

# Genetics of Cancer

## Lecture 33

### Alterations in different kinds of Genes cause Cancer

#### Oncogenes

dominant gain-of-function mutations  
promote cell transformation

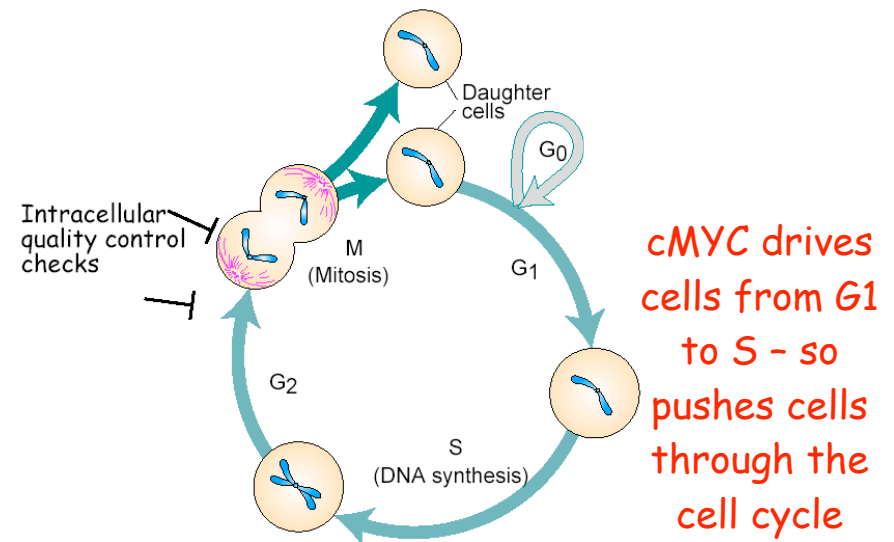
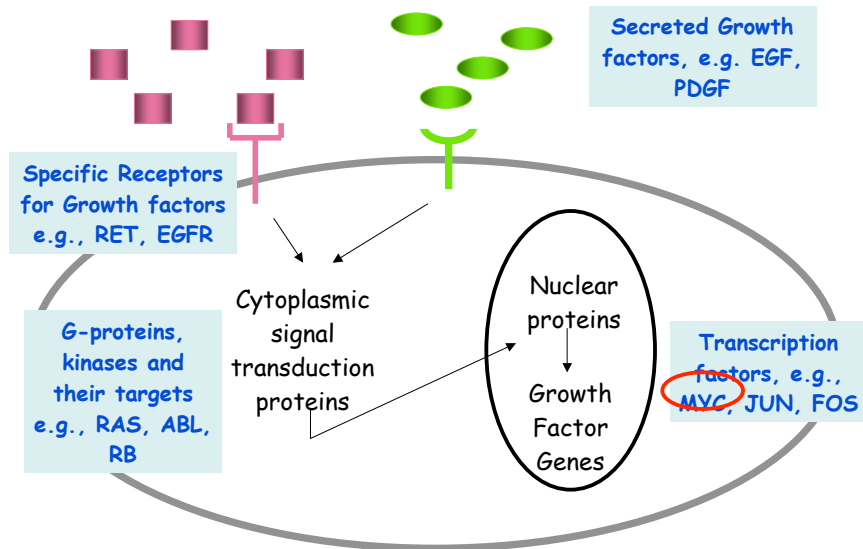
#### Tumor suppressor genes

recessive, loss-of-function mutations  
promote cell transformation

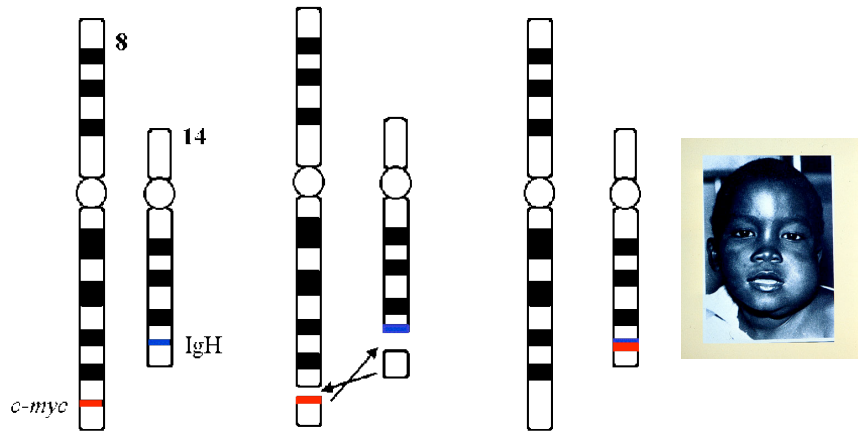
#### Mutator genes

Usually recessive, loss-of-function mutations  
that increase spontaneous and environmentally

### Signal Transduction and Growth Regulation



Burkitt's Lymphoma: A chromosome translocation  
 → cMYC to be expressed inappropriately in B-cells



cMYC drives cells from G1 to S

Another way that oncogenic transcription factors can be up-regulated: Gene Amplification



