

...the growth cone may be regarded as a sort of club or battering ram, endowed with exquisite chemical sensitivity, with rapid ameboid movements, and with certain impulsive force, thanks to which it is able to proceed forward and overcome obstacles met in its way, forcing cellular interstices until it arrives at its destination.

Santiago Ramón y Cajal (1909)

Histologie du système nerveux de l'homme
et des vertébrés, p.599

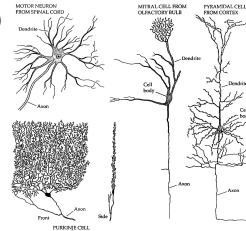
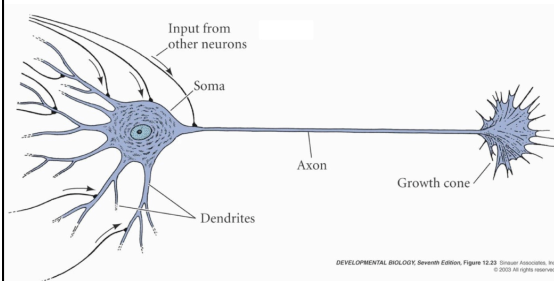
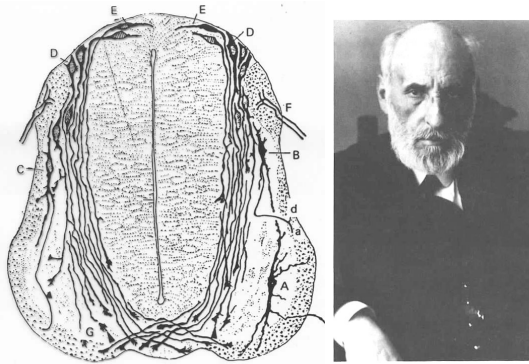
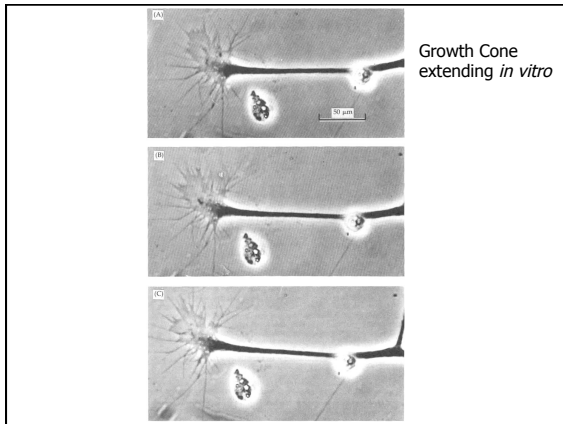


Figure 12.23 (modified) Diagram of a Motor Neuron



Growth Cones discovered (Ramón y Cajal, 1890)





Mechanisms of Growth Cone Guidance

Contact-mediated (short-range)

- requiring direct cell-cell or cell-substrate contact

Contact attraction

Contact repulsion

Diffusible (' long-range')

Chemoattraction

Chemorepulsion

(Tessier-Lavigne & Goodman, 1996 "Molecular Biology of Axon Guidance")

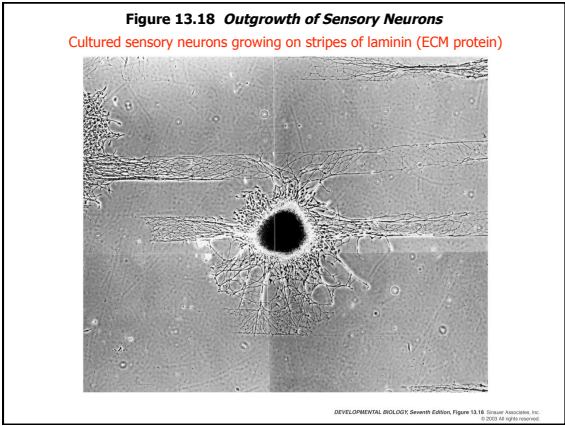
Mechanisms of Growth Cone Guidance

Contact mediated (short-range) - requiring direct cell-cell or cell-substrate contact

Contact attraction - mediated by two classes of proteins

Direct Cell-Cell Adhesion: Cell Adhesion Molecules (CAMs)

Cell - Substrate Adhesion: Extracellular (ECM) Matrix proteins
and their cellular receptors



Mechanisms of Growth Cone Guidance

Cell Adhesion Molecules (CAMs)

Major classes:

Calcium-dependent Adhesion proteins: Cadherins
 e.g. N-cadherin, E-cadherin, P-cadherin

Immunoglobulin Superfamily Adhesion proteins:
Ig-CAMs
 e.g. NCAM, L1, Fasciclin II (FasII), etc.

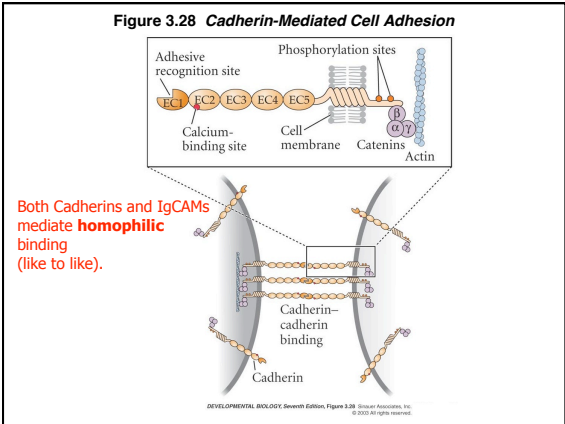
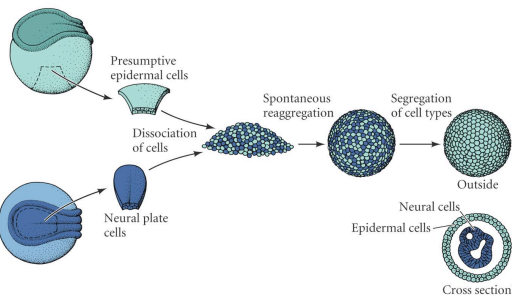


