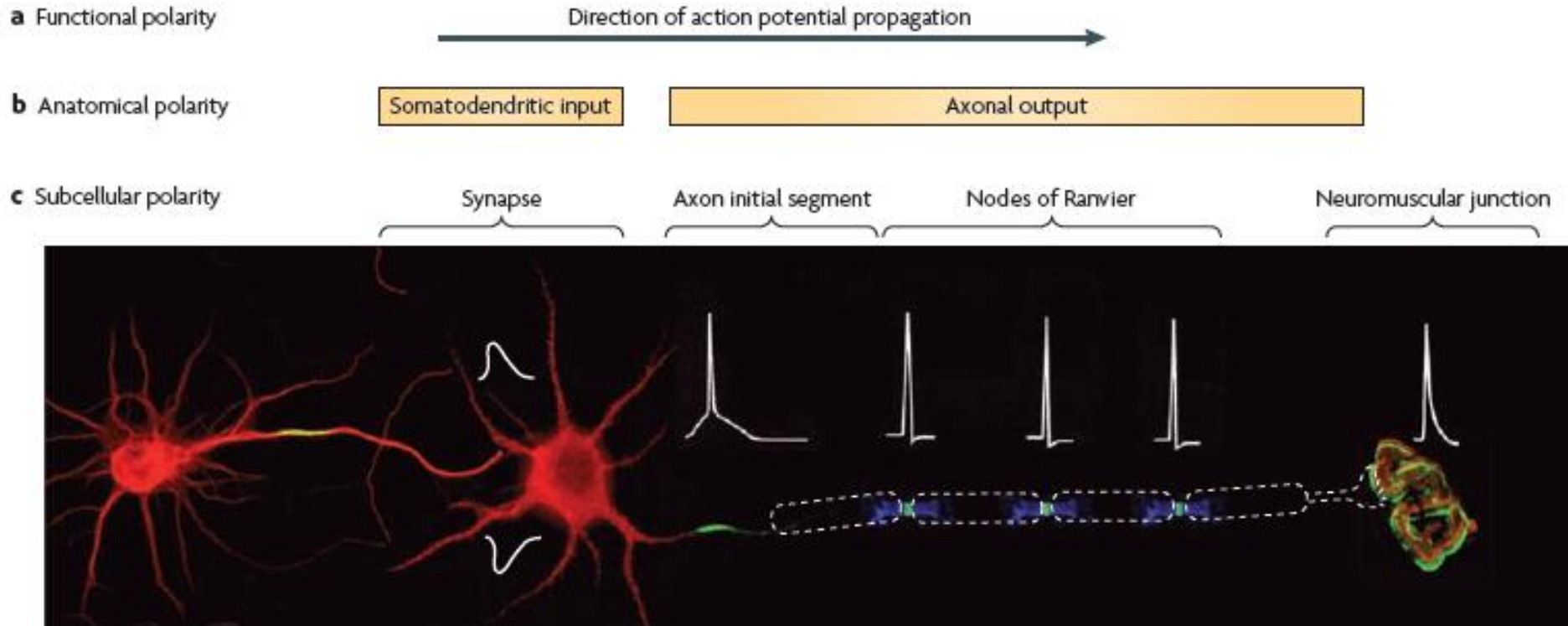


Az elektromosan ingerlékeny membrán  
szerkezete és a mielinizáció

# Az idegi sejtmembrán polarizáltsága

The axon initial segment and the maintenance of neuronal polarity

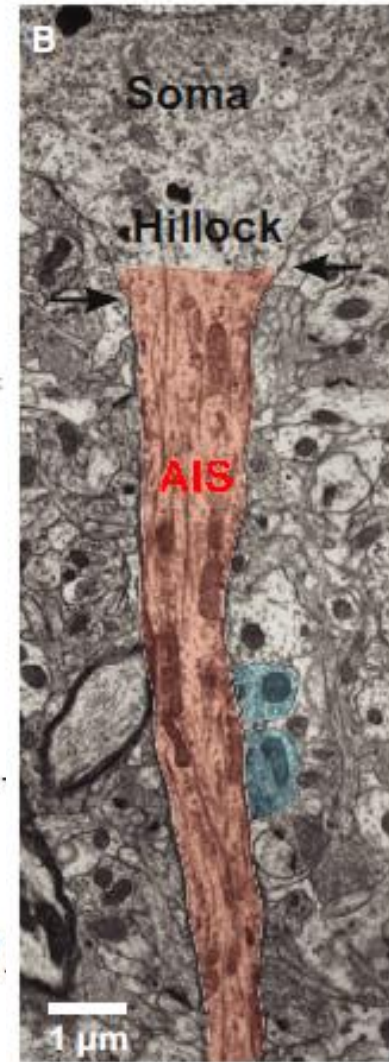
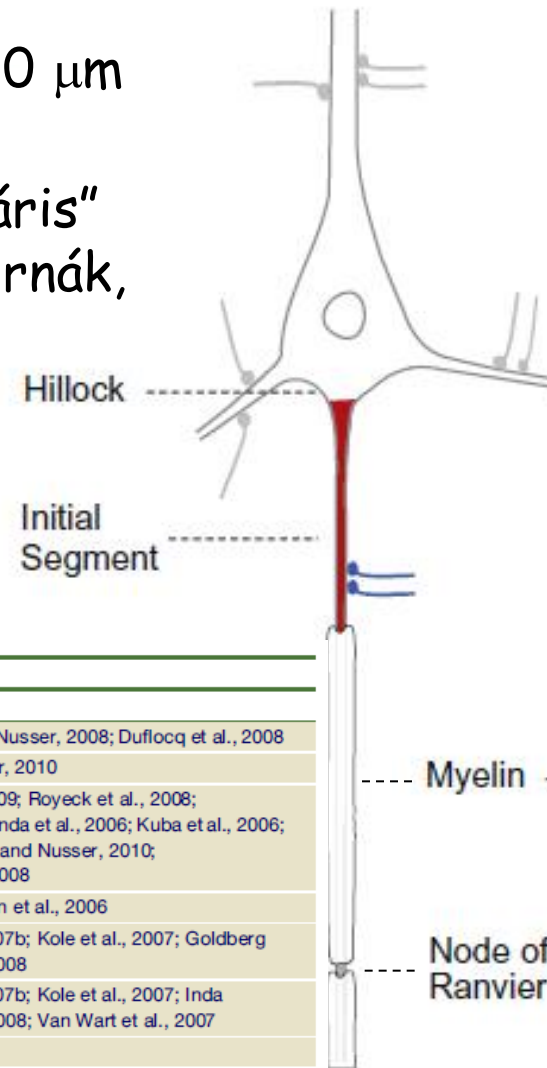
*Matthew N. Rasband*



**Figure 1 | Neurons are highly polarized cells.** **a** | Neurons are functionally polarized because action potentials propagate in a single direction. Excitatory and inhibitory synaptic inputs are integrated at the axon initial segment (AIS). The resulting action potentials then propagate along the axon through the activity of ion channels clustered at nodes. Finally, neurotransmitter is released at the nerve terminal. **b** | Neurons are also anatomically polarized, as they can be subdivided into a somatodendritic input domain and an axonal output domain. The AIS separates these two domains. **c** | Neurons have a high degree of subcellular polarity, and synapses, the AIS, nodes of Ranvier and the neuromuscular junction are the main subcellular domains. Each of these domains is enriched in specific types of ion channels, receptors, adhesion molecules and molecular scaffolds that allow for the unidirectional propagation of action potentials. Each of these subcellular domains can also elicit unique electrophysiological responses (shown in white).

# Az axon kezdeti (iniciális) szakasza (AIS)

- gerincesekben, nem mielinizált, 10-60  $\mu\text{m}$  hosszú membránszakasz
- plazmamembrán alatti denz, „granuláris” réteg ~ Ranvier befűződés: ioncsatornák, állványfehérjék
- gyakori GABAerg szinapszisok
- ER:  $\text{Ca}^{2+}$  raktár (~spine apparatus)



(B) Electron micrograph of the AIS of a cortical pyramidal neuron. The soma, axon hillock, and AIS (red) are indicated. Blue areas indicate presynaptic terminals onto the AIS. Arrows (black) indicate the onset of the dense granular layer beneath the surface membrane indicating the start of the AIS. Adapted from Peters et al. (1968).

Table 1. Ion Channel Expression Patterns in the Axon Initial Segment

Current	Channel	Cell Type	Reference
$I_{\text{NaT}}$	$\text{Na}_v1.1$	IN, RGC, MN	Van Wart et al., 2007; Lorincz and Nusser, 2008; Duflocq et al., 2008
	$\text{Na}_v1.2$	PC	Hu et al., 2009; Lorincz and Nusser, 2010
	$\text{Na}_v1.6$	PC, DG, RGC, PN, IN, MN	Van Wart et al., 2007; Hu et al., 2009; Royeck et al., 2008; Boiko et al., 2003; Catterall, 1981; Inda et al., 2006; Kuba et al., 2006; Lorincz and Nusser, 2008; Lorincz and Nusser, 2010; Kress et al., 2010; Duflocq et al., 2008
$I_{\text{NaP}}$	$\text{Na}_v1.7$	PC	Stuart and Sakmann, 1995; Astman et al., 2006
$I_{\text{D}}$	$\text{K}_v1.1$	PC, MNTB, IN	Dodson et al., 2002; Shu et al., 2007b; Kole et al., 2007; Goldberg et al., 2008; Lorincz and Nusser, 2008
	$\text{K}_v1.2$	PC, RGC, MNTB, IN	Dodson et al., 2002; Shu et al., 2007b; Kole et al., 2007; Inda et al., 2006; Lorincz and Nusser, 2008; Van Wart et al., 2007
	$\text{K}_v2.2$	MNTB	Johnston et al., 2008
$I_{\text{A}}$	$\text{K}_v1.4$	PC	Ogawa et al., 2008
$I_{\text{M}}$	$\text{K}_v7.2 / \text{K}_v7.3$	PC	Pan et al., 2006; Shah et al., 2008
	$\text{K}_v7.2$	MN, DG	Devaux et al., 2004; Klinger et al., 2011
$I_{\text{Ca}}$ (T/R-type)	$\text{Ca}_v2.3 / \text{Ca}_v3.2 / \text{Ca}_v3.1$	CC, PC, PN	Bender and Trussell, 2009
$I_{\text{Ca}}$ (P/Q/N-type)	$\text{Ca}_v2.1/\text{Ca}_v2.2$	PC; PN	Calleraert et al., 1996; Yu et al., 2010

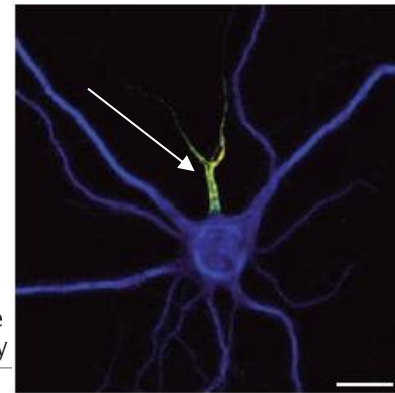
IN, interneuron; PC, pyramidal cell; DG, dentate granule cell (hippocampus); PN, Purkinje neuron; RGC, retinal ganglion cell; MNTB, medial nucleus of the trapezoid body; CC, cartwheel cell; MN, spinal cord motoneuron.

