

By the time I was born, more of me had died than survived. It is no wonder I cannot remember; during that time I went through brain after brain for nine months, finally contriving the one model that could be human, equipped for language.

Lewis Thomas (1992)

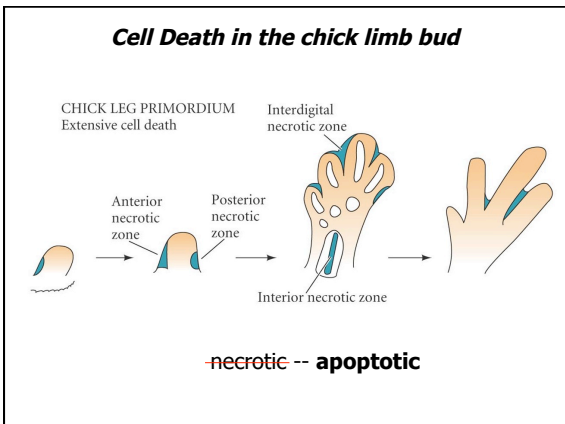
Cell Death in Development

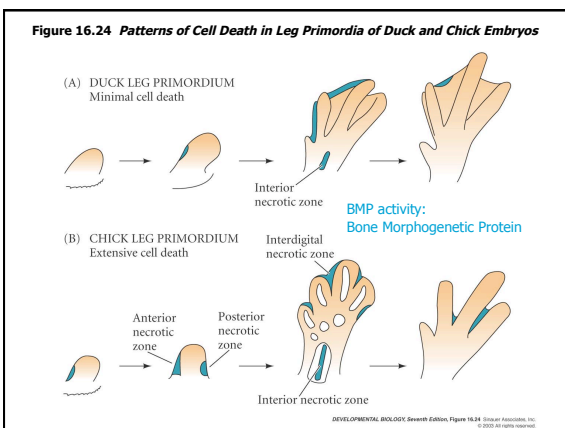
Programmed Cell Death / Apoptosis

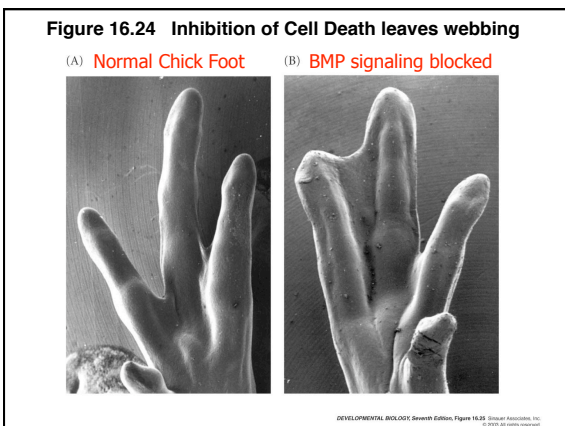
Cell Death in Development

Cell death plays an important role in morphogenesis.

Example: Interdigital death in limb bud.







Cell death is (in the vertebrates) prominent in

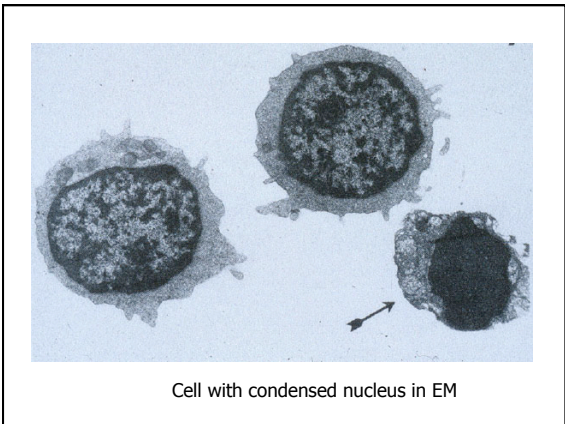
- developing nervous system
- developing and mature immune system

1. Immune cells recognizing 'self' die during immune system development.
2. Immune challenge results in proliferation of cells; when these cells are no longer needed, they die.