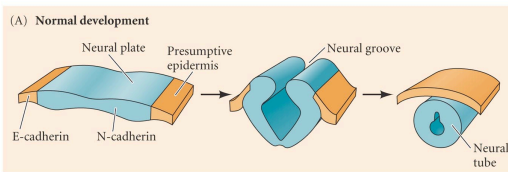
Figure 3.23 Reaggregation of Cells From Amphibian Neurulae



Cadherins mediate tissue-specific adhesion

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Fig 12.6 Expression of N- and E-cadherin Adhesion Proteins During Neurulation

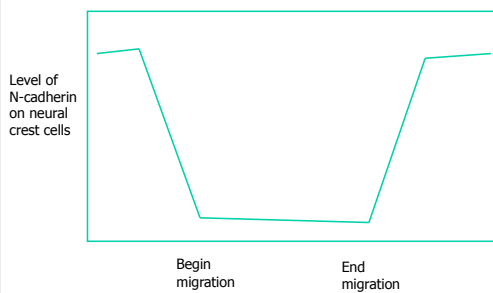


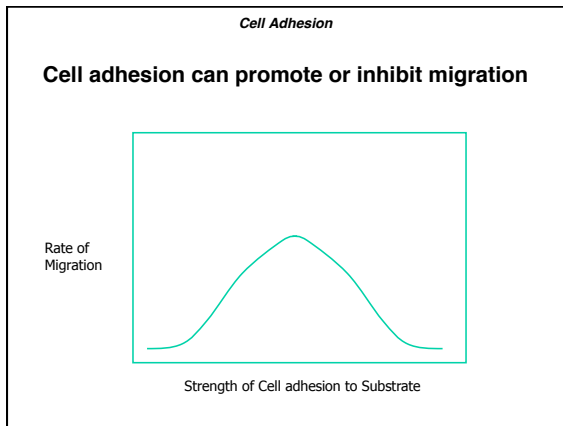
Cadherins aid separation of epidermal & neural ectoderm

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N-cadherin plays a role in neural crest cell migration

N-Cadherin level is high prior to neural crest cell migration, low during migration, and high again after migration ends.





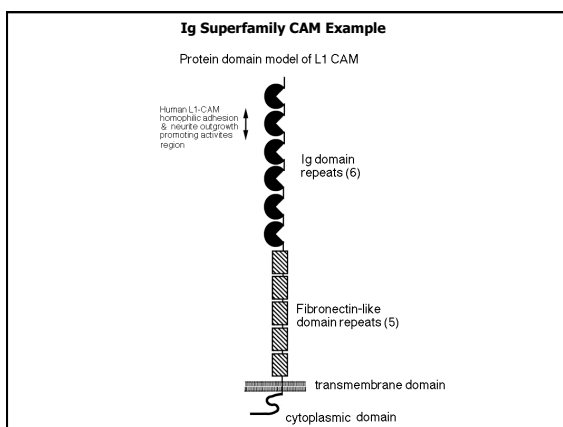
Mechanisms of Growth Cone Guidance

Cell Adhesion Molecules (CAMs)

Immunoglobulin Superfamily Adhesion proteins: Ig-CAMs
 NCAM, L1, Fasciclin II, etc.

NCAM - more general nervous system "glue"

- used to direct retinal growth cone outgrowth (1 function)
- two different forms based on glycosylation:
 - high SA (lower adhesion)
 - low SA (higher adhesion)
 - [SA = sialic acid]
- high SA found during growth, low after reaching target



Mechanisms of Growth Cone Guidance

Cell Adhesion Molecules (CAMs)

Immunoglobulin Superfamily Adhesion proteins: Ig-CAMs
NCAM, L1, Fasciclin II, etc.

L1 - more specific vertebrate CNS CAM
- found in limited number of tracts, regions of brain & s.c.

Human mutations cause neurological disorders such as
X-linked hydrocephalus, MASA syndrome

L1 disruption by ethanol may cause part of Fetal Alcohol Syndrome birth defects

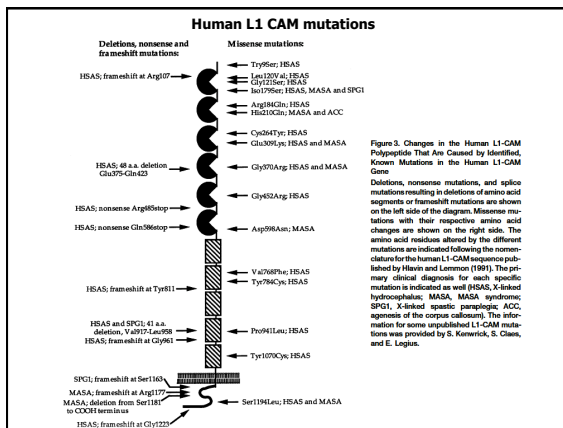


Figure 21.13(2) Possible Mechanisms Producing Fetal Alcohol Syndrome (FAS)

