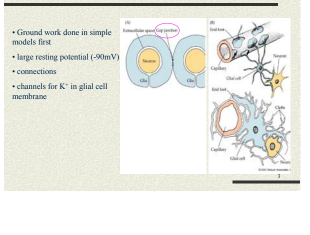
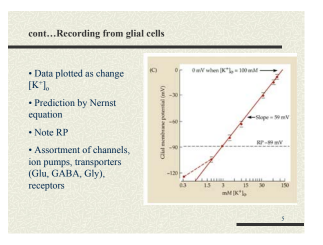
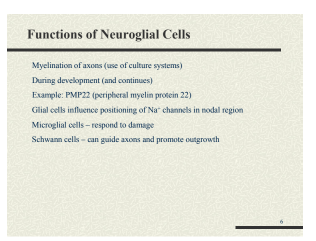


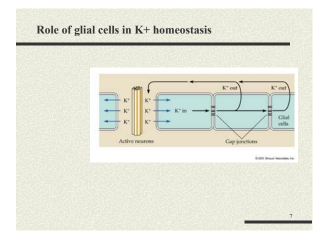
Fig. 8.1



Connectivity amongst glia, neurons, and capillaries



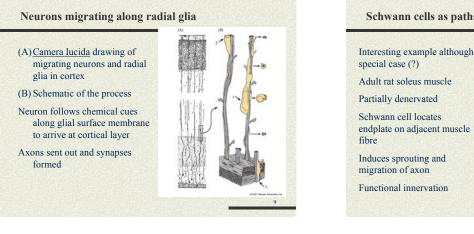




# cont...Functions of Neuroglial Cells

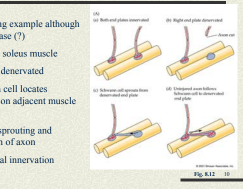
# During development

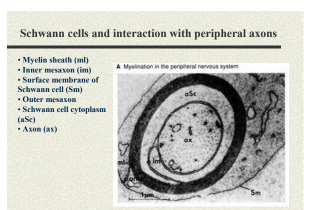
- Groupings of neurons into nuclei these nuclei are first outlined by glial cells (neurons arriving later)
- Radial glial cells used to guide migrating neurons (in cerebral cortex, hippocampus, cerebellum) (next slide)



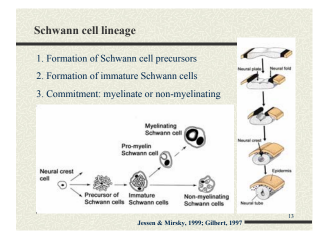
11

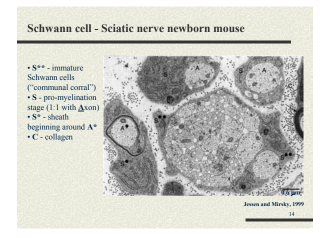
# Schwann cells as paths for outgrowth in PNS

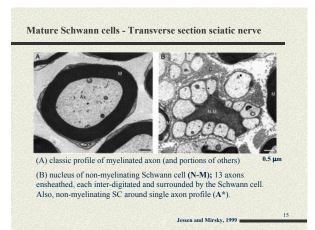




# Schwann Cells - the basics eripheral glial cells - myelinating and non-myelinating eilance on signalling from axons eneron-derived signals during development and when mature ew evidence supports glial-derived signalling as critical for neuronal survival during specific periods of development egulate molecular and f'al specialization's of axons; maturation of perineurial sheath







#### Summary points on Schwann cells

- glial lineage arises from neural crest (NC) cells
- major peripheral myelin protein (Po) found to be earliest glial cell marker (found in migrating NC cells)\*\*
- Po expression modulated by axons (up and down...)
- narrow window for transition from precursors to Schwann cell (E14/E15--E17 rat (mouse E15)) ("no" precursors in mature nerves)
- β-neuregulins (axonal) bias NC cells to differentiate to glial cell, although some controversy needs to be resolved

\*\*Bhattacharyya et al. 1991; Lee et al. 1997; etc.

