
**III. DEGENERATION: Signs, Timecourse**

**IV. REGENERATION**

A. Neurons in the PNS can regenerate their axons. How?

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



**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

**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?

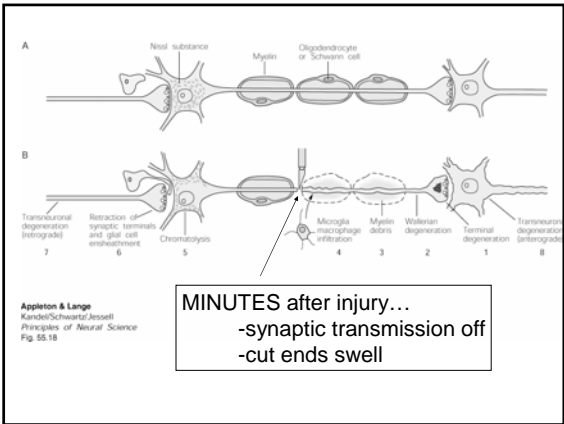
B. Neurons in the CNS have a limited capacity to regenerate axons. Why?

**V. EXPERIMENTAL STRATEGIES TO PROMOTE REPAIR AND RECOVERY OF FUNCTION: Principles, examples**

**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



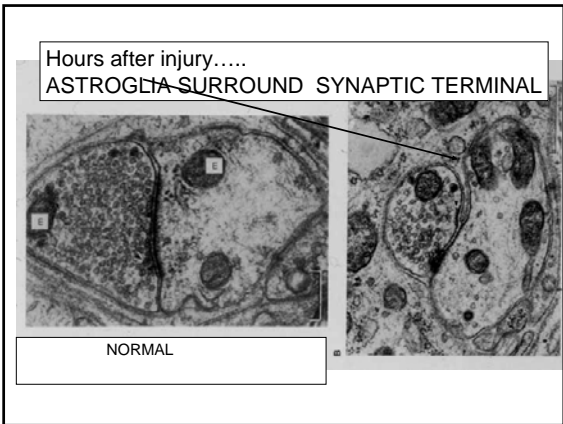
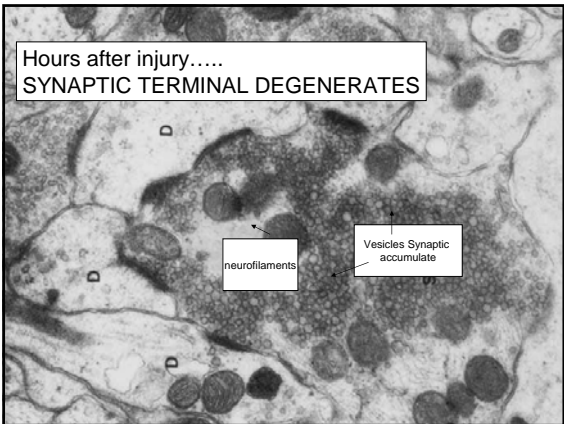
**REACTIONS TO INJURY WITHIN THE NEURON:**

**Immediately -**

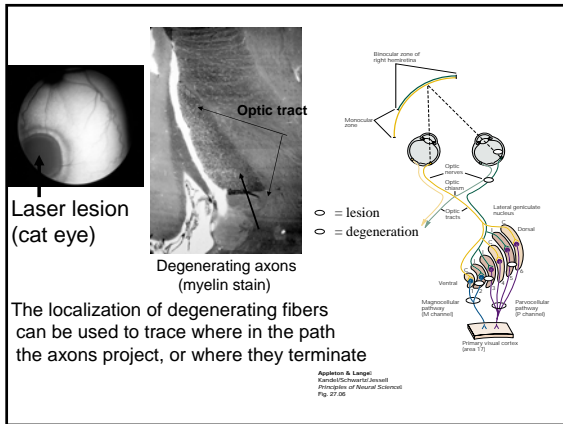
1. Synaptic transmission off
2. Cut ends pull apart and seal up, and swell, due to axonal transport in both directions