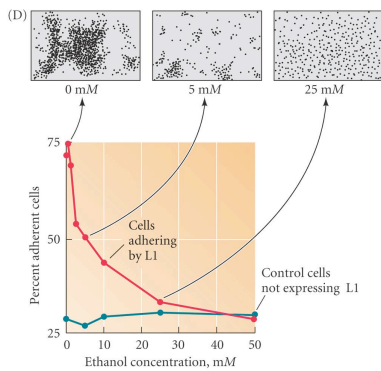
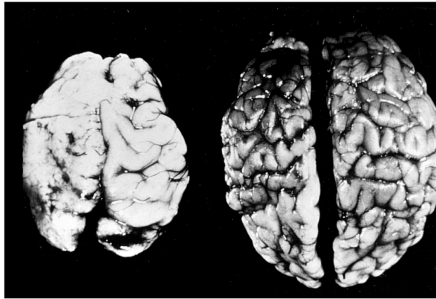
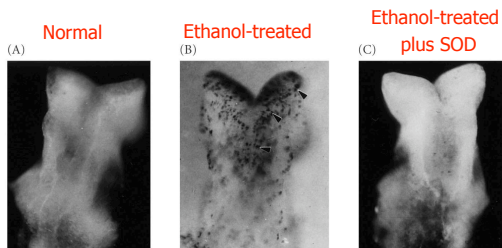


Figure 21.12 Comparison of a Brain from an Infant with Fetal Alcohol Syndrome (FAS, Left) With a Brain From a Normal Infant of the Same Age (Right)



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Figure 21.13(1) Possible Mechanisms Producing Fetal Alcohol Syndrome (FAS)



**Nile-Blue stained dying cells
in 9d mouse embryo brain**

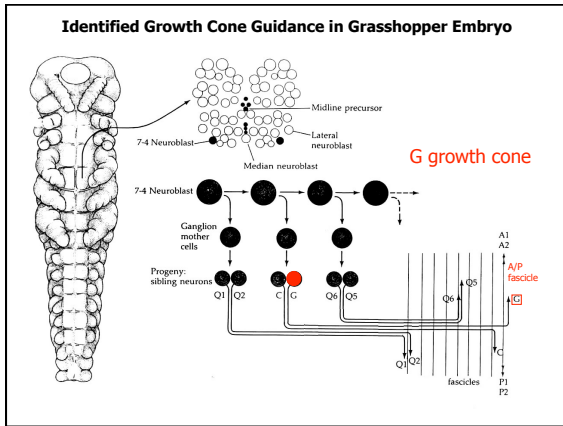
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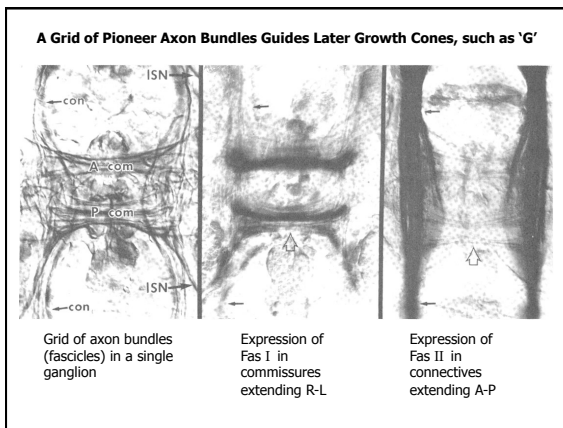
Mechanisms of Growth Cone Guidance

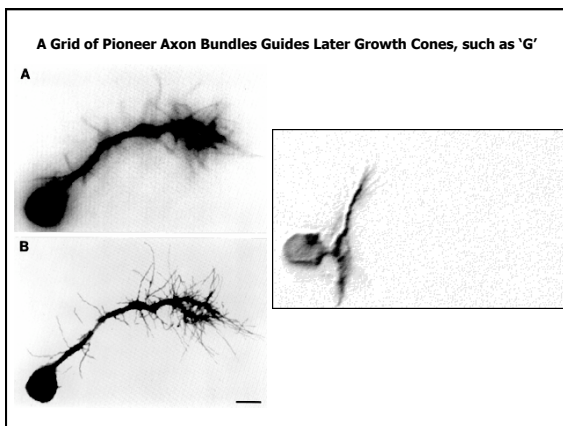
Contact attraction by CAMs

The "Labeled Pathways" Hypothesis -

Most growth cones are guided along axons already laid down (by "pioneer neurons") - each with a unique molecular marker or set of markers







A Grid of Pioneer Axon Bundles Guides Later Growth Cones, such as 'G'

Experiments:

Kill cells that make A/P bundle:
G growth cone guidance disrupted

Block function of Fas II (found on A/P bundle):
G growth cone guidance disrupted

Mechanisms of Growth Cone Guidance

Contact attraction by CAMs

Intermediate cell recognition by pioneer neurons:
"guidepost cells" in grasshopper limb bud