Neuron 73, January 26, 2012

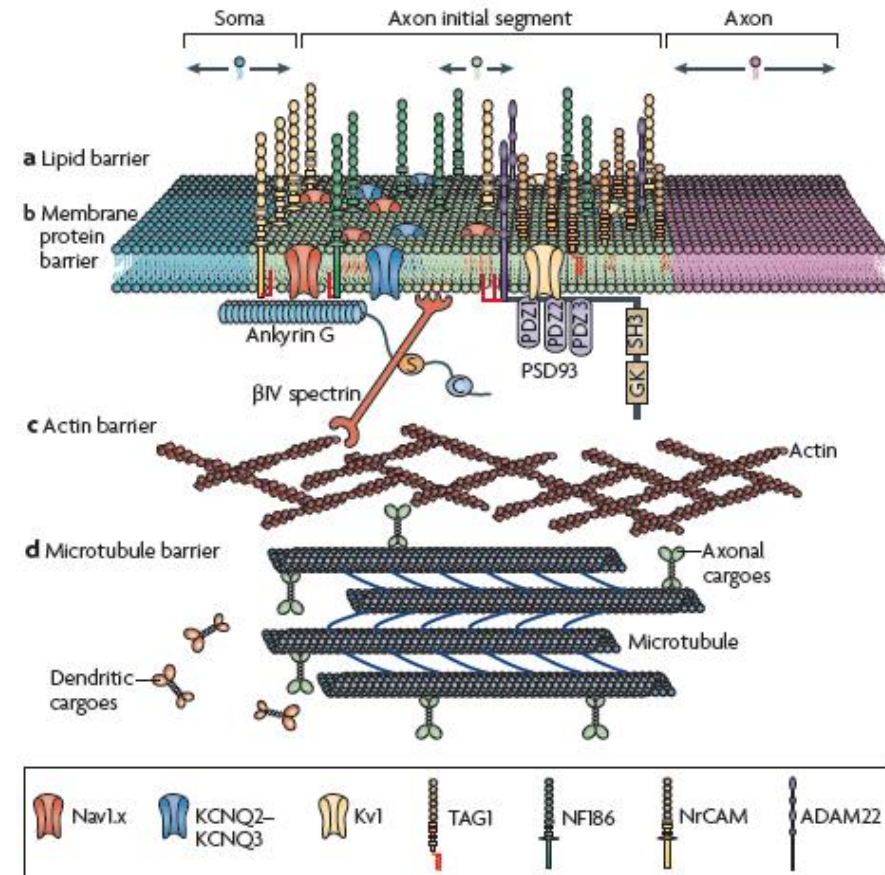
Signal Processing in the Axon Initial Segment

# Az axon kezdeti (iniciális) szakasza (AIS)

- fizikai „gát” az axonális és szomatodendritikus membránkomponensek laterális diffúziója ellen
  - + citoplazmás szűrő (aktin: méret alapján?)
  - + vezikulák szelektív átengedése (MT)
- axon/dendrit differenciációval párhuzamosan alakul ki
  - szinaptikus kapcsolatok előtt
- fesz.függő  $\text{Na}^+$  ( $\text{Na}_V$ ) és  $\text{K}^+$  ( $\text{K}_V$ ) csatornák nagy denzitásban + kis átmérő  $\rightarrow$  akciós potenciál
  - 35-45  $\mu\text{m}$  hosszan
  - $>50\times$  sűrűség (kevesebb?)
- Ranvier-befűződéshez hasonló szerkezet, de az AIS kialakulása gliasejt-független



ankyrinG / neurofascin 186

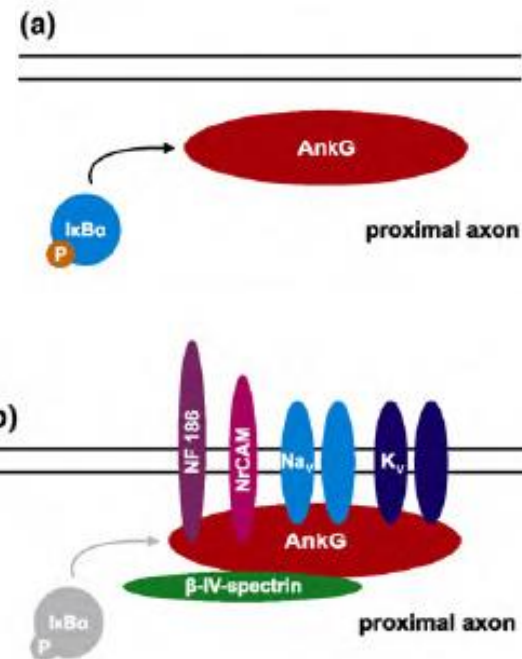


# Az AIS kialakulása

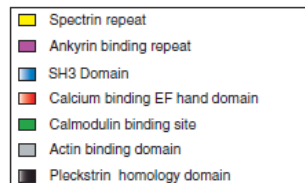
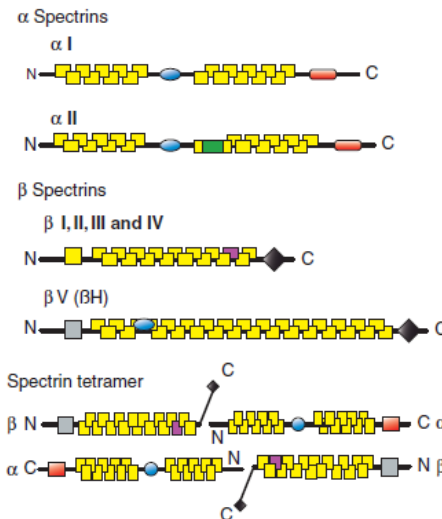
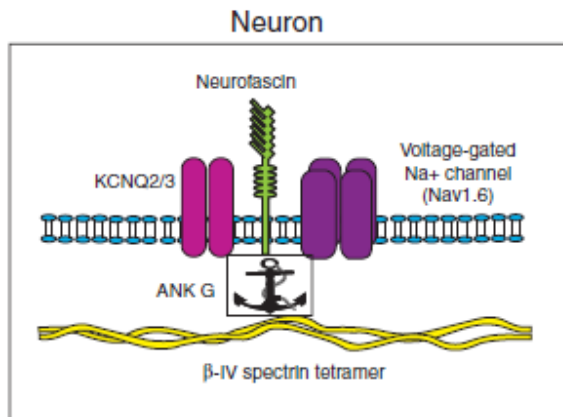
1. proximális axonban ankyrinG felhalmozódás  
( $I\kappa B\alpha$  foszforiláció elősegíti?)

2. ankyrinG: scaffold fehérje

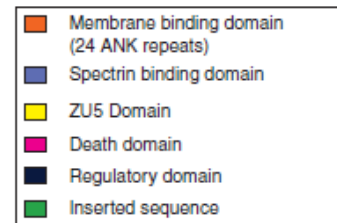
- adhéziós molekulák (NF186 [neurofascin], NrCAM)
- $\beta$ -IV spectrin  $\rightarrow$  aktin



Building and maintaining the axon initial segment  
Matthew S Grubb and Juan Burrone



## Ankyrins



Canonical ankyrins: B, G and R

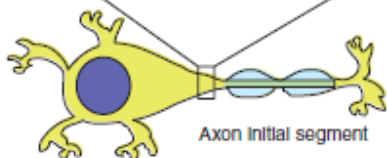


Neuronal variants: B and G



Membrane Domains Based on Ankyrin and Spectrin Associated with Cell-Cell Interactions

Vann Bennett and Jane Healy

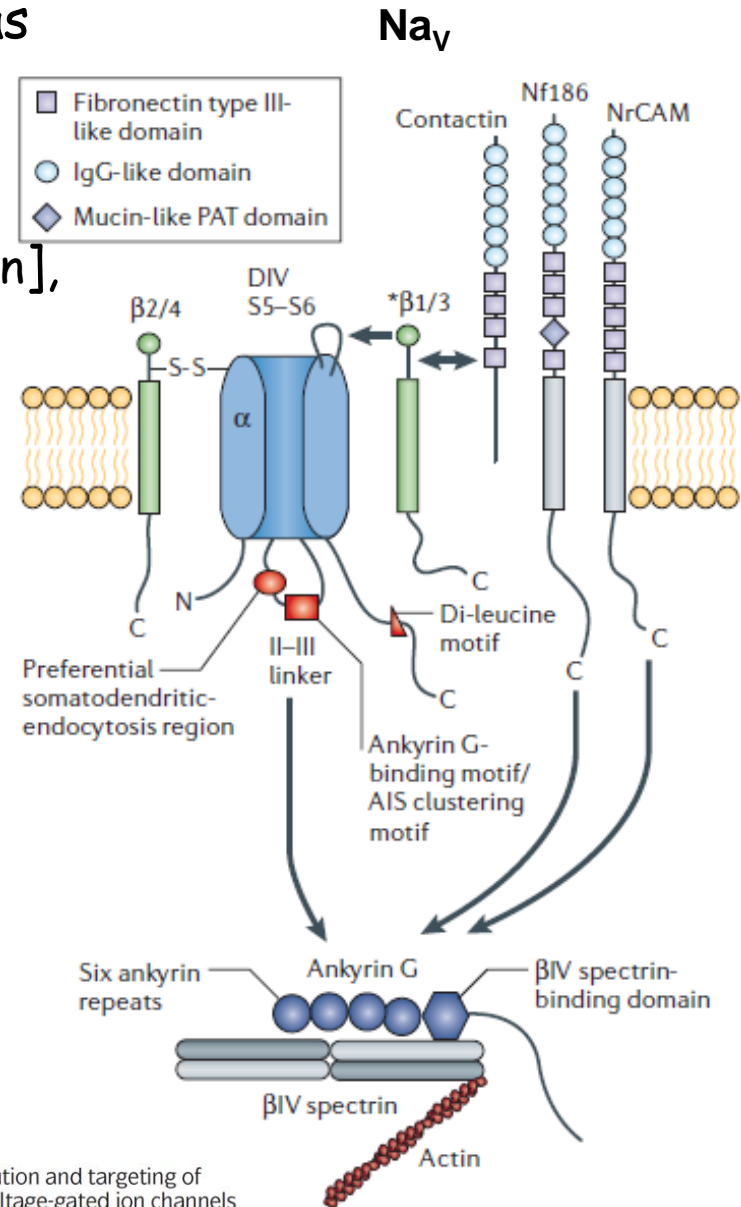
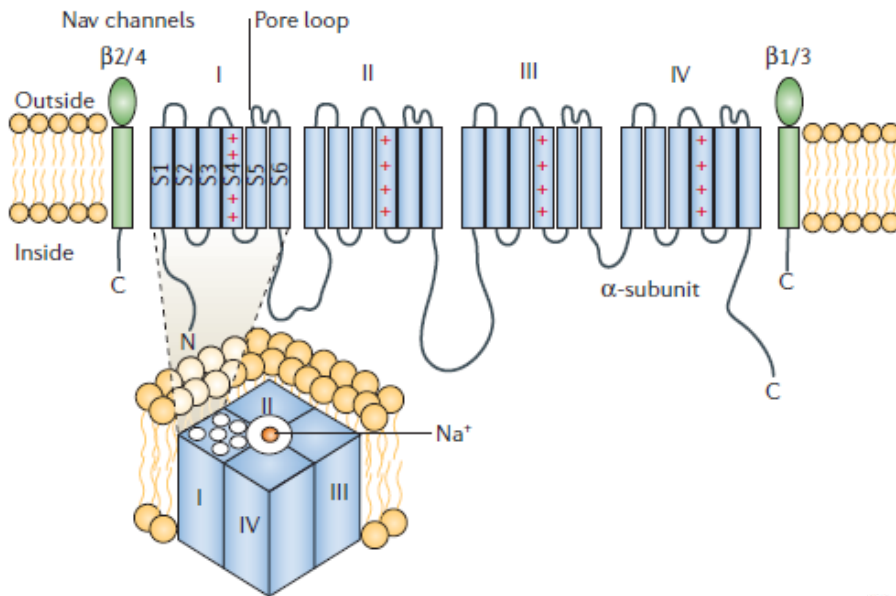


# Az AIS kialakulása

1. proximális axonban ankyrinG felhalmozódás  
( $I\kappa B\alpha$  foszforiláció - inaktiváció)

2. ankyrinG: scaffold fehérje

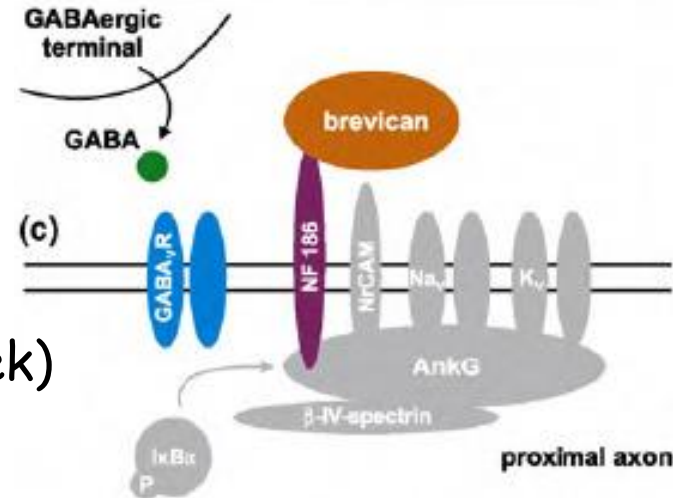
- adhéziós molekulák (NF186 [neurofascin], NrCAM)
- $\beta$ -IV spectrin  $\rightarrow$  aktin
- ioncsatornák



# Az AIS kialakulása

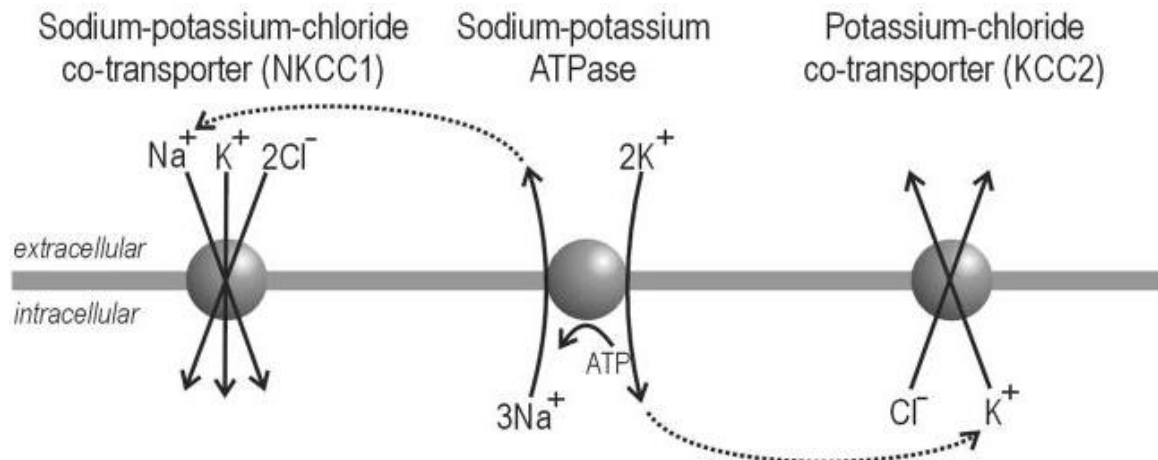
## 3. neurofascin (NF186):

- ECM szervezés (brevican, phosphocan, tenascin)
- GABAerg szinaptikus bemenet → axo-axonikus szinapszis (kandeláber sejtek)
  - általában igen hatékony gátlás
  - kortikális piramissejtek: serkentés!