Two types of cell death:

Necrosis - caused by acute injury, involves cell lysis
- undesirable because cell contents are released

Apoptosis / Programmed Cell Death
- stereotyped pattern of events including
→ nuclear condensation



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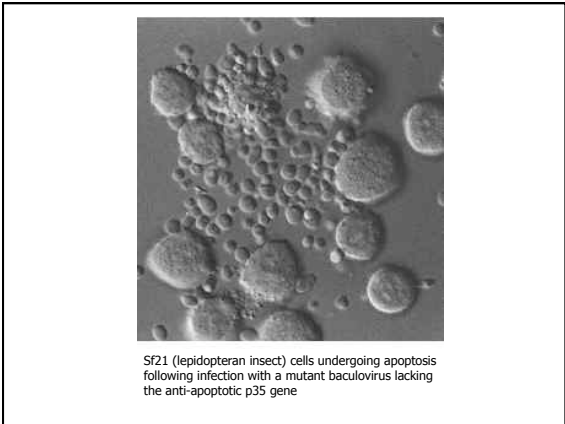
Apoptosis / Programmed Cell Death
- stereotyped pattern of events including nuclear condensation
→ chromosome fragmentation

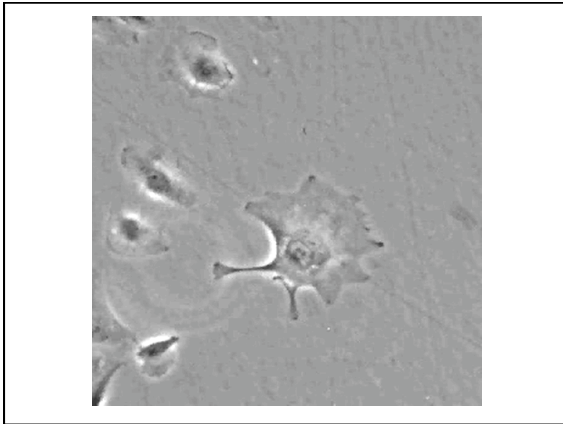
"TUNEL" - TdT-mediated dUTP Nick End Labeling
- shows 'ends' of chromosomes
-- few ends in normal cells
-- many in apoptotic cells undergoing fragmentation

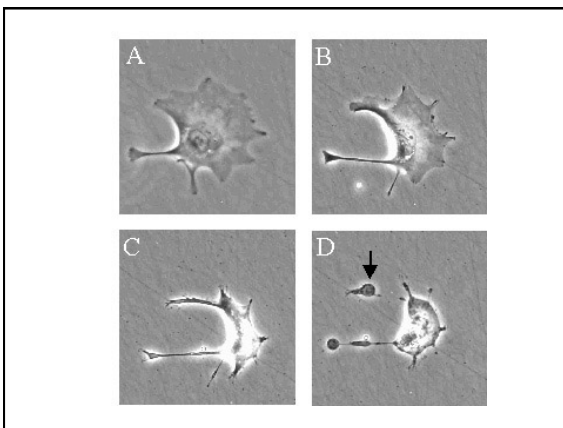
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→ cell membrane blebbing







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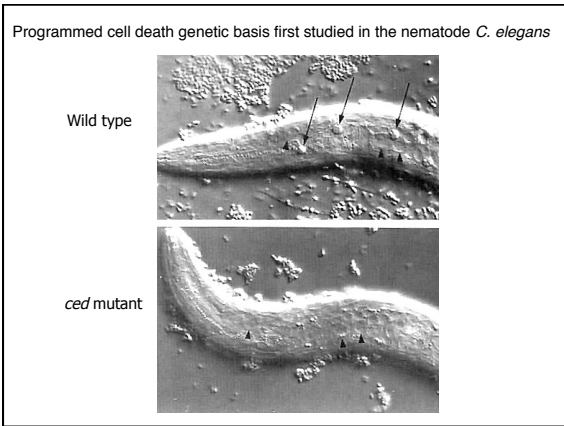
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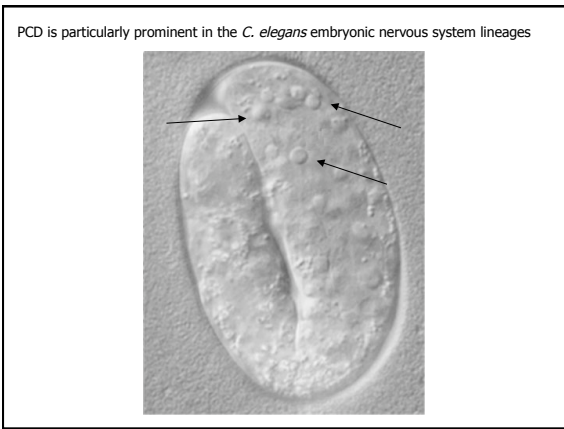
Apoptosis / Programmed Cell Death