Pioneer Neurons in the Grasshopper Limb Bud

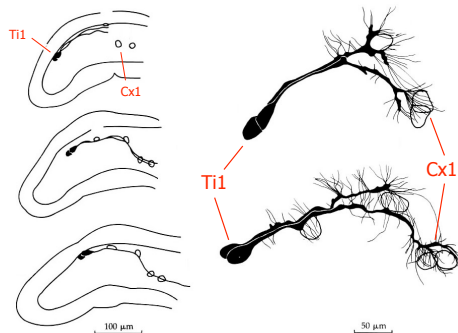
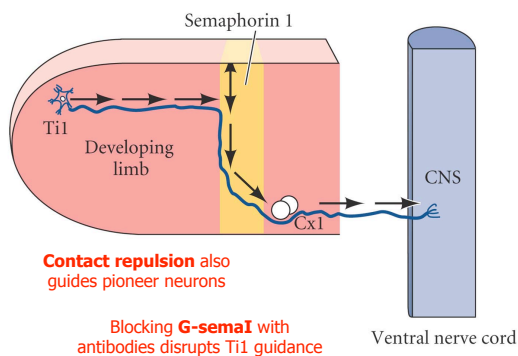
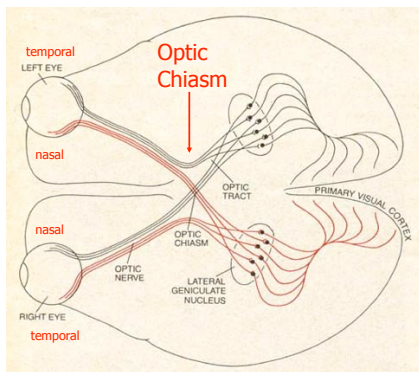


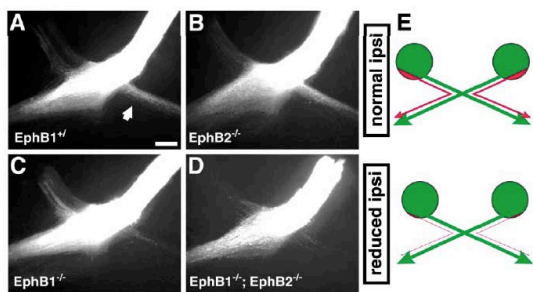
Fig. 13.20 The Action of Semaphorin 1 in the Developing Grasshopper Limb

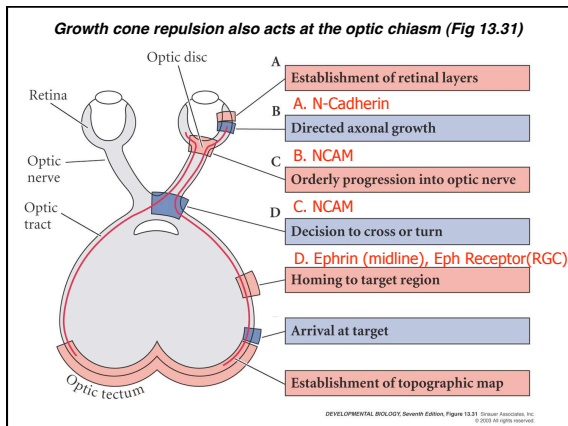


Crossing of retinal axons at the optic chiasm



Ephrins act in contact repulsion at the optic chiasm





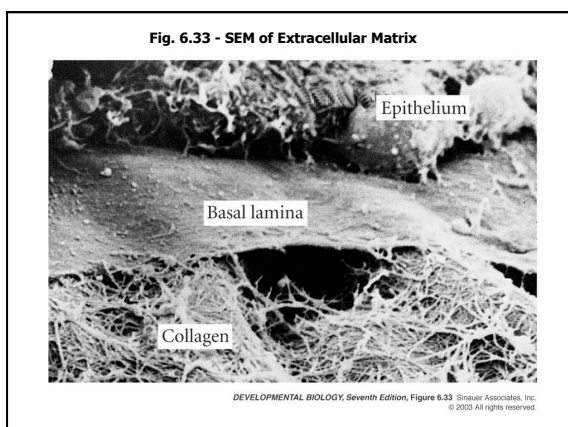
Mechanisms of Growth Cone Guidance

Extracellular Matrix proteins:

Collagen, Fibronectin, Laminin, etc.

ECM Receptors:

Integrins (α and β subunits)



There are many different types of Collagen (at least 16)

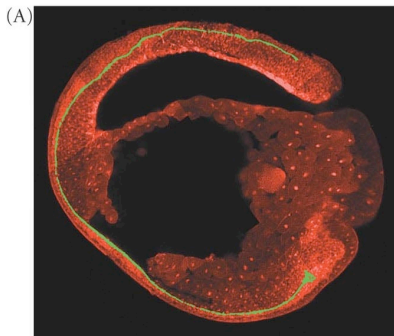
Table 12-2 Types of Collagen and Their Properties

Type	Molecular Formula*	Polymerized Form	Distinctive Features	Tissue Distribution
I	$(\alpha1(I))_2\alpha2(I)$	fibril	low hydroxylysine low carbohydrate broad fibrils	skin, tendon, bone, ligaments, cornea, internal organs (accounts for 90% of body collagen)
II	$(\alpha1(II))_3$	fibril	high hydroxylysine high carbohydrate usually thinner fibrils than type I	cartilage, intervertebral disc, notochord, vitreous body of eye
III	$(\alpha1(III))_3$	fibril	high hydroxyproline low hydroxylysine low carbohydrate	skin, blood vessels, internal organs
IV	$(\alpha1(IV))_3$ (controversial)	basal lamina	very high hydroxylysine high carbohydrate probably retains procollagen extension peptides	basal laminae
V	$(\alpha1(V))_2\alpha2(V)$	unknown	high hydroxylysine high carbohydrate	widespread (in small amounts)

*The seven different α -chains are designated $\alpha1(I)$ through $\alpha1(V)$, $\alpha2(I)$, and $\alpha2(V)$.