Building and maintaining the axon initial segment  
Matthew S Grubb and Juan Burrone

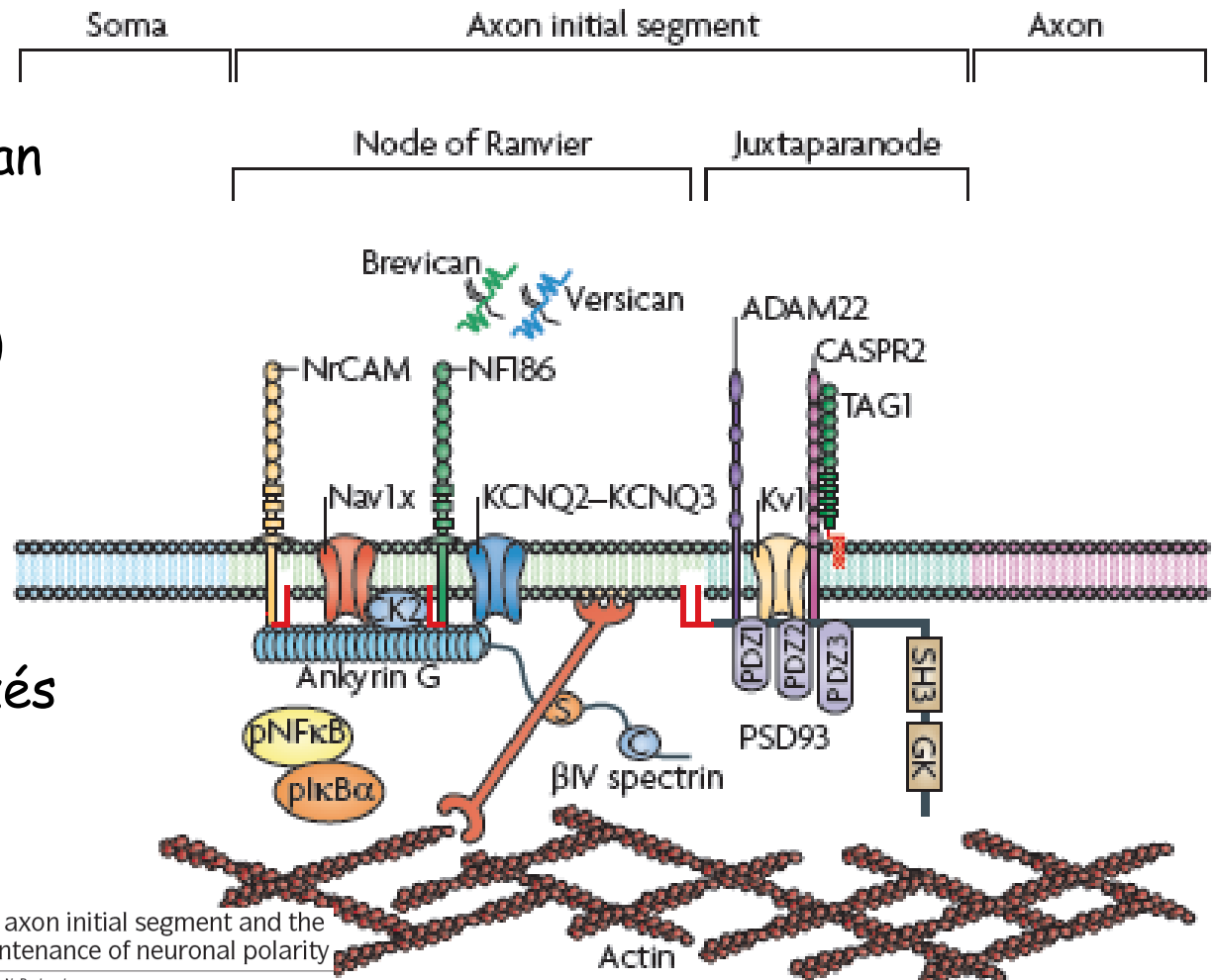
- KCC2 (*K-Cl* kotranszporter) denzitás az AIS-ben alacsony → magas  $[Cl^-]_{IC}$  →  $GABA_A$ R aktivációja depolarizáló hatású



# Az AIS (és a Ranvier-féle befűződés) szerkezete

- master scaffold: ankyrinG +  $\beta$ IV spectrin, PSD-93
- feszültség-függő  $\text{Na}^+$  és  $\text{K}^+$  csatornák [ $\text{Na}_v1.X$ ;  $\text{KCNQ2-KCNQ3}$ ,  $\text{K}_v1.X$ ]
- adhéziós molekulák: NrCAM, neurofascin, TAG1
- metalloproteáz: ADAM22
- ECM: brevican, versican
- szabályozó faktorok:
  - CK2 (*kazein kináz2*)
  - $\text{NF}\kappa\text{B}$  /  $\text{I}\kappa\text{B}\alpha$

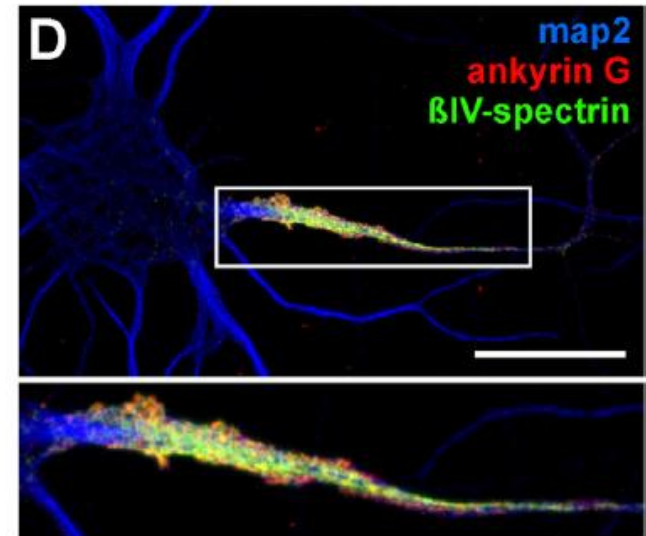
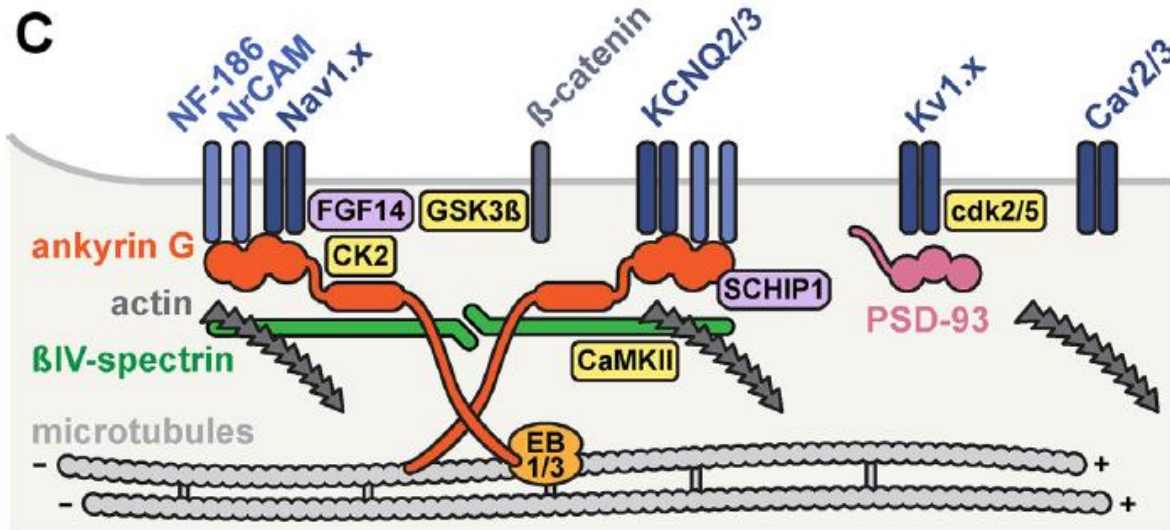
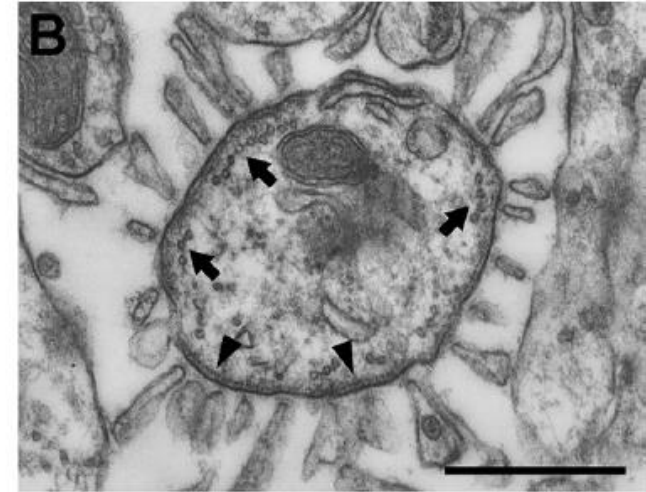
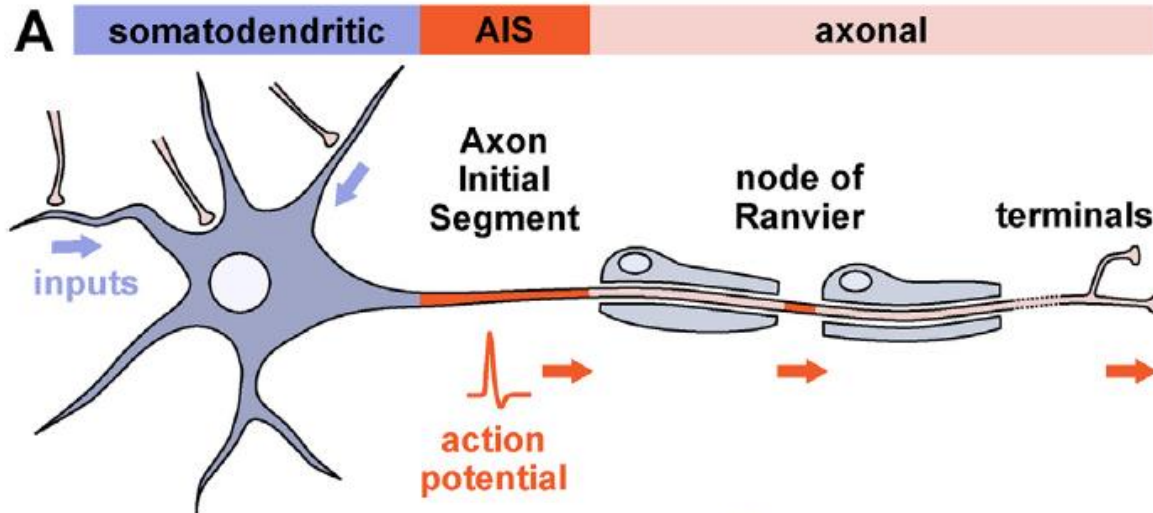
- Ranvier-befűződés: szabályozott elrendezés (ld. később)