**Hours -**

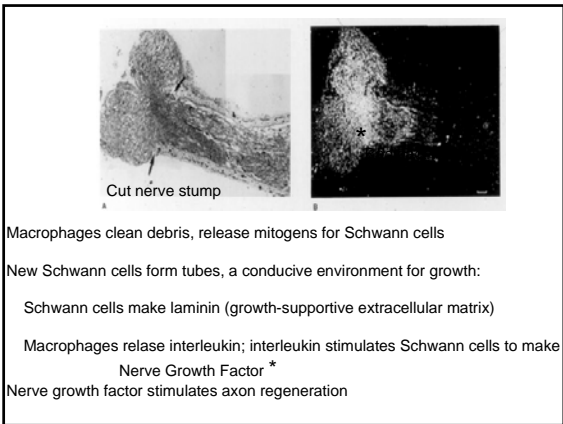
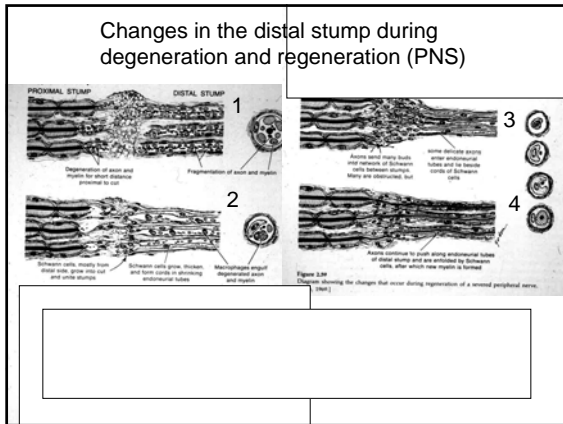
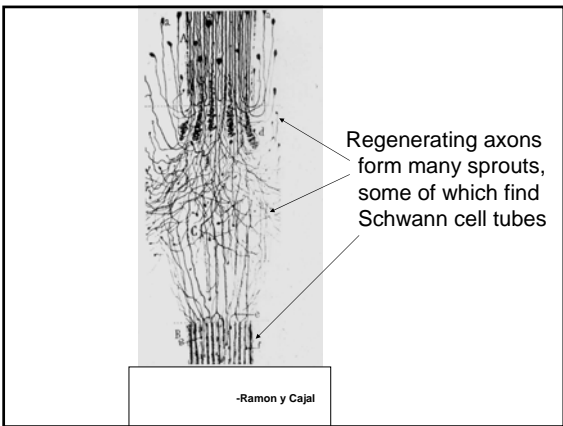
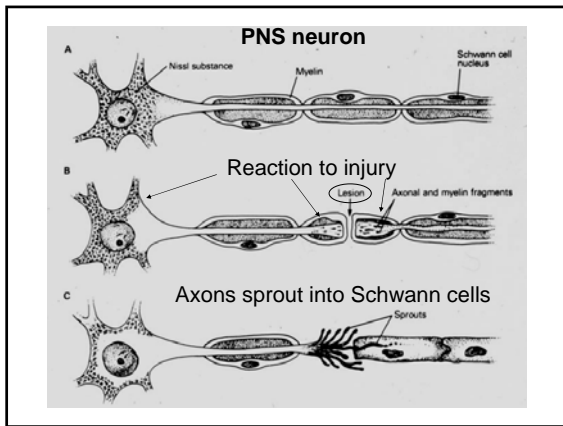
3. Synaptic terminal degenerates - accumulation of neurofilaments, vesicles.
4. Astroglia surround terminal normally; after axotomy, astroglia interpose between terminal and target and cause terminal to be pulled away from postsynaptic cell.