80-90% of collagen is types I-III

Fig. 6.23(A - rotated) - **Fibronectin** in the ECM of *Xenopus* Gastrula



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Mechanisms of Growth Cone Guidance

ECM Receptors:

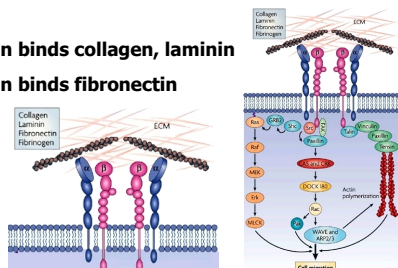
Integrins (α and β subunits)

Different combinations of α and β subunits make receptors that bind different ECM proteins:

Examples -

$\alpha_1\beta_1$ integrin binds collagen, laminin

$\alpha_5\beta_1$ integrin binds fibronectin



Mechanisms of Growth Cone Guidance

Chemoattraction by diffusible substances

Chemoattraction of growth cones first shown with NGF

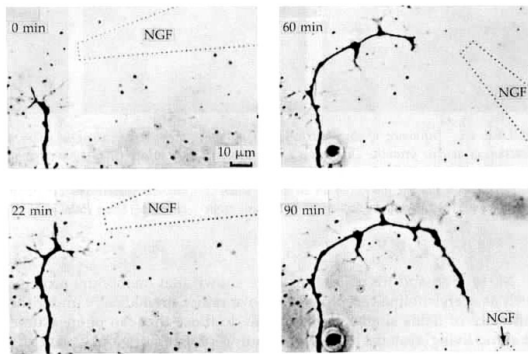
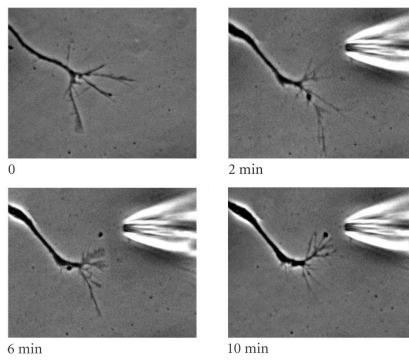
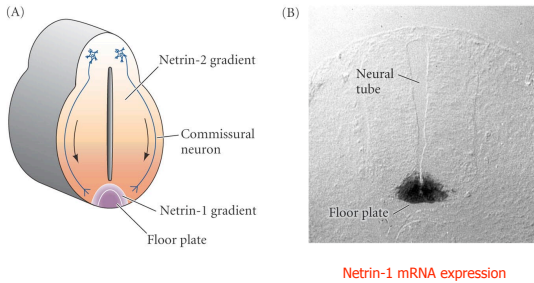


Fig 13.27 Embryonic Axon From a Rat Dorsal Root Ganglion Turn in Response to NT-3



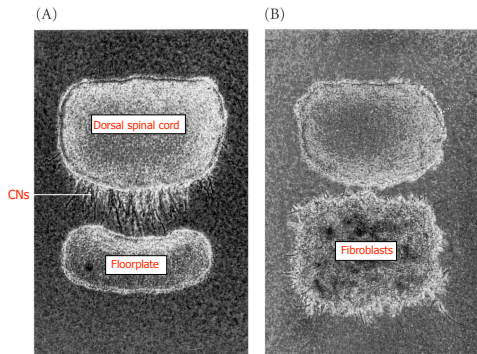
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Figure 13.22 Trajectory of the Commissural Axons in the Rat Spinal Cord



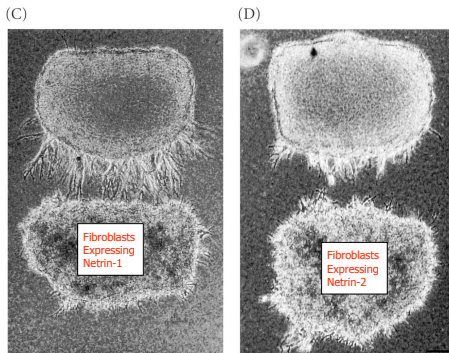
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Fig. 13.23 Transformed Chick Fibroblast (COS) Cells Secreting Netrins Elicit Axon Outgrowth of Commissural Neurons from 11-d Emb. Rat Dorsal Spinal Cord Explants



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Fig. 13.23 Transformed Chick Fibroblast (COS) Cells Secreting Netrins Elicit Axon Outgrowth of Commissural Neurons from 11-d Emb. Rat Dorsal Spinal Cord Explants



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Signals can be attractive or repulsive, depending on the receptor

Netrin-mediated dorsal/ventral signaling is an ancient feature of animal nervous systems.

Netrin signaling also patterns growth cone guidance and cell migration in the nematode *C. elegans*.

Mutants with abnormal nervous system function ("unc" - uncoordinated) were found to be affected in netrin-mediated growth cone guidance.

The first example of a netrin receptor mediating *chemorepulsion* was discovered in *C. elegans* (the *unc-5* gene).

Chemorepulsion by netrins is also found in the vertebrates.

Figure 13.24 C.e. Netrin Expression And Function in Axonal Guidance

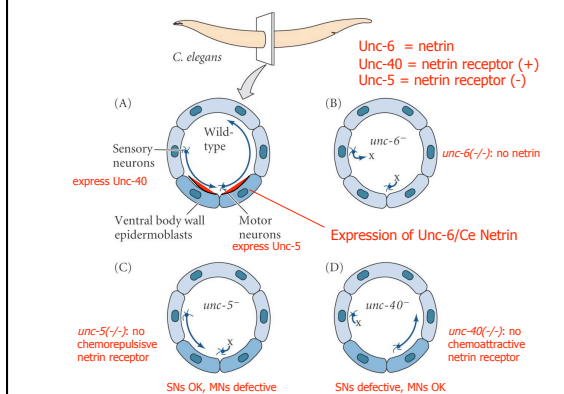
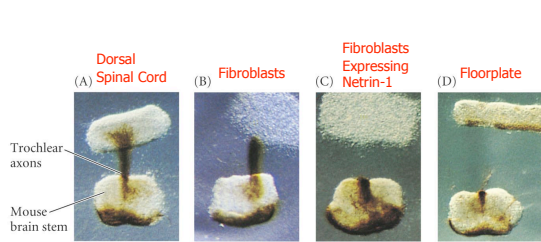