The axon initial segment and the maintenance of neuronal polarity



# Az AIS (és a Ranvier-féle befűződés) szerkezete



No Pasaran! Role of the axon initial segment in the regulation of protein transport and the maintenance of axonal identity

Christophe Leterrier\*, Bénédicte Dargent

# Az AIS (és a Ranvier-féle befűződés) szerkezete

**Fig. 1.** The axon initial segment (AIS). (A) The neuron integrates inputs received in the somatodendritic compartment (blue). The AIS (red), located at the beginning of the axon, generates the action potential that propagates up to the terminals (light red), and is regenerated at nodes of Ranvier (red) across internodes. (B) The AIS of a Purkinje cell (transverse section, right). This electron microscopy image demonstrates two morphological features of the AIS: microtubule fascicles (arrows) and the membrane undercoat (arrowheads). Adapted with permission from Synapse Web (J. Spacek and K. Harris, PI, <http://synapses.clm.utexas.edu>). Scale bar, 0.5  $\mu\text{m}$ . (C) The AIS of a cultured rat hippocampal neuron labeled for ankG (red) and  $\beta\text{IV}$ -spectrin (green). Map2 (blue) is excluded from the axon and delineates the somatodendritic compartment. Scale bar, 20  $\mu\text{m}$ . (D) The AIS components. The main AIS scaffold is ankG (orange), linked to  $\beta\text{IV}$ -spectrin (green) that in turn binds to actin filaments (dark gray). AnkG binds to Nav1.x channels and Kv7.2/7.3 channels (dark blue), as well as adhesion proteins NF-186 and NrCAM (blue). EB1/3 proteins (light orange) link ankG to microtubules (light gray). AnkG also binds to SCHIP1 (purple). Other channels present at the AIS are Kv1.x channels linked to PSD-93 (pink), and Cav2/3 channels. Kinases are shown in yellow (see details in main text).

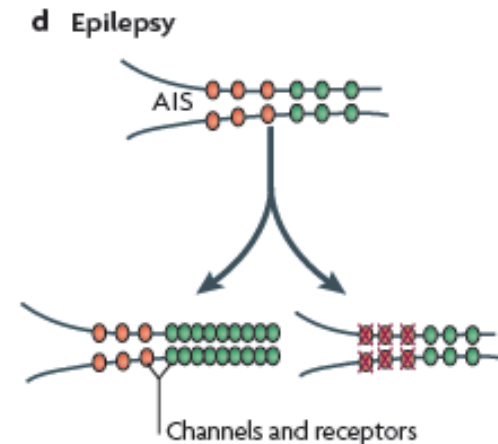
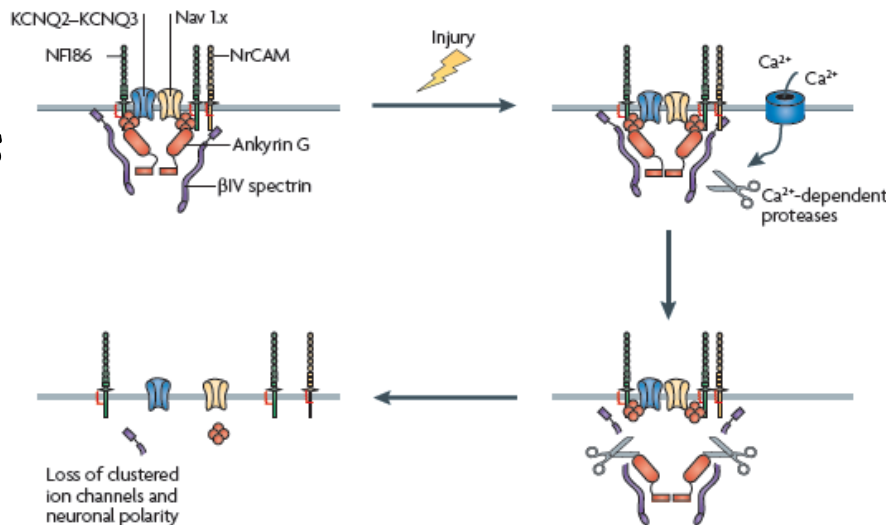
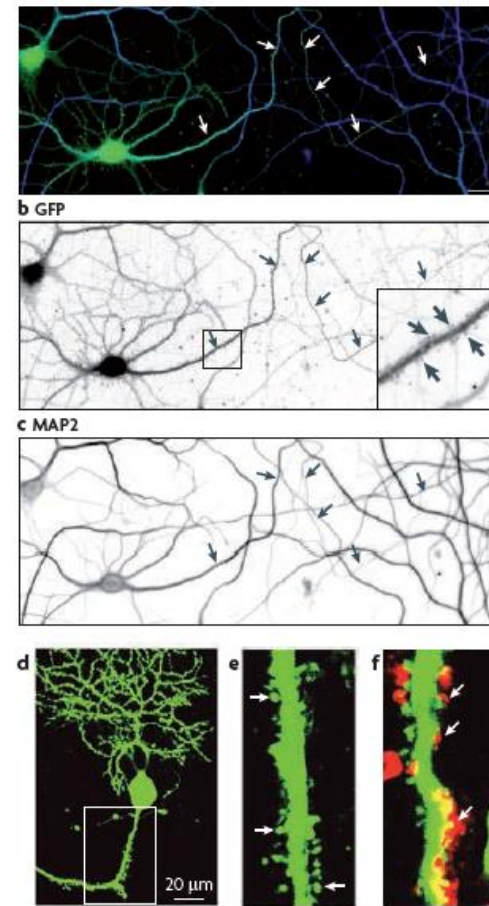
No Pasaran! Role of the axon initial segment in the regulation of protein transport and the maintenance of axonal identity

Christophe Leterrier\*, Bénédicte Dargent

# Az AIS működése és szerepe

- **plaszticitás:** térbeli pozíció az ingerlékenységet befolyásolja (disztálisabb -> magasabb küszöb)
- **ankyrinG silencing** -> „dendritizáció”
- **axonális sérülés:** az AIS épségétől (vagy a MT stabilitásától) függ a nyúlvány regenerációja
- **skizofrénia, epilepszia, MS:** aberráns AIS és/vagy receptor-eloszlás
- **ischemia:**  $Ca^{2+}$ -függő proteolízis -> AIS szétesés, disztális irányú eltolódás

ankyrinG siRNA



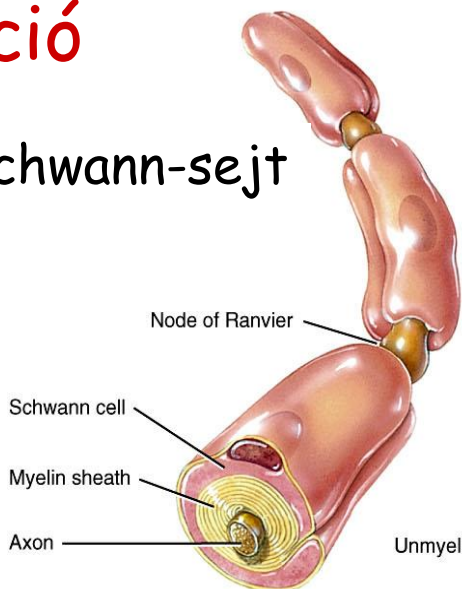
# Az AIS működése és szerepe

Figure 5 | **Nervous system injury and disease alters neuronal polarity.** **a** | Ischaemic injury (that is, stroke) disrupts the axon initial segment (AIS) cytoskeleton. The micrograph shows AIS stained for the cytoskeletal protein  $\beta$ IV spectrin (green), neurons labelled with NeuroTrace (red) and nuclei labelled with Hoechst stain (blue). The dashed line indicates the transition from the injured brain, in which the AISs are missing, to the area not exposed to ischaemia, where the AISs remain intact. The scale bars represent 50  $\mu$ m. **b** | The cascade of events leading to loss of neuronal polarity after injury. Injury increases cytoplasmic  $Ca^{2+}$ , which activates the  $Ca^{2+}$ -dependent cysteine-protease calpain. Calpain (indicated by the scissors) cleaves ankyrin G (AnkG, also known as ANK3) and  $\beta$ IV spectrin, leading to the declustering of ion channels and loss of polarity. **c** | The type and location of axonal injury determines the consequence for neuronal polarity. Axonal transection near the cell body causes a dendrite-to-axon identity switch (left panel), whereas transection far from the cell body does not (right panel). **d** | Disruption of channel and receptor density, function or location can cause or result from altered nervous system function. For example, epilepsy can increase channel densities (left) or result from altered AIS channel function (right). Demyelination leads to altered subcellular polarity of axons. Parts **a** and **b** are modified, with permission, from REF. 64 © (2009) The Society for Neuroscience.

# A mielinizáció

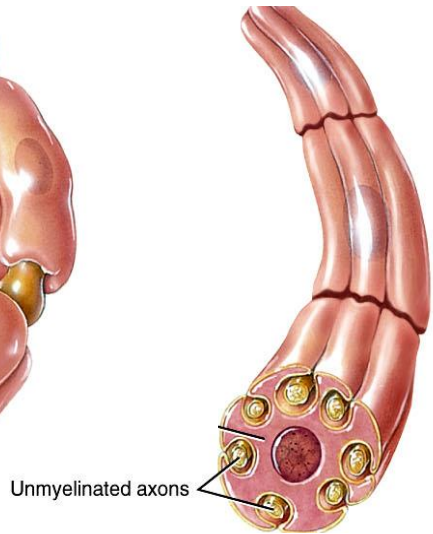
perifériás idegrendszer

Schwann-sejt

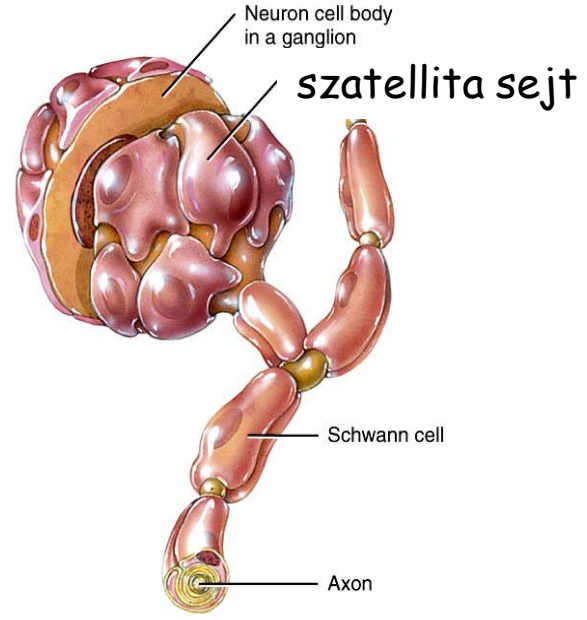


velőshüvelyű axon

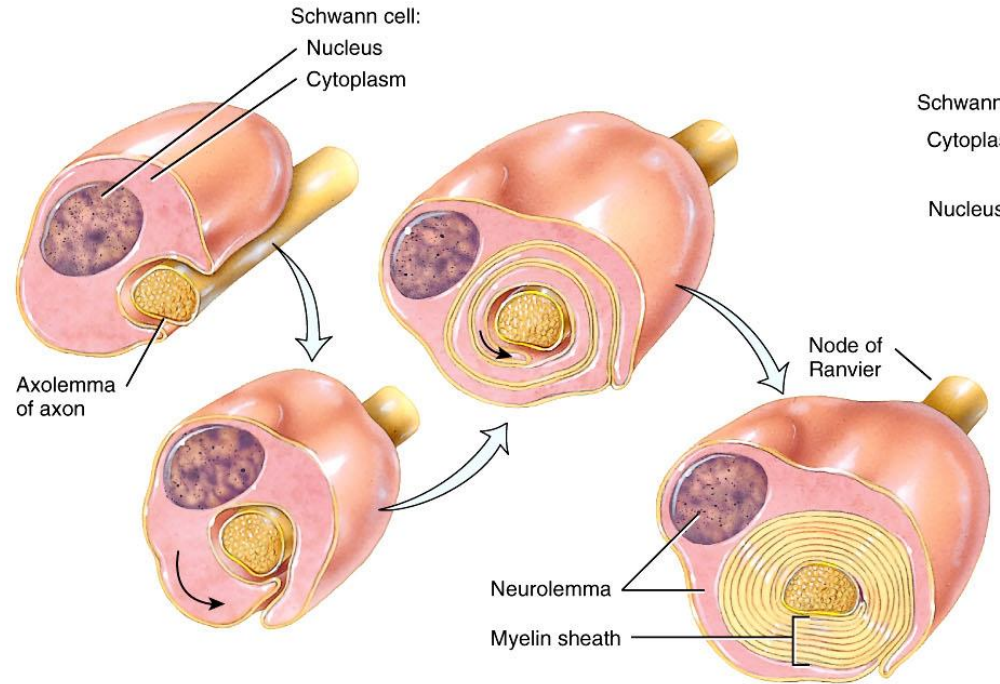
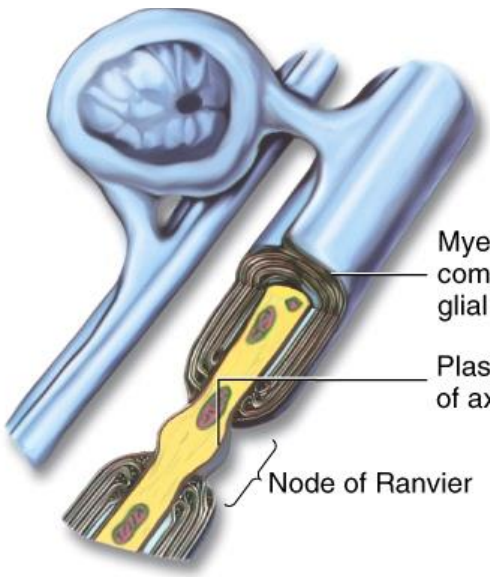
Remak-sejt