# cont. summary points...1

- $\bullet$  dependence on signalling from axons for survival ( $\beta$ -neuregulin)
- Evidence: in vitro cultures and KO's,  $\beta$ -neuregulin essential for precursor cell survival and the change from precursor to glial cell
- period from about birth to 3 weeks get final differentiation step
- membrane synthesis, up and down regulation of genes
- transection of nerve leads to changes which revert glial phenotype to immature state
- · environment formed which would promote axonal re-growth
- ... and new evidence from new technology

# cont. summary points...2

#### Knock-out of Erbb3 gene

- a major receptor for β-neuregulin in crest cells and early glia
- initially number of DRG and motor neurons normal during embryogenesis (ca. E12)
- these mice lack Schwann-cell precursors and Schwann cells
- $\bullet$  by E14, 80% of DRG neurons lost; by E18, 80% motorneurons were lost (as late as E16 all OK)
- chimeric experiments (Erbb3 in neurons but not 'glia')

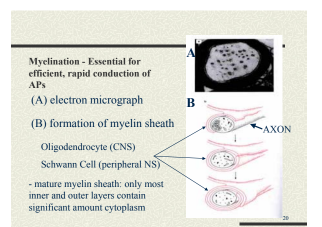
# cont. summary points...2

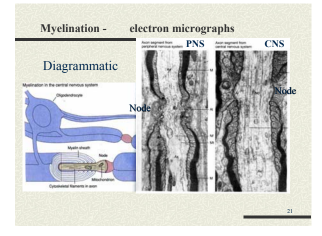
- DRG RIP too early to receive trophic signals from targets
- motorneurons last until E18 then die why?

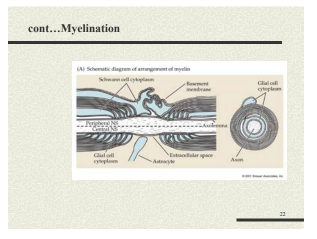
\* initial survival and migration to target independent of signals from immature glial cells

\* BUT: target-derived and glial signals required for survival

\* Note timing: link to transformation of glial precursors to immature glial cells usually occurs just prior to E18







# cont. Myelination

- Myelin interrupted at nodes of Ranvier (1 1.5mm spacing)
- Measurements made indicate CV for fibres  $> 11 \mu m$  is 6 times axon diameter; fibres  $< 11 \mu m$  about 4.5 X

• Balance: thickness of myelin (increases R) and cross-sectional area of axon (decreases - causes increase in internal longitudinal R) - compromise: axon diameter 0.7 x overall fibre diameter

Distance between nodes optimized

#### cont. Myelination

• Single Schwann cell makes myelin in one internode region (*ca.* 500 needed for single peripheral axon); oligodendrocyte can do several axons

- Formation of myelin by Schwann cells appears to be axon dependent-signaling; oligodendrocytes rely on astrocytes for signaling
- Myelin Basic Proteins found in both; group of 7 related proteins (alternative splicing variants)

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# cont. Myelination

- Classic experiments done by Ritchie and co-workers (mostly on rabbit nerves)
- Location of V-gated channels not what you might expect!
- $\textit{or}_{\bullet}$  Na<sup>+</sup> channels conc'd in nodes of Ranvier; none paranodal
  - K<sup>+</sup> channels conc'd under sheath (between nodes)
- V/C showed nodes displayed only inward currents and repol'n **NOT** by an increase of  $G_{K}$ + then what?

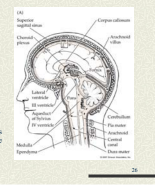
• Chronic demyelination by **diphtheria toxin** - Na<sup>+</sup> channels eventually populate demyelinated region and then get continuous conduction through the area, but poor substitute

# The blood brain barrier (BBB)

#### 3 main compartments

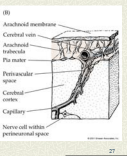
- Blood in capillariesCSF surrounds bulk of NS,
- contained in ventricles
- Intercellular clefts

 Endothelial cells of capillaries specialized to be less permeable
 Most substances blocked; not lipophilic or gases (dissolved)
 Choroid plexus: specialized epithelial cells surround cp capillaries. These cells produce and secrete CSF.
 Intercellular clefts (20 nm): gateway to neurons