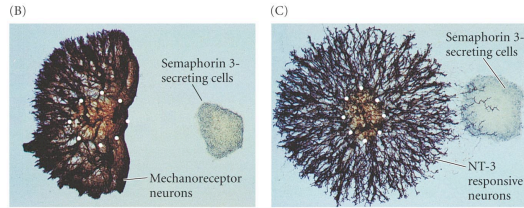
Figure 13.25 Netrins inhibit Outgrowth of Trochlear Axons From Explants of the Mouse Brain Stem



A vertebrate example of Chemorepulsion by a netrin

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Figure 13.21(2) Semaphorin 3 as a Selective Inhibitor of Axonal Projections into the Ventral Spinal Cord



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Engrailed-2 as a secreted chemorepulsive and chemoattractive agent for *Xenopus* Retinal Neurons

Abstract

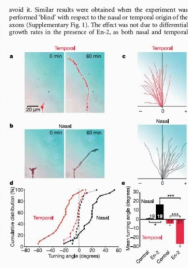
Vol 42(3) November 2005;doi:10.1002/net.20480

The transcription factor Engrailed-2 guides retinal axons

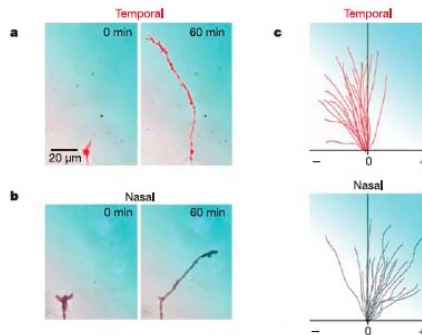
Isabelle Brunet^{1,2}, Christine Wein¹, Michael Piper¹, Alain Tremblau¹, Michel Volovitch¹, William Harris¹, Alain Prochiantz¹ & Christine Hall¹

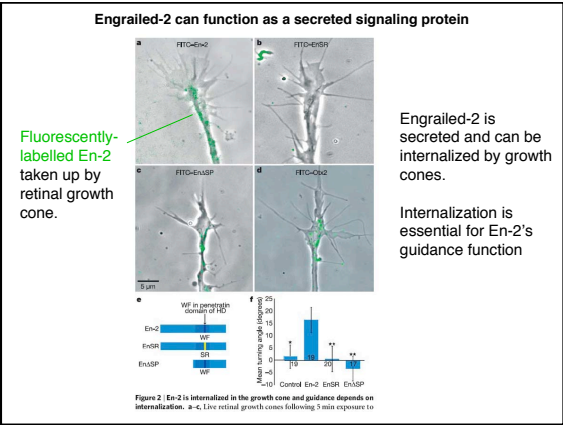
Engrailed-2 (En-2), a homeodomain transcription factor, is expressed in a caudal-ventral gradient in the developing midbrain, where it has an instructive role in patterning the optic tectum—the target of topographic retinal input^{1,2}. In addition to its well-known role in regulating gene expression through its DNA-binding domain, En-2 may also have a role in cell-cell communication, as suggested by the presence of other domains involved in nuclear export, secretion and internalization³. Consistent with this possibility, here we report that an external gradient of En-2 protein strongly repels growth cones of *Xenopus* axons originating from the temporal retina and, conversely, attracts nasal axons. Fluorescently tagged En-2 axons invade growth cones within minutes of exposure, and a neutral form of the protein that cannot enter cells fails to elicit axon turning. Once internalized, En-2 stimulates the rapid phosphorylation of proteins involved in translation initiation and triggers the local synthesis of new proteins. Furthermore, the turning responses of both nasal and temporal growth cones in the presence of En-2 are blocked by inhibitors of protein synthesis. The differential guidance of nasal and temporal axons reported here suggests that En-2 can participate directly in topographic map formation in the vertebrate visual system.

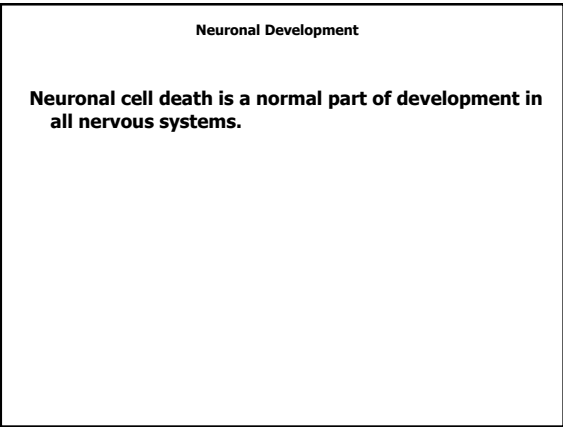
The new molecular insight into the mechanism of topographic mapping by the retinal tectum is a causal gradient of En-2, whose chemorepulsive properties promote matching gradients of receptors and ligands within the retina and tectum. The first candidate molecule fulfilling the requirements of this hypothesis, the Ephrins, were identified in the tectum and found to be repulsive to retinal axons expressing EphA receptors^{4,5}. Temporal axons expressing high levels of EphA receptors map to the nasal tectum and avoid the Ephrins-rich caudal tectum. En-2, which is also expressed in a caudal-ventral gradient in the developing tectum, has been shown to promote the expression of local Ephrins^{6,7} and, through its transcriptional activity, is thought to have a major role in setting up the Ephrins gradient. However, work in knockout mice