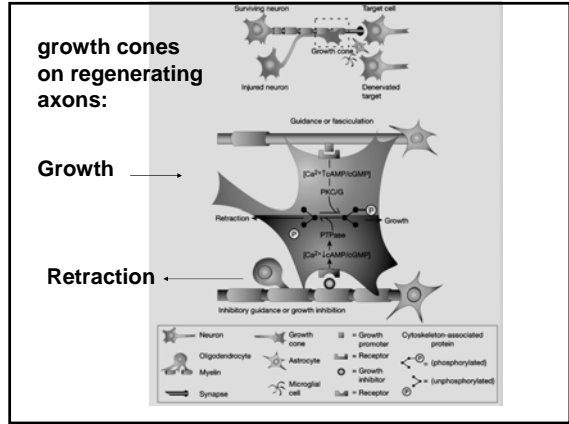
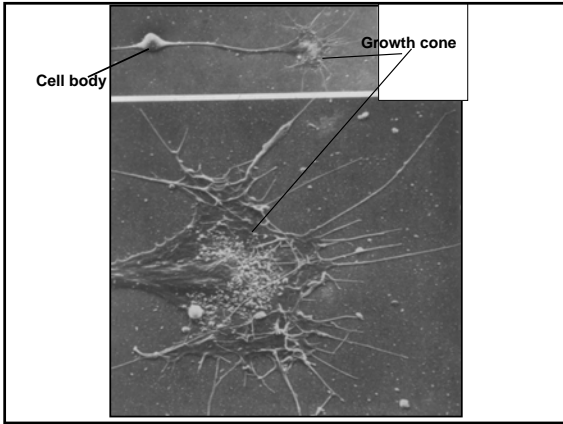






- I. RESPONSE OF THE NEURON TO INJURY
- II. GLOSSARY OF GLIAL CELLS: Normal function, response to injury
- III. DEGENERATION: Signs, Timecourse, applications of "reading" trans-synaptic degeneration
- IV. REGENERATION
  - A. Neurons in the PNS can regenerate their axons. How?
  - B. Neurons in the CNS 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)**

- After degeneration of distal axon and myelin, macrophages clean up debris.
- Macrophages release mitogens that induce Schwann cells to divide
- The myelin-forming Schwann cells repopulate the nerve sheaths;
- Schwann cells make laminin
- Macrophages make interleukin, which induces Schwann cells to make Nerve Growth Factor.

- Axons sprout, and some sprouts enter new Schwann cell tubes
- Axonal growth cones successfully grow

**I. RESPONSE OF THE NEURON TO INJURY**

**II. GLOSSARY OF GLIAL CELLS: Normal function, response to injury**

**III. DEGENERATION: Signs, Timecourse**

**IV. REGENERATION**

- 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**

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

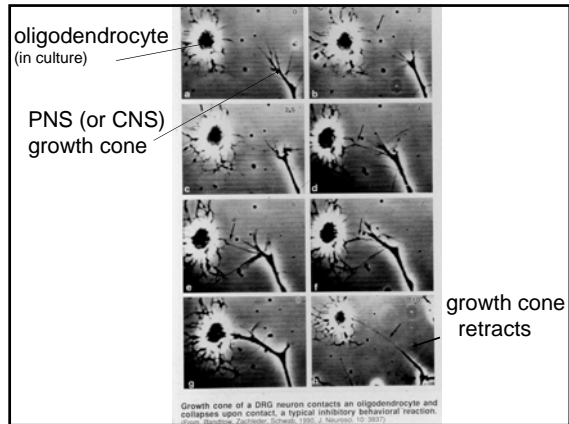
**CNS axons can regrow, but...**

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





**The exciting news: CNS neurons can sprout or grow.**

**Challenges:**

- \*Overcome the "bad" glial environment:
  - combat glial scars, inhibitory extracellular matrix
  - add blockers of myelin
  - repopulate with neurons and "good" glia
- \*Help axons regrow:
  - add neurotrophins (increase cAMP levels to prime neurons to ignore myelin-inhibitors).
  - re-express "youth" proteins - GAP-43
- \*Induce reformation of synapses (least understood step);
  - how do normal synapses form?

**To determine whether axons have regenerated....**

Descriptive tests based on microscopy.

Functional tests, including behavioral assays.