több, nem mielinált axon



központi idegrendszer  
oligodendroglia

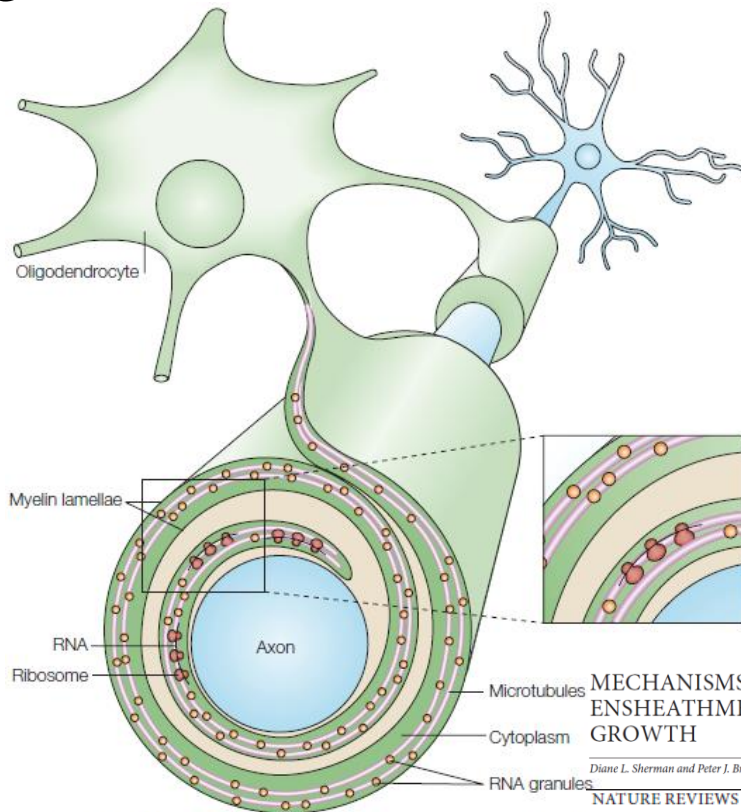


Schwann cell  
Cytoplasm  
Nucleus

# A mielin hüvely kialakulása

- mielin bázikus fehérje szintézis a periférián -> lokalizált RNS szemcsék
  - MT-hez rögzítve, ill. szállítva
  - oligodendroglia: „feltekert” MT, rés-kapcsolatok
  - Schwann sejt: mikrovillusok, Cajal-sávok (disztrofin-periaxin komplex)

CNS

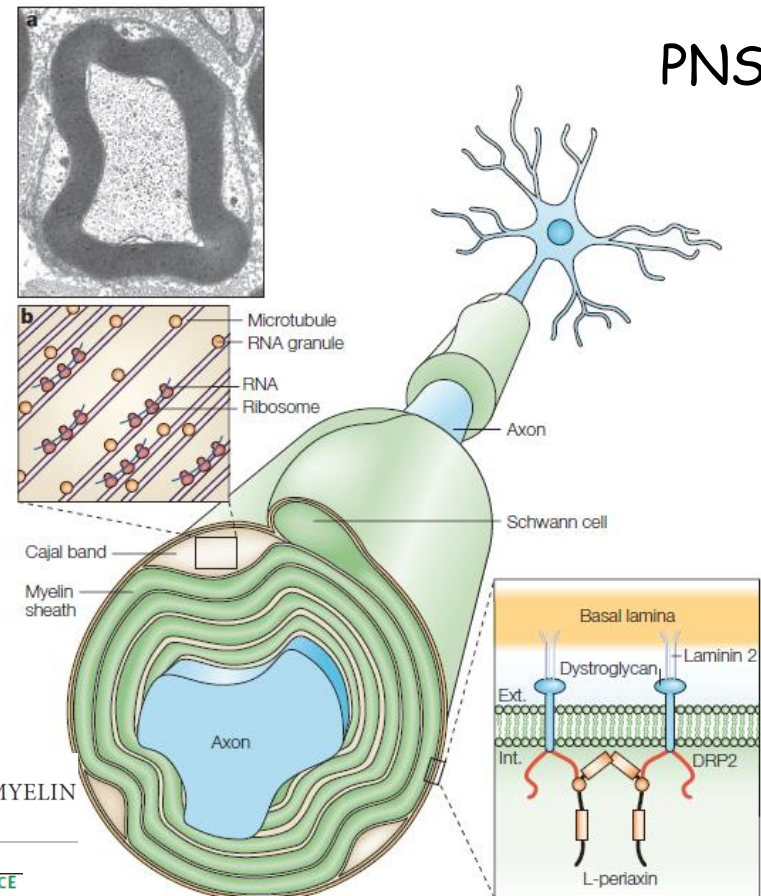


MECHANISMS OF AXON  
ENSWHEATHMENT AND MYELIN  
GROWTH

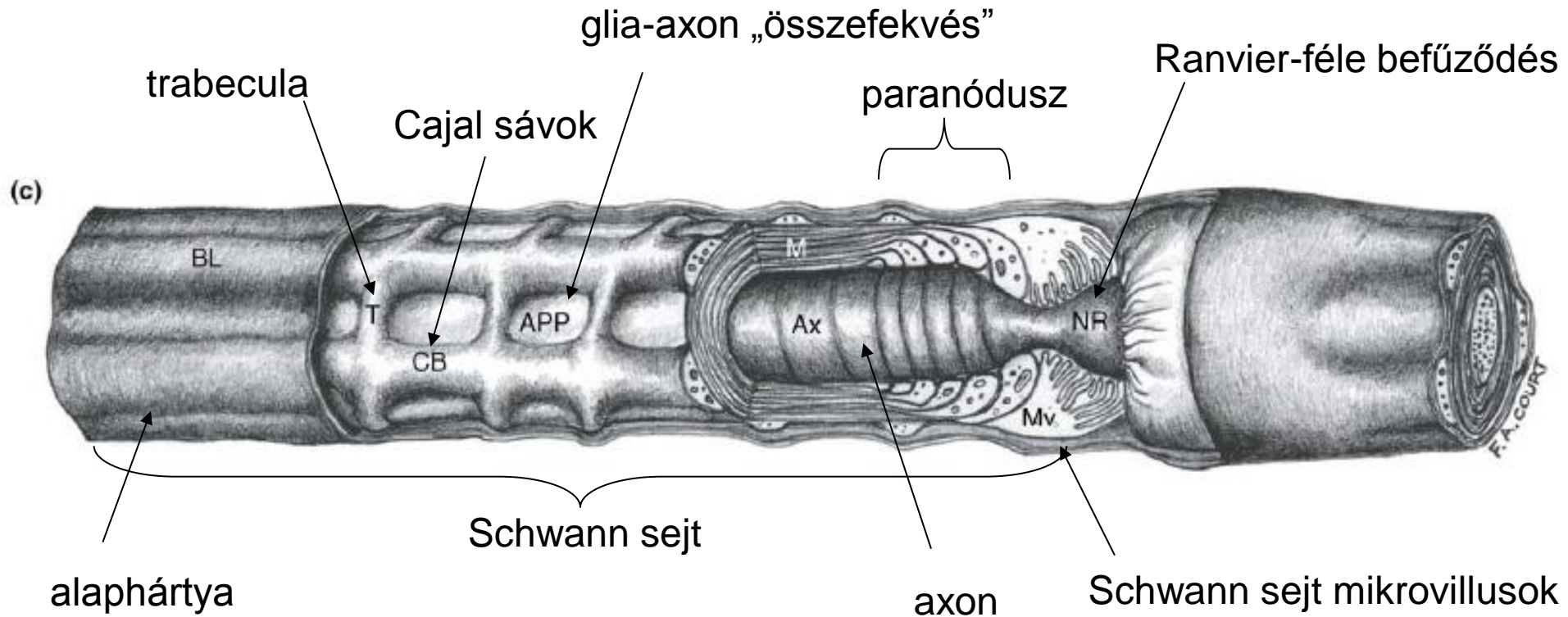
Diane L. Sherman and Peter J. Brophy

NATURE REVIEWS | NEUROSCIENCE

PNS



# Mielin hüvely a PNS-ben

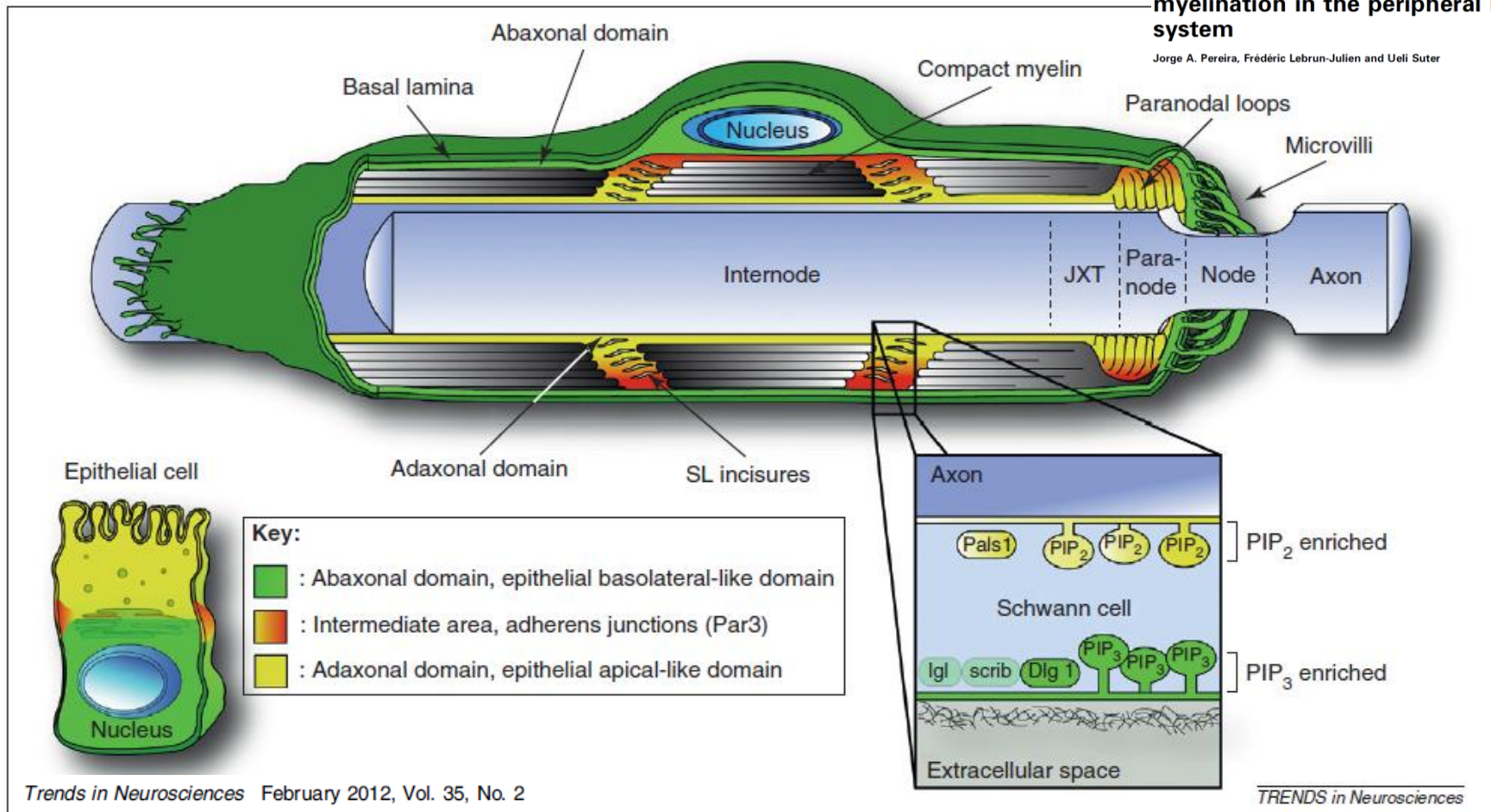


Schematic drawing with cut away sections displays the architecture of the myelinated fiber. Abbreviations: App: apposition; Ax: axon; BL: basal lamina; CB: Cajal Band; M: myelin; Mv: microvilli; NR: node of Ranvier; T: trabecula.