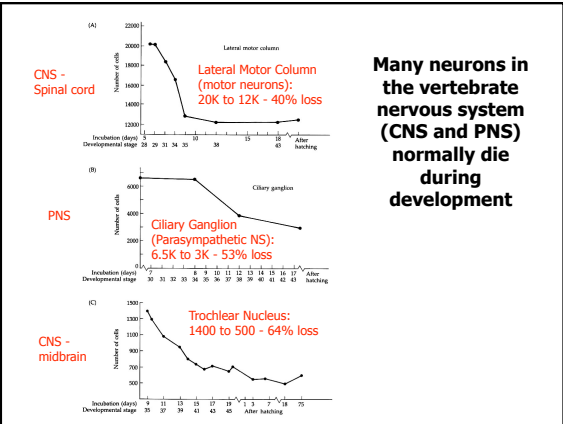


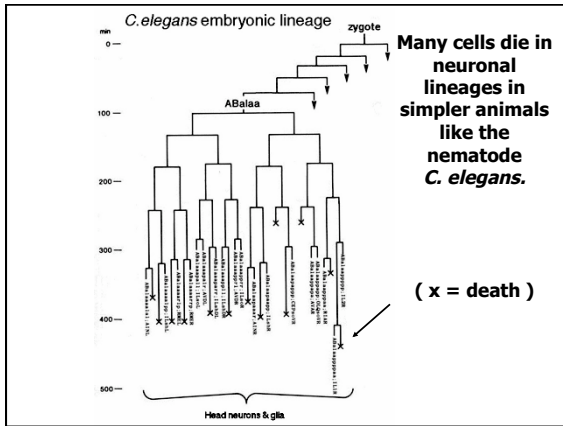
Temporal *Xenopus* Retinal Neurons are repulsed by gradient of En-2; Nasal Retinal Neurons are attracted.

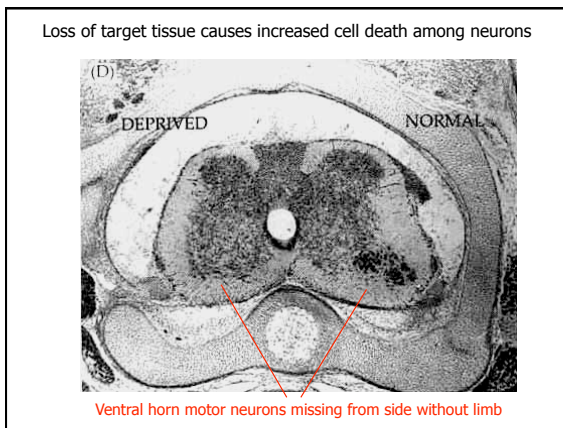


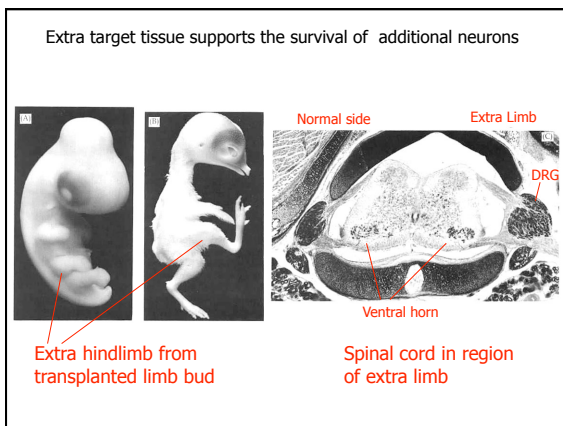


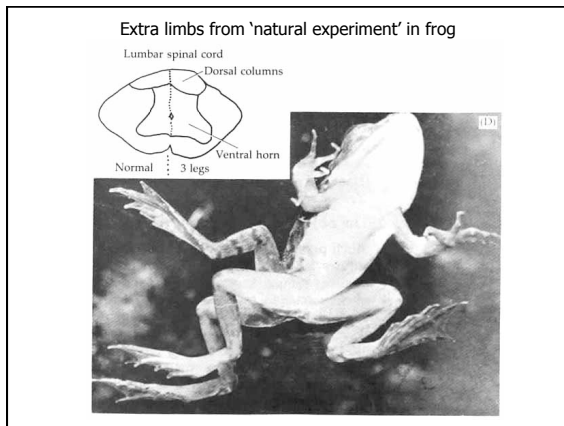


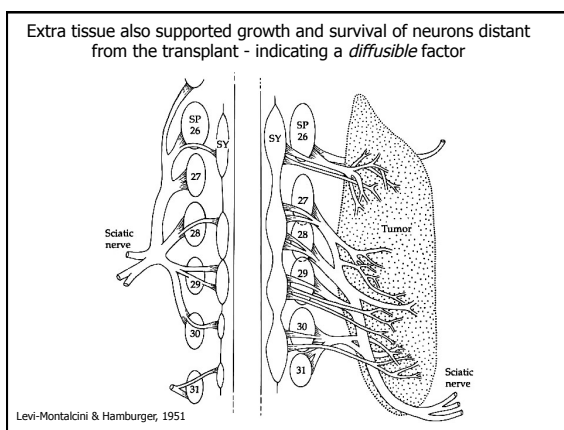


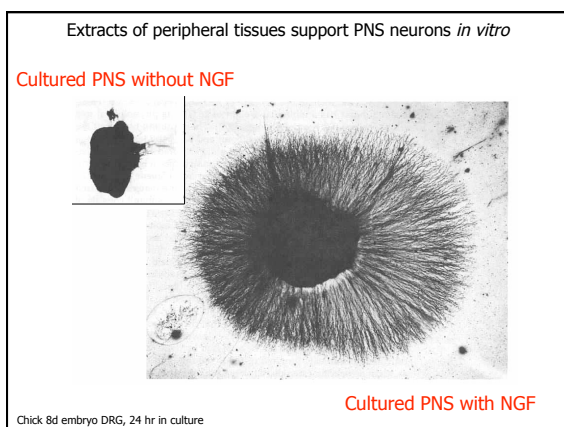












Discoverers of Nerve Growth Factor (NGF)