**Therapeutic Strategies:**

1. Implant
  - lengths of peripheral nerve (a natural "bridge")
  - Or
  - artificial plastic tubes lined with supportive glia

- Sciatic nerve (PNS) is cut and axons degenerate: Schwann cells repopulate nerve
- Nerve length sutured to cut optic nerve
- Retinal axons regrow in grafted nerve
- Retinal axons reestablish synapses (radioactive label transported)

(work of Aguayo et al.)

Chapter 55-20  
Kandel et al.

**Retinal axons regenerate through the PNS nerve graft and transmit signals successfully**

Chapter 55-20  
Kandel et al.

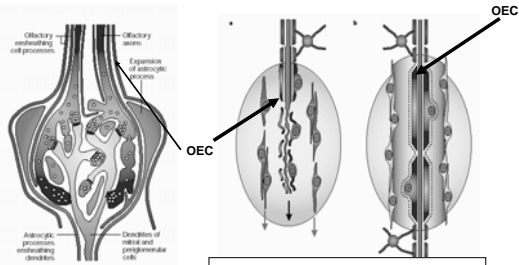
20 msec

**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\*



Olfactory ensheathing cells, with properties of CNS and PNS glia, transplanted into transected corticospinal tract



And recovery of function occurs after transplantation (caveat: some axons might be "spared"...)

(Rev: Ralsman, 2001, Nat. Rev. Neurosci. 2: 369; Also Li et al., 2003, J. Neurosci. 23:7783)

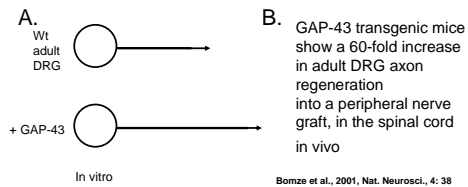
### Therapeutic Strategies:

- Gene transfer via
  - retroviruses
  - injection of RNA,
  - anti-sense oligonucleotides

Example of Gene transfer 1:

Instigate events that occur during *development* by gene transfer genetically:

GAP-43 transgenic mice:

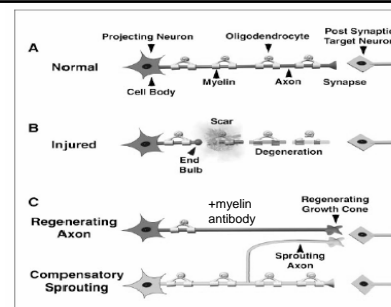


### Therapeutic Strategies:

- Direct delivery of growth factors to promote axon regrowth

### Therapeutic Strategies:

- Application of "neutralizing" activity (e.g., antibodies) to "combat" inhibitory glia/myelin components



Axons can regenerate if myelin/oligodendrocytes are "neutralized" by antibody application (M. Schwab)

**COMBINATION OF APPROACHES:**

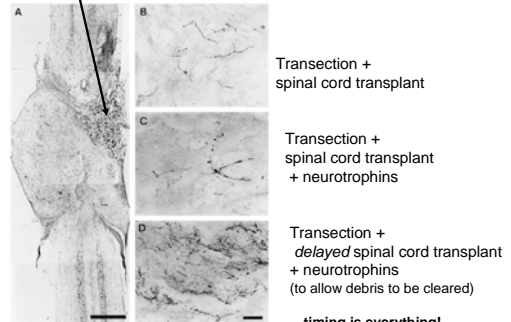
#2. Cellular Transplants

Transplant embryonic spinal cord

Plus....

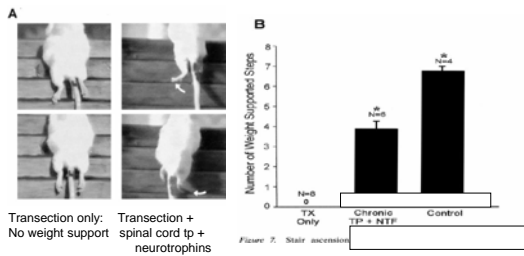
#4. Delivery of **growth factors**

**TRANSPLANT OF EMBRYONIC SPINAL CORD IN LESION SITE**



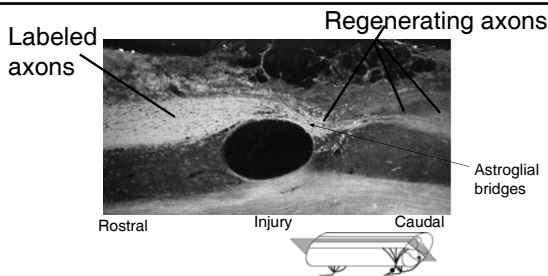
Coumans et al., 2001, J. Neurosci. 21:9334