# Mielin hüvely a PNS-ben

Molecular mechanisms regulating myelination in the peripheral nervous system

Jorge A. Pereira, Frédéric Lebrun-Julien and Ueli Suter

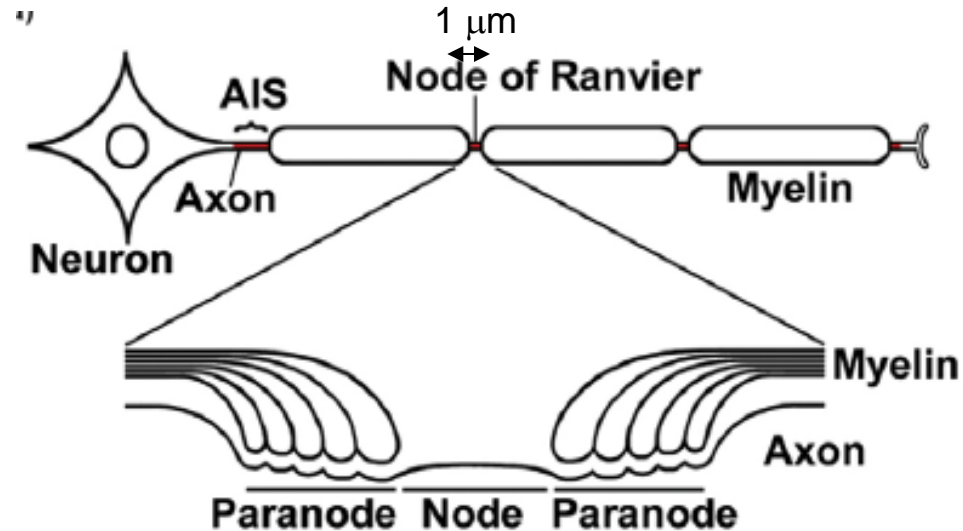
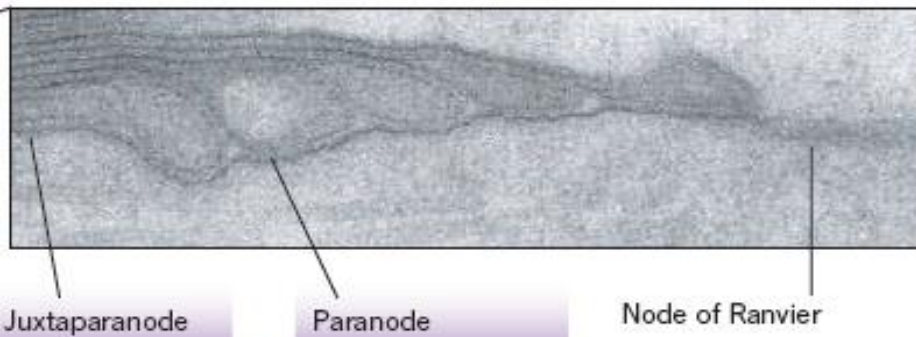


**Figure 2.** Polarized structure of the adult myelinating SC. Myelinating SCs cover a segment of the axon, designated the internode, and organize their subcellular domains in a polarized fashion, both in the longitudinal and radial axes. Longitudinally, SCs display the nucleus at the center. At the edge of the internode, cytoplasm-filled SC paranodal loops tether the internode to the axon and define the juxtapanodal region (JXT). SC microvilli project into the nodal area. Radial polarity of SCs is also striking, with the nucleus being localized in the outermost wrap of the myelin sheath (abaxonal domain), followed by compact myelin and the innermost wrap facing the axon (adaxonal domain). Cytoplasm-filled spiral-shaped channels, Schmidt-Lanterman (SL) incisures, connect the adaxonal and abaxonal cytoplasm. The abaxonal domain is in tight contact with the basal lamina, a thin layer of highly organized ECM components synthesized by SCs, whereas the adaxonal domain is in close contact with the axolemma. This radial polarity organization shows similarities to epithelial cell polarization with an apical and a basolateral domain. Adult myelinating SCs show distributions of intrinsic polarity-regulatory proteins similar to those of epithelial cells, with Dlg1 being enriched in the abaxonal domain (basolateral-like), Pals1 concentrated in the adaxonal domain, SL incisures and paranodal loops (apical-like), and Par3 being localized to adherens junctions in outer regions of paranodal loops and SL incisures (between both domains). The asymmetric distribution of polarity proteins is coupled to a polar distribution of phosphoinositides. In mature myelinating SCs, PIP<sub>2</sub> is enriched in the adaxonal domain and PIP<sub>3</sub> is concentrated at the abaxonal domain. These enriched distributions of polarity proteins and PIPs relate to adult myelinating SCs. Note that when SC polarity is established and further enhanced during the myelination process, localization of these proteins and lipids is dynamic with different effects on signaling, as exemplified by Par3 recruitment to the SC-axon interphase at myelination initiation (Figure 3).



# A mielinhüvely és a Ranvier-féle befűződés

- „szigetelés”: membrán ellenállás  $\uparrow$ , kapacitancia  $\downarrow$  -> gyorsabb konduktancia ( $> 100\times$ )
- energia-felhasználás  $\downarrow$  ->  $< 0.1\%$  membrán-felszínen van Na/K ATPáz
- szakaszok:
  - nódusz  $\sim$  AIS
  - paranódusz: mielin-hüvely széle; legnagyobb adherens komplex
  - juxtapanódusz: speciális barrier (5-15  $\mu\text{m}$ ;  $K_v + \text{Caspr2}$ , TAG-1)
  - internódusz: mielin burok, közeli axon-glia oppozíció

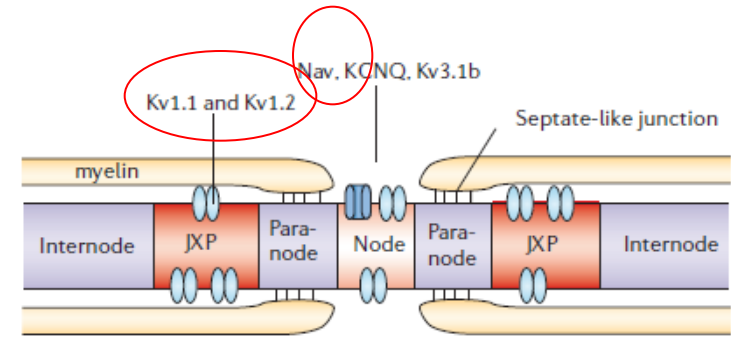


# A mielinhüvely és a Ranvier-féle befűződés

The distribution and targeting of neuronal voltage-gated ion channels  
 Helen C. Lai\* and Lily Y. Jan\*

- ioncsatornák szabályozott eloszlása az axonon

- $Na_V$ : akciós potenciál terjedés a nóduszban
- $K_V$ : hiperpolarizáció → akciós potenciál gátlása a juxtaparanodális területen



- adhéziós rendszerek

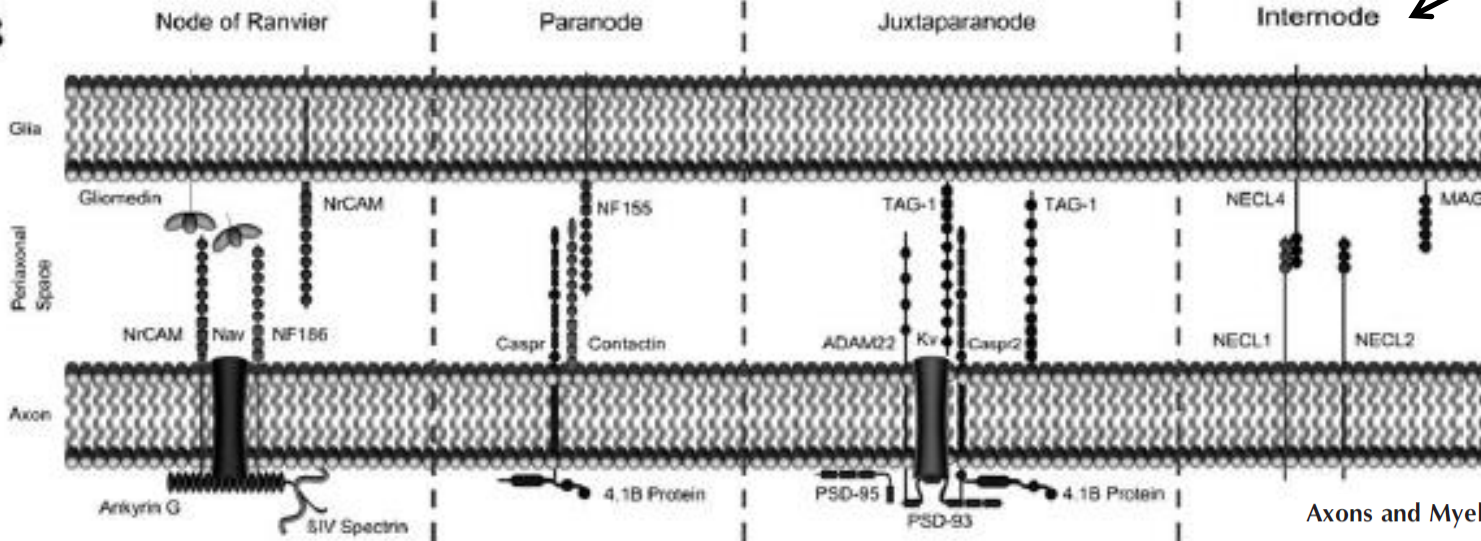
$K_V$  és  $Na_V$  csatornák elválasztása; „válaszfalak”

$K_V$  csatornák lokalizációja

PNS: mikrobolyhok  
 CNS: NG2 nyúlvány

mielin rögzítése, növekedése

**B**



Axons and Myelinating Glia: An Intimate Contact

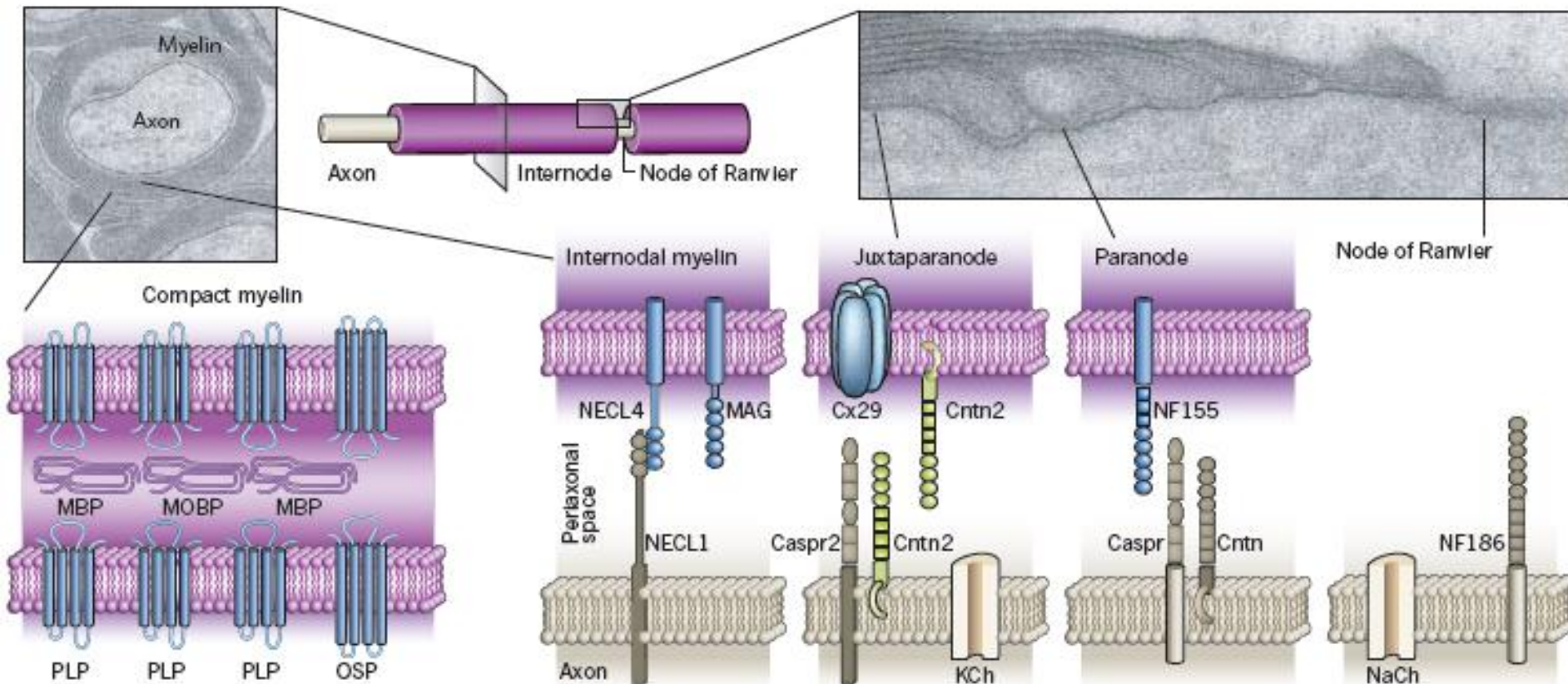
Lida Zoupi, Maria Savvaki and Domna Karageorgis