Viktor Hamburger
1900 - 2001



1965



Rita Levi-Montalcini
b. 1909



The Nobel Prize in Physiology or Medicine 1986

"for their discoveries of growth factors"



Stanley Cohen

1/2 of the prize
USA

Vanderbilt University
School of Medicine
Nashville, TN, USA

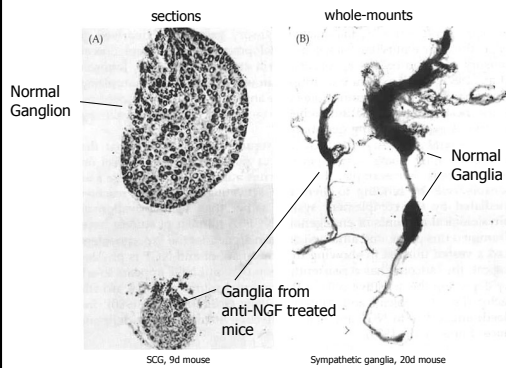


Rita Levi-Montalcini

1/2 of the prize
Italy and USA

Institute of Cell Biology of
the C.N.R.
Rome, Italy

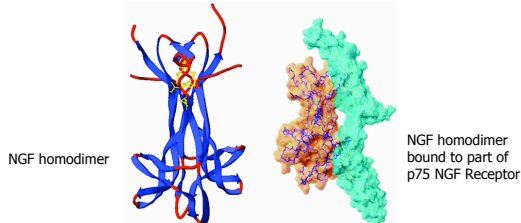
NGF function is essential for neuronal survival *in vivo*



Levi-Montalcini, 1972

Neurotrophic Factors support the survival of specific types of neurons

Neurotrophins - family of NGF-related proteins
NGF - Nerve Growth Factor
BDNF - Brain-derived neurotrophic factor
NT3 - Neurotrophin 3
NT4 - Neurotrophin 4



Neurotrophic Factors support the survival of specific types of neurons

Neurotrophins - family of NGF-related proteins
NGF - Nerve Growth Factor
BDNF - Brain-derived neurotrophic factor
NT-3 - Neurotrophin 3
NT-4 - Neurotrophin 4

13 kDa secreted glycoproteins (as homodimer)

Bind to receptor tyrosine kinases: TrkA, TrkB, TrkC

e.g., NGF receptors - TrkA
BDNF, NT-4 receptor - TrkB
NT-3 receptor - TrkC
(NT-3 also binds to TrkA, TrkB)

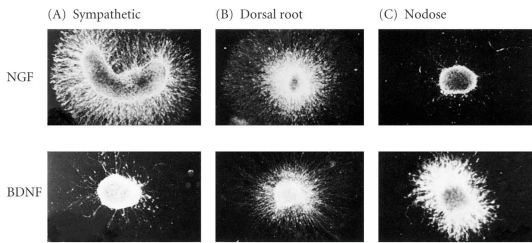
Neurotrophic Factors support the survival of specific types of neurons

Neurotrophins - family of NGF-related proteins
NGF - Nerve Growth Factor
BDNF - Brain-derived neurotrophic factor
NT3 - Neurotrophin 3
NT4/5 - Neurotrophin 4/5

Other Neurotrophic and Differentiation Factors:

GDNF- Glial-derived neurotrophic factor
CDF/LIF - Cholinergic differentiation factor/
Leukemia inhibiting factor

Figure 13.29 Effects of NGF And BDNF On Axonal Outgrowth



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Neurotrophic Factors support the survival of specific types of neurons

Three potential functions of neurotrophic and related factors

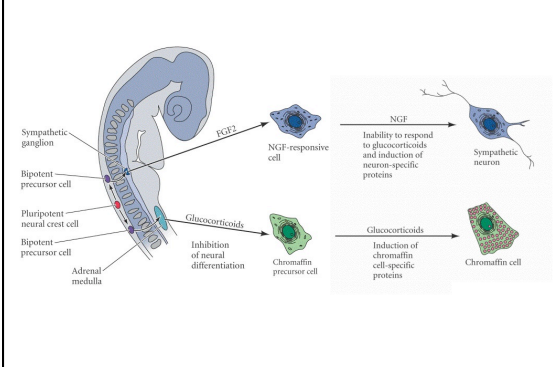
1. *Neurotrophic* factor
2. *Chemoattractive* factor
3. *Instructive* (inductive/signaling) factor

NGF shows all three functions

Experiment showing NGF's instructive role:

NGF can convert presumptive adrenal medulla cells (would become chromaffin cells) into neurons.

Figure 13.7 Differentiation of trunk neural crest cells



Neuronal Differentiation Factors

Example: CDF/LIF

CDF is made by heart muscle cells.

Some NC-derived neurons that innervate heart initially make norepinephrine (NE).

Interaction with heart converts cells to use ACh.

CDF has other roles in development:

LIF (immune system)

Essential for implantation of blastocyst