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



**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
6. Molecules that increase, decrease during inflammation, degeneration, regeneration
7. Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor



**Therapeutic approach:** stimulate animals' own immune system by injection of spinal cord homogenate to generate antibodies that block the inhibitory factors on myelin / adult CNS cells.  
*Practicalities of immunizing humans with myelin?*

JACK MARTIN and ASIF MAROOF, COLUMBIA - P&S  
 Huang et al., 1999, Neuron 24: 629. See also work of M. Schwartz

**Molecular mechanisms underlying regeneration:**

1. Vaccination to combat myelin (cont.)

Mice immunized with spinal cord cells show functional recovery

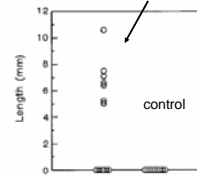


Figure 2. Maximum Length of Regenerating Axons  
 The maximum distance that injured corticospinal tract axons regenerated caudal to the site of lesion in mice immunized with mouse spinal cord homogenate in IFA (IM) and controls injected with IFA

Huang et al., 1999, Neuron 24: 639

**Molecular mechanisms underlying regeneration**  
**2. Prime cells with neurotrophins**

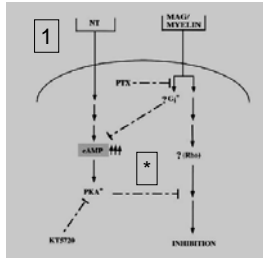
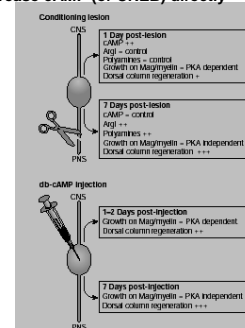


Figure 11. Model to Explain How Priming with Neurotrophins Blocks Inhibition by MAG or Myelin

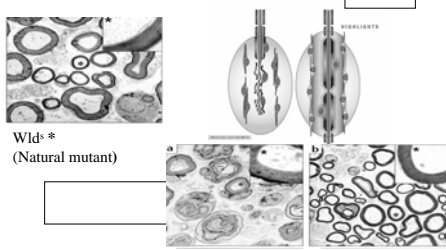
If neurotrophins are presented before the neuron "sees" myelin, cAMP increases and inhibition by myelin is blocked \*

**Molecular mechanisms underlying regeneration**  
**2. (cont.) Prime cells with neurotrophins, or increase cAMP (or CREB) directly**



M.Filbin, 2003, Nat. Rev. Neurosci. 4: 1; Neuron, 2004, 44:609

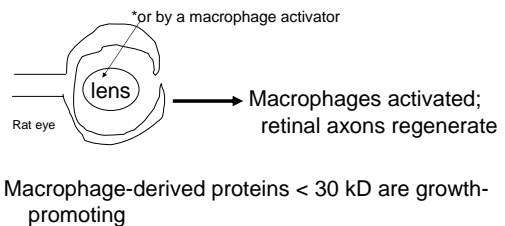
**Molecular mechanisms underlying regeneration:**  
**3. Identification of a gene underlying Wallerian degeneration**



\*encodes nuclear ubiquitination factor E4B; leads to neuroprotection by altering pyridine nucleotide metabolism or by changing ubiquitination.