# A mielinhüvely felépítése

- lipid tutajok: magas koleszterin, galaktolipid tartalom

Myelination and support of axonal integrity by glia

Klaus-Armin Nave<sup>1</sup>

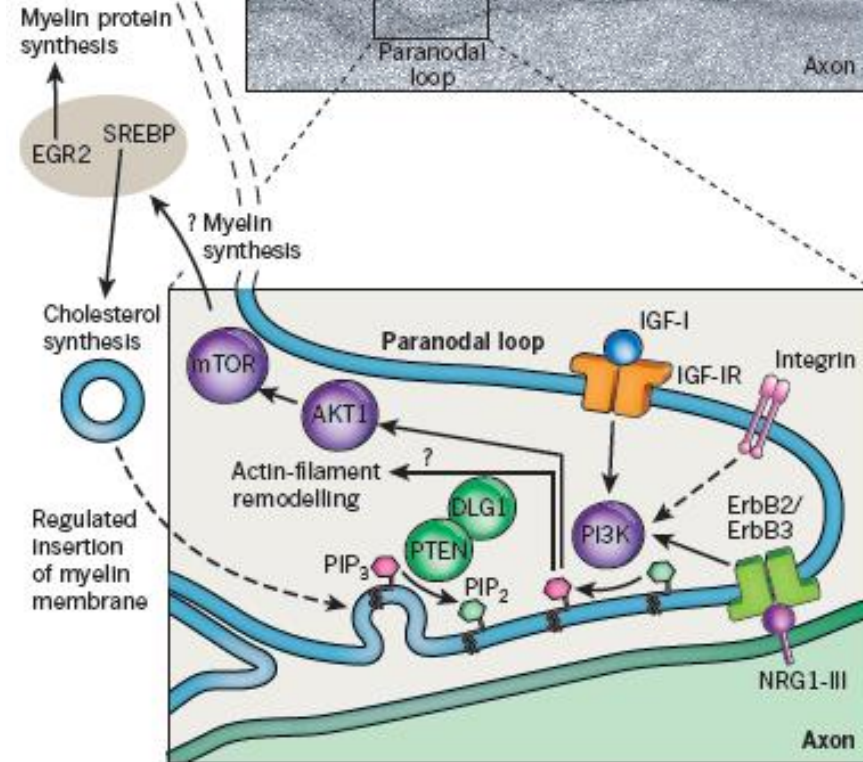
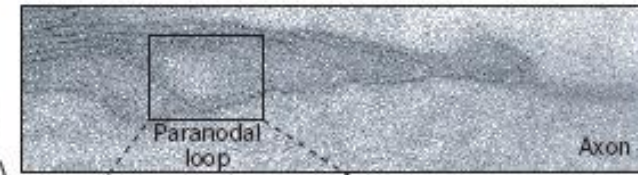


**Caspr**: contactin-associated protein; **Cntn**: contactin (Cntn2 is also known as Tag1); **Cx29**: connexin 29 kDa; **KCh**: fast potassium channels; **MAG**: myelin-associated glycoprotein; **MBP**: myelin basic protein; **MOBP**: myelin oligodendrocyte basic protein; **NaCh**: voltage-gated sodium channels; **NECL**: nectin-like protein/synCAM; **NF155/186**: neurofascin 155 kDa/186 kDa; **OSP**: oligodendrocyte-specific protein; **PLP**: proteolipid protein

# A mielin hüvely kialakulása

Myelination and support of axonal integrity by glia

Klaus-Armin Nave<sup>1</sup>



**NRG1-III**: neuregulin-1 type III; **ErbB**: epidermal growth factor receptor; **PI3K**: phosphatidylinositol-3-kinase; **DLG1**: mammalian discs large homolog 1; **PTEN**: phosphatase and tensin homologue; **BACE**:  $\beta$ -site amyloid precursor protein-cleaving enzyme 1; **EGR2**: early growth response protein 2 (also known as Krox20); **ErbB2/ ErbB3**: heterodimeric NRG1 receptor tyrosine kinases; **IGF-1R**: IGF-1 receptor; **PIP2**: phosphatidylinositol-4,5-bisphosphate; **SREBP**: sterol regulatory element binding protein.

- g arány: axon átmérő / (axon+mielin) átmérő;  
~0,6 - 0,7 (remielinizáció után kisebb)

## ➤ PNS-ben axonális hatásra:

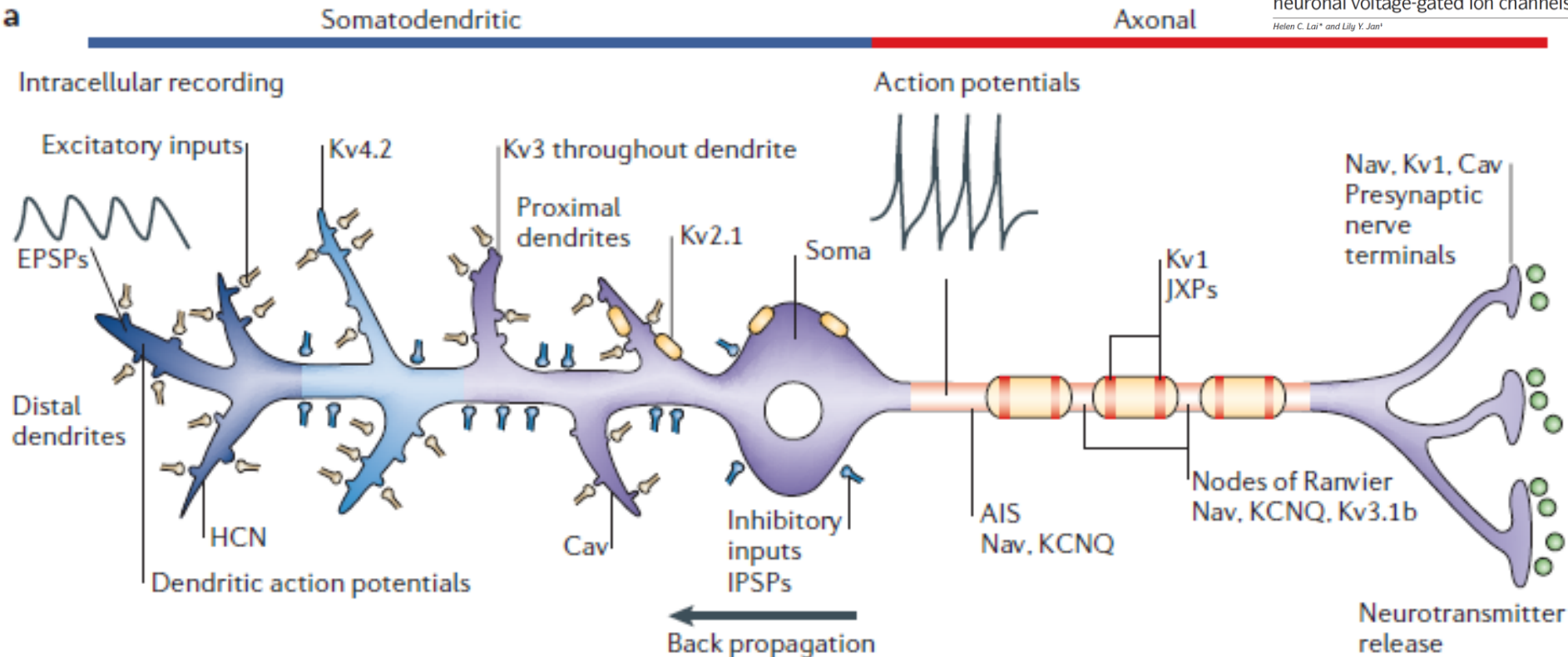
- neuregulin-1 / ErbB  $\rightarrow$  mielin szintézis  $\uparrow$ , aktin polimerizáció, átalakulás a Schwann sejtben
- axonális szignál nélkül nincs mielinizáció: induktív hatás

## ➤ CNS:

- NRG/ErB vagy elektromos aktivitás hiányában is, fixált axon körül is  $\rightarrow$  axonális szignál nem induktív
- számos szignál: BDNF, CNTF, PDGF, LIF...
- axon membrán: permisszív hatás

# Az ingerlékeny membrán mozaikossága

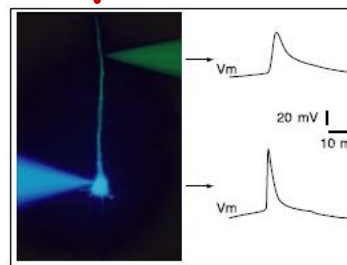
The distribution and targeting of neuronal voltage-gated ion channels  
Helen C. La\* and Lily Y. Jan\*



**General localization of voltage-gated ion channels in a model neuron.** In general, **Nav channels** are found in the axon initial segment (AIS), nodes of Ranvier and presynaptic terminals. Voltage-gated potassium **Kv1 channels** are found at the juxtaparanodes (JXPs) in adult myelinated axons and presynaptic terminals. The Kv channel **KCNQ** is found at the AIS and nodes of Ranvier, and **Kv3.1b channels** are also found at the nodes of Ranvier. Canonically, excitatory and inhibitory inputs (EPSPs and IPSPs — excitatory and inhibitory postsynaptic potentials; yellow and blue presynaptic nerve terminals, respectively) from the somatodendritic region spread passively to the AIS where action potentials are generated by depolarization, and travel by saltatory conduction to the presynaptic nerve terminals to activate voltage-gated calcium (**Cav**) channels that increase intracellular calcium levels, thereby triggering neurotransmitter release. Hyperpolarization-activated cyclic-nucleotide-gated (**HCN**) channels have a gradient distribution that increases in density from the soma to the distal dendrites (dark blue shading). **Kv2.1 channels** are found in clusters on the soma and proximal dendrites (light yellow ovals). **Kv3 channels** are found throughout the dendrite. **Kv4.2 channels** are located more prominently on distal dendrites (light blue shading). Kv channels in the dendrites contribute to controlling back propagation. Strong enough inputs in the dendritic region can generate dendritic action potentials. **Dendritic Cav channels** increase in density toward the proximal dendrites and the soma.