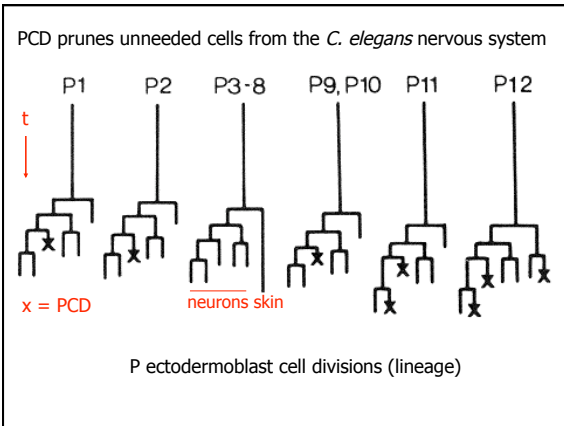
- stereotyped pattern of events including
nuclear condensation
chromosome fragmentation
cell membrane blebbing

→ phagocytosis by nearby cells

- active suicide program, often requiring gene activation (transcription & translation)







Programmed cell death genetic basis first studied in the nematode *C. elegans*

Steps in cell death process identified by mutants:

Decision: *ced-9, egl-1*

Execution: *ced-3, ced-4*

Engulfment: *ced-1, ced-2*

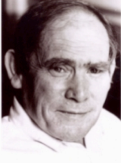
Digestion of DNA: *nuc-1*

Vertebrates have similar proteins: *bcl-2* is homolog of *ced-9*


Nobel Prize for work in the nematode *C. elegans* (including cell death genetics)

2002 Nobel Prize awarded to Brenner, Horvitz & Sulston for Studies of the Genetic Regulation of *C. elegans* Development


- [The Nobel Prize in Physiology or Medicine 2002](#) - official Nobel Committee website
- [Nobel Committee Press Release](#)



Sydney Brenner



Bob Horvitz

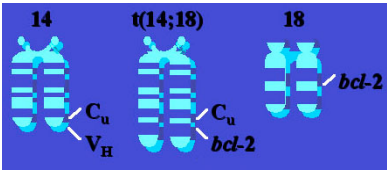


John Sulston

Mammalian Apoptosis Genes

bcl-2 was first discovered as an oncogene in B cell lymphomas.

bcl-2 protein coding sequence translocated from chromosome 18 to 14 (t(14;18)) in front of Ig Heavy Chain promoter.



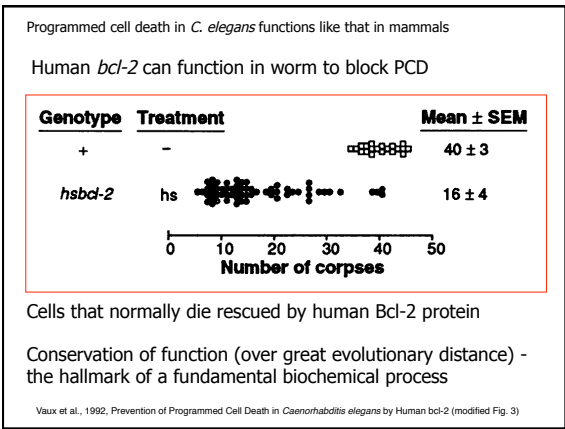
bcl-2 homology to *C.e. ced-9* gene gave a clue to its function.

bcl-2 gene was permanently ON in these B cells, blocking apoptosis, immortalizing them - pre-disposing cells to cancer.

Mammalian Apoptosis Genes

bcl-2 was shown able to block apoptosis in IL-3 deprived B cells.

bcl-2 gene inserted and activated in *C. elegans* could rescue worm cells from programmed cell death.

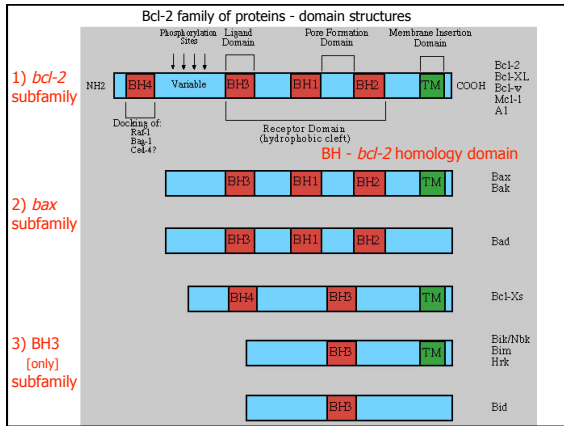


Mammalian Apoptosis Genes

Mammals have many *bcl-2*-like genes regulating apoptosis.