# cont...The blood brain barrier (BBB)

Fluid movement thru intercellular spaces, not thru glia (experiment: inject HRP into, product from peroxidase rx electron dense, look at distribution)



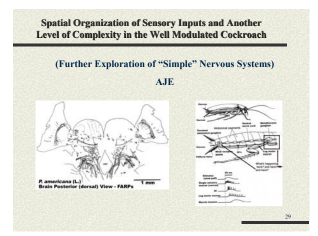
# Parting shots at glial cells

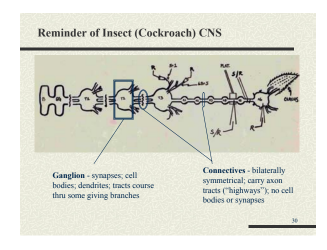
Glial cells act to separate individual or groups of neurons

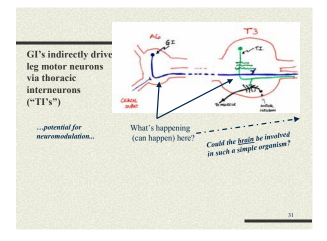
- Help regulate [K<sup>+</sup>] in extracellular environment
- Transmitters can act on glial membranes role ?

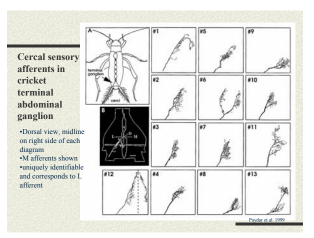
Glutamate transporter in glial cells

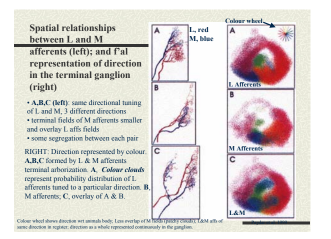
What if persistent high [Glu]<sub>0</sub>? (mice that lack gene for astrocytic glu-transporter (GLT-1) develop epilepsy and increased susceptibility to convulsants)

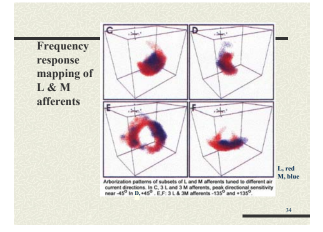


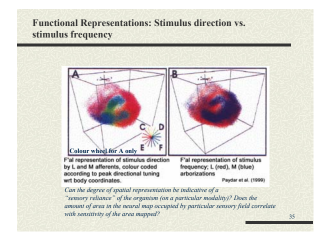


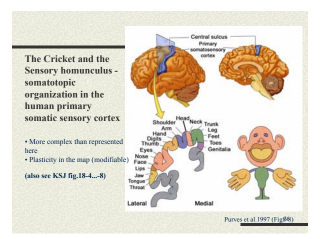












# Continuing saga of the "well modulated" cockroach

#### Recall,

neurons

Octopamine (OA), dopamine (DA), and serotonin (5-HT) as putative neuromodulators

· effects on thoracic interneurons that drive motor neurons

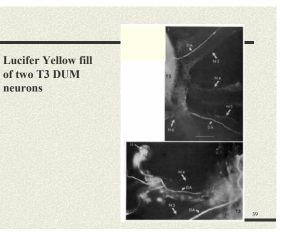
# Role of FMRFamide-like peptides

- peripheral innervation of skeletal muscles
- · central release sites
- release into haemolymph (blood)

#### Identifying specific neurons involved in modulation of activity in T3 and in skeletal muscle (Dorsal Unpaired Median (DUM) Neurons (peripheral and central

(?) connections)

# Patterns of innervation by DUM neurons in T3 • DUM3,5,6; DUM3,4,5,6; DUM3,5; DUM5,6 • Where do they go and what do they do? · Central vs. peripheral roles Ckrch T3 DUM Neurons (cell bodies not shown



## **Recruitment of DUM Neuron during WP-evoked** escape response