# A backpropagating akciós potenciál (BAC)

- az AIS-en kialakuló AP a dendritekre is visszaterjed; függ:



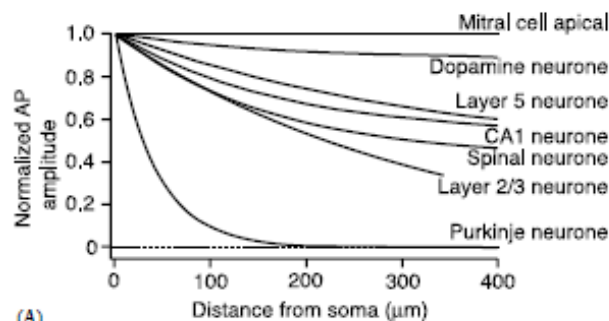
Backpropagating action potentials in neurones: measurement, mechanisms and potential functions

Jack Waters<sup>\*1</sup>, Andreas Schaefer<sup>1</sup>, Bert Sakmann

Progress in Biophysics and Molecular Biology 87 (2005) 145–170

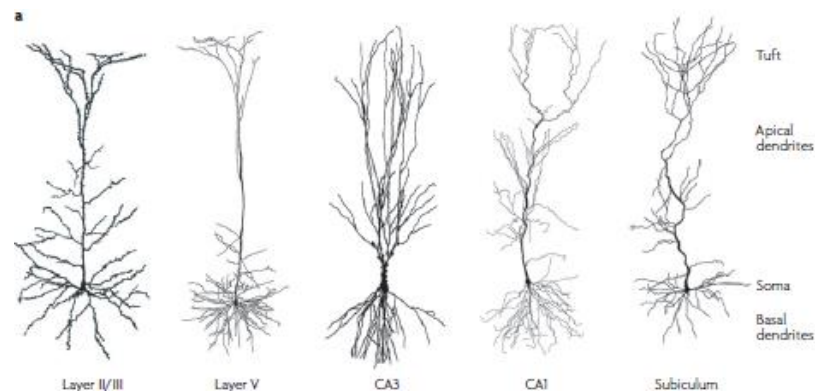
- sejttesttől mért távolság

- sejt típus



(A)

- dendritfa alakja



- ioncsatorna eloszlás és típus

- HCN [hiperpolarizáció aktiválta kation csatorna]: disztális dendritek felé növekvő sűrűségben
- fesz. függő  $\text{Na}^+$ ,  $\text{Ca}^{2+}$  és  $\text{K}^+$  csatornák

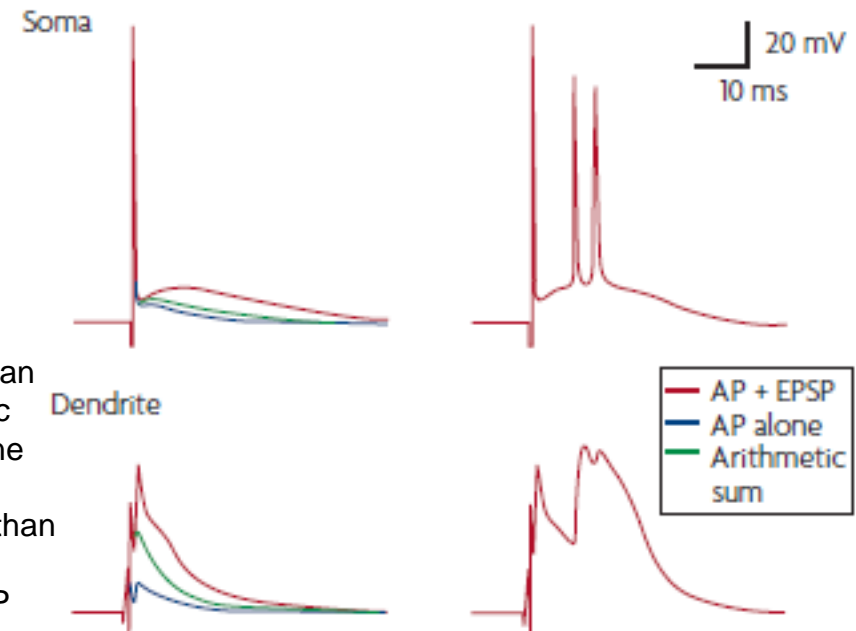
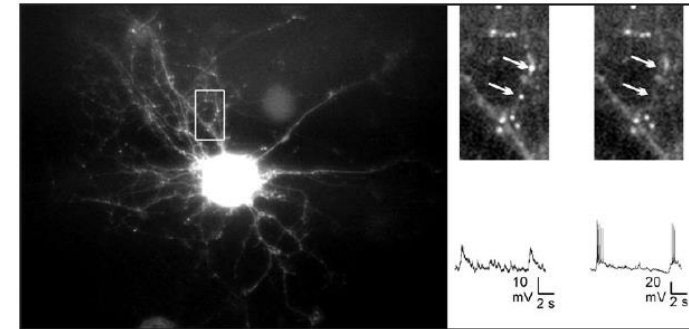
# A backpropagating akciós potenciál (BAC)

• szerepe van:

- a neuron output-járól retrográd szignál: koincidencia-detekció; pl. LTP
- dendritikus  $[Ca^{2+}]_{IC}$  emelése: pl. BDNF release
- szinaptikus integráció befolyásolása

## Back-propagating action potential

A key contributor in activity-dependent dendritic release of BDNF  
[Communicative & Integrative Biology 1:2, 153-155;]



Amplification of backpropagating action potentials by dendritic EPSPs can lead to bursting. The two lefthand plots show somatic (top) and dendritic (bottom) responses to an antidromic action potential (AP) activated alone (blue trace) or in combination with a dendritic EPSP-like response to dendritic current injection. The combined response (red trace) is larger than the linear sum of the action potential and the EPSP separately (green trace). The two right-hand plots are from another trial in which the EPSP triggered a second action potential that backpropagated even more effectively, leading to a large dendritic spike that triggered another action potential and hence a burst.

Pyramidal neurons: dendritic structure and synaptic integration

Nelson Spruston

Ábrák



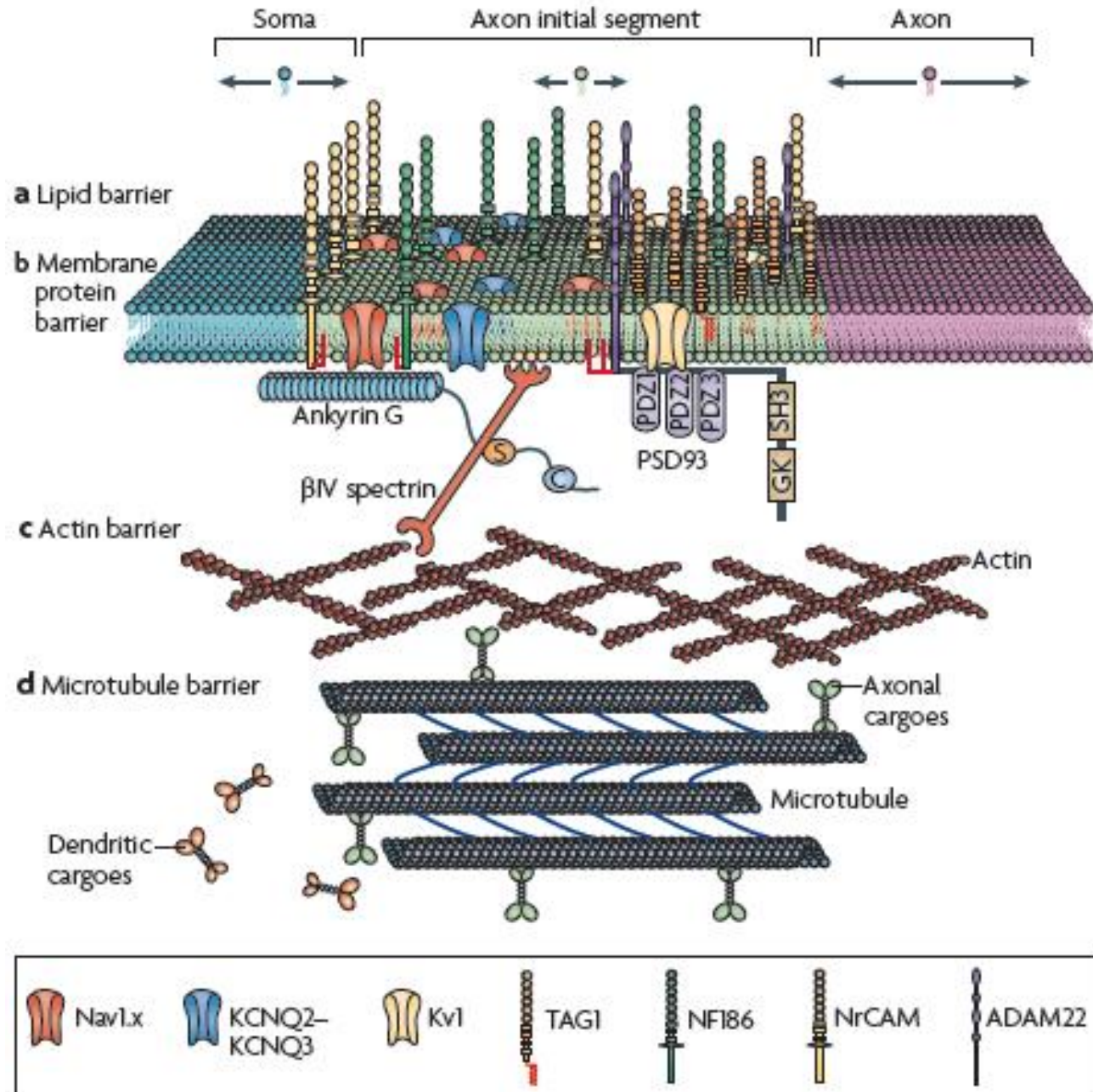
# Az AIS

The axon initial segment and the maintenance of neuronal polarity

Matthew N. Rasband

## Molecular substrates of the axon initial segment barrier.

Four potential mechanisms have been proposed to contribute to the axon initial segment (AIS) barrier. **a** | The lipid composition of the AIS plasma membrane may directly influence mobility and diffusion rates in the AIS. The lipid composition can be directly regulated by post-translational modifications of membrane and cytoplasmic proteins or by phospholipid–cytoskeleton interactions. **b** | The high density of membrane proteins creates a membrane diffusion barrier that limits the lateral mobility of other transmembrane proteins and lipids at the AIS. The high density of these proteins is established through binding to ankyrin G. **c** | Actin filaments can contribute to the maintenance of neuronal polarity and the AIS barrier by limiting the entry of cytoplasmic proteins into the axon and by contributing to the maintenance of protein density at the AIS membrane through interactions with  $\beta$ IV spectrin. **d** | Microtubule fascicles with unique cross-bridges at the AIS allow axonal cargoes but not dendritic cargoes to enter the axon, thus contributing to the maintenance of neuronal polarity. ADAM22, a disintegrin and metalloproteinase domain-containing protein 22; GK, guanylate kinase; NF186, neurofascin 186; NrCAM, neuronal cell adhesion molecule; PDZ, PSD95/discs large/zonula occludens; PSD93, postsynaptic density protein 93; SH3, SRC homology 3; TAG1, transient axonal glycoprotein 1 (also known as contactin 2).



# Az AIS és a Ranvier-féle befűződés szerkezete

The axon initial segment and the maintenance of neuronal polarity

Matthew N. Rasband

**The axon initial segment and nodes of ranvier are prototypical examples of subcellular polarity.**

a | A cultured hippocampal neuron with high densities of ankyrin G (AnkG, also known as ANK3) (green) and neurofascin 186 (NF186) (the overlap between AnkG and NF186 is shown in yellow) at the axon initial segment (AIS).

Microtubule-associated protein 2 (MAP2) (blue) is confined to the somatodendritic domain. The scale bar represents 20  $\mu\text{m}$ . b | Nodes of Ranvier show a highly polarized organization that includes nodal  $\text{Na}^+$  channels (green), paranodal contactin-associated protein (Caspr) (red) and juxtapanodal  $\text{K}^+$  channels (blue). The scale bar represents 5  $\mu\text{m}$ .

c | The molecular organization of the AIS, which has many features in common with nodes of Ranvier and juxtapanodes. These domains are comprised of ion channels (Nav1.x, KCNQ2–KCNQ3 and Kv1.x), cell adhesion molecules (neuronal cell adhesion molecule (NrcAM), neurofascin 186 (NF186), a disintegrin and metalloproteinase domain-containing protein 22 (ADAM22), transient axonal glycoprotein 1 (TAG1, also known as contactin 2) and CASPR2), extracellular matrix molecules (brevican and versican), cytoskeletal scaffolds (AnkG,  $\beta\text{IV}$  spectrin and postsynaptic density protein 93 (PSD93)) and other signalling proteins, some with unknown roles (casein kinase II (CK2), phosphorylated nuclear factor- $\kappa\text{B}$  (pNF $\kappa\text{B}$ ) and phosphorylated inhibitor of  $\kappa\text{B}$  (plk $\beta\alpha$ )). Intriguingly, both nodal and juxtapanodal proteins are located at the AIS, but they occupy mutually exclusive domains in myelinated axons. GK, guanylate kinase; PDZ, PSD95/discs large/zonula occludens; SH3, SRC homology 3.