Three subfamilies:

- 1) Bcl-2 subfamily (e.g., *bcl-2*, *Bcl-XL*) promotes cell survival.
- 2) Bax subfamily (e.g., *Bax*, *Bix*) is pro-apoptotic.
- 3) BH3 subfamily (e.g., *Bad*, *Bik*) is pro-apoptotic.



Mammalian Apoptosis Genes

Other worm *ced* genes also have mammalian homologs.

egl-1 gene first discovered as a dominant mutation causing PCD in the egg laying neuron HSN.

Egl-1 protein is a homolog of pro-apoptotic BH3 subfamily proteins.

Mammals have 10 Caspases functioning in apoptosis:

- [Caspase-1: ICE protease, the first discovered in mammals]
- Caspase-9: closest *ced-3* homolog
- Caspase-9 (-) mutant in mice is PCD-deficient, especially in the CNS.

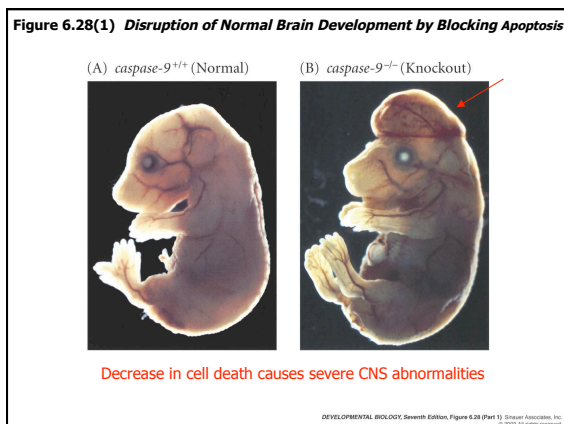
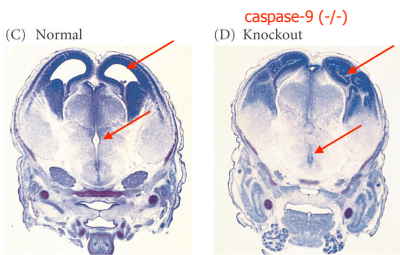


Figure 6.28 Disruption of Normal Brain Development by Blocking Apoptosis



Note lack of spaces (ventricles) found in normal brain

DEVELOPMENTAL BIOLOGY, Seventh Edition, Figure 6.28 (Part 2) © 2003 Sinauer Associates, Inc. and W. H. Freeman & Co.

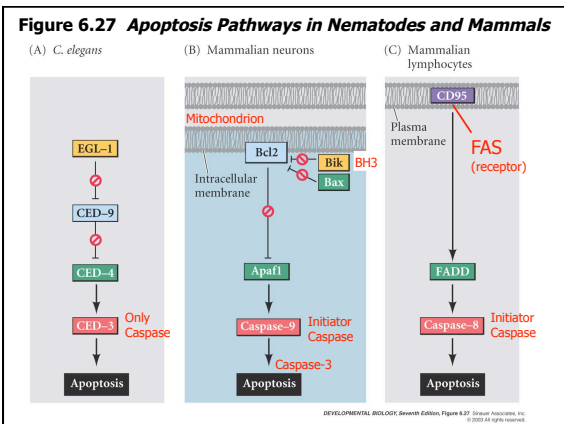
Mammalian Apoptosis Genes

Other worm *ced* genes also have mammalian homologs
Caspases function as initiator and effector caspases
All caspases synthesized as inactive pro-caspases
Cleavage activates caspases to be functional enzymes
Caspase-9 (Ced-3 homolog) is an initiator caspase; one function:
cleave & activate caspase-3 (effector)
Caspase-3 and other effectors begin digestion of cell contents
Ced-4 homolog: Apaf-1
Apaf-1 binds Pro-caspase-9, promotes its auto-cleavage to active caspase

Mammalian Apoptosis Genes

Basic molecular pathway conserved from worms to humans
Mammals have multiple pathways to activate a caspase cleavage cascade
(vs. nematodes with only one pathway)
Some mammalian pathways bypass bcl-2/mitochondrial pathway





"Death Receptors"

Fas (aka CD95 or Apo-1) is a 'death receptor' in the TNF receptor superfamily (TNF = Tumor Necrosis Factor)

Death receptors mediate active killing signals by (e.g.) cytotoxic T cells.

Death receptors mediate killing of auto-reactive immune cells.