Mack et al., Nat. Neurosci. 4: 1199 (2001)

**Molecular mechanisms underlying regeneration**  
**4. Increase (good) microglia in eye by stabbing lens\***

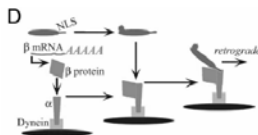


Macrophage-derived proteins < 30 kD are growth-promoting

Yin...and Benowitz, 2003, J. Neurosci. 15: 2284

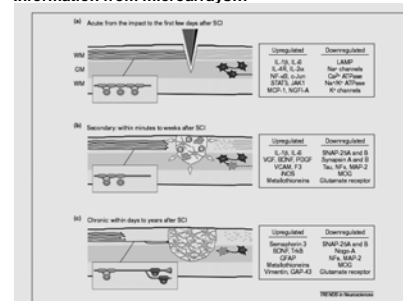
**Molecular mechanisms underlying regeneration**  
**5. Signals that travel from injury site back to nucleus**

Importinβ increases after injury and binds to a nuclear localization signal (nls); the entire complex travels retrogradely to modulate the regenerative response



Hanz and...Fainzilber, 2003, Neuron 40:1095; See also work of R. Ambron, Columbia P&S

**Molecular mechanisms underlying regeneration**  
**6. Molecules that increase, decrease during inflammation, degeneration, regeneration**  
**\*\*Information from microarrays...**



Bareyre and Schwab, 2003, TINS 26: 555

**Molecular mechanisms underlying regeneration**  
 6. (cont.) Molecules that increase, decrease during inflammation, degeneration, regeneration  
 \*\*Information from microarrays...

**Brainstem lesion**

-antibody to myelin proteins

+antibody to myelin proteins

Bareyre and Schwab, 2003, TINS 26: 555

**Molecular mechanisms underlying regeneration:**  
 7. Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

Figure 2. Structure of Nogo isoforms and the NgR Receptor

Work of S. Strittmatter (Yale);  
 Also Tae-Wan Kim; Joseph Gogos (Columbia - P&S)

**Molecular mechanisms underlying regeneration:**  
 7. Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

Nogo  
 Mag (Myelin-associated glycoprotein)  
 Omgp (Oligodendrocyte myelin glycoprotein)

Filbin, 2003, Nat.Rev.Neurosci. 4:1

**Molecular mechanisms underlying regeneration:**  
 7. (cont.) Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

All 3 myelin proteins (Nogo, Mag, Omgp) interact with the Nogo receptor (NgR)

McGee and Strittmatter, 2003, TINS 26: 193

**Molecular mechanisms underlying regeneration:**  
 7. (cont.) Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

The three known myelin proteins:  
 MAG (myelin-associated glycoprotein)  
 NOGO  
 Omgp (Oligodendrocyte myelin glycoprotein)

interact with the Nogo Receptor (NgR), which, in turn, interacts with the P75 receptor, a known "negative" receptor, leading downstream to growth inhibition

Figure 1. Schematic Diagram Showing Signaling by Myelin-Derived Growth Inhibitory Proteins

McGee and Strittmatter, 2003, TINS 26: 193

**Molecular mechanisms underlying regeneration:**  
 7. (cont.) Molecular identification of 3 myelin-associated factors, their common receptor and co-receptor

P75 receptor also counteracts neurotrophin-Trk interactions

McGee and Strittmatter, 2003, TINS 26: 193

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

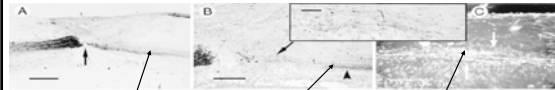
The case of Christopher Reeves...

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)

### Molecular mechanisms underlying regeneration: 1. Vaccination to combat myelin



(A) Micrograph of a longitudinal section of the lesioned corticospinal tract in a mouse immunized with spinal cord homogenate in IFA. WGA-HRP labels the tract rostral and caudal to the lesion (arrow). Many regenerated axons can be seen caudally.

*Therapeutic approach:* stimulate animals' own immune system by injection of spinal cord homogenate to generate polyclonal antibodies that block the inhibitory factors on myelin / adult CNS cells.  
*Practicalities of immunizing humans with myelin?*

Huang et al., 1999, Neuron 24: 639;  
See also work of M. Schwartz