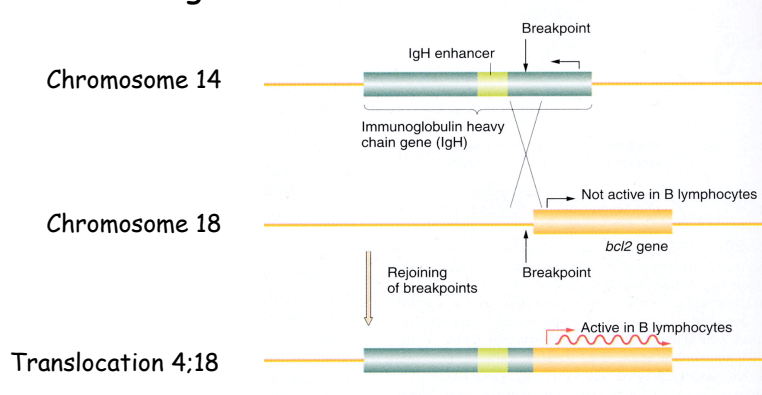
Blue - staining of all chromosomes

Red - staining of chromosome 4

Green - staining of the N-MYC gene

(N-MYC and cMYC share many similar properties)

One more example - with an interesting twist  
 A translocation between Chr 14 and Chr 18 to put the BCL2 gene under the strong IgH promoter



The BCL2 protein **PREVENTS** programmed cell death, B cells live longer than normal leading to B-cell Lymphomas

What chromosomal events convert proto-oncogenes to dominantly acting oncogenes

- Point mutations (e.g., RAS)
- Partial deletion mutations (e.g., RTKs)
- Chromosomal translocations that produce novel fusion proteins (e.g., Bcr-Abl)
- Chromosomal translocation to juxtapose a strong promoter upstream and the proto-oncogene such that it is inappropriately expressed (e.g., cMyc, Bcl2)
- Gene amplification resulting in overexpression

# Alterations in different kinds of Genes cause Cancer

## Oncogenes

dominant gain-of-function mutations promote cell transformation

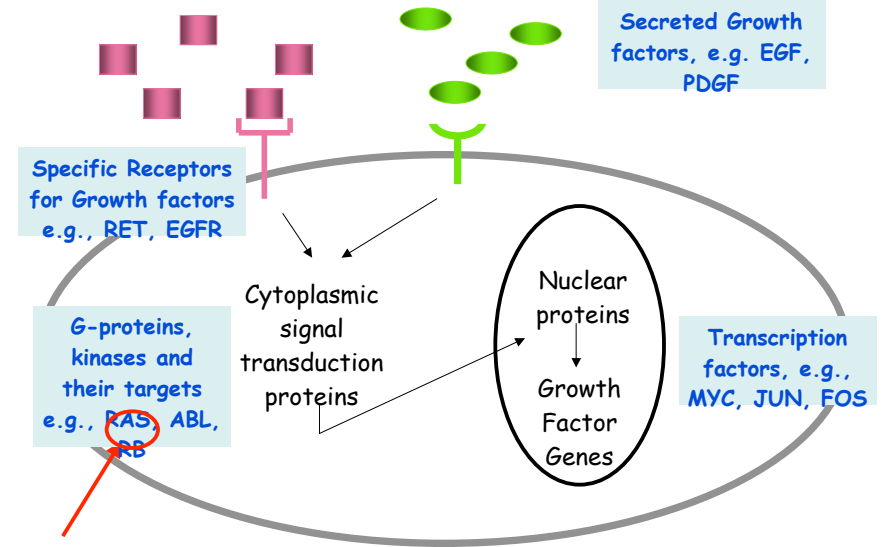
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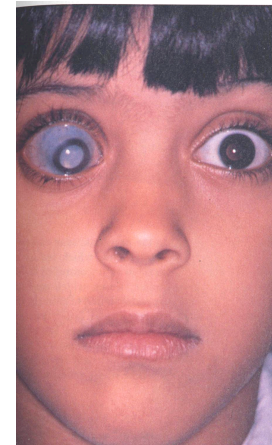
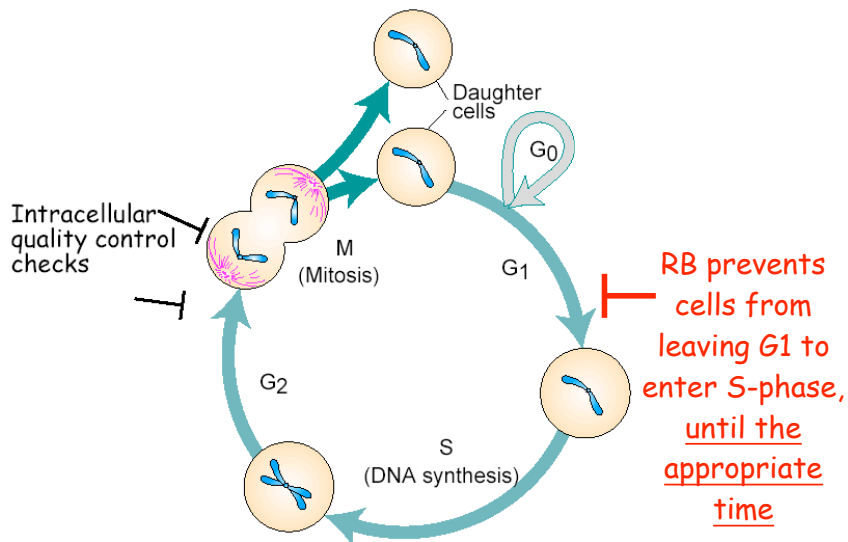
## Mutator genes

Usually recessive, loss-of-function mutations that increase spontaneous and environmentally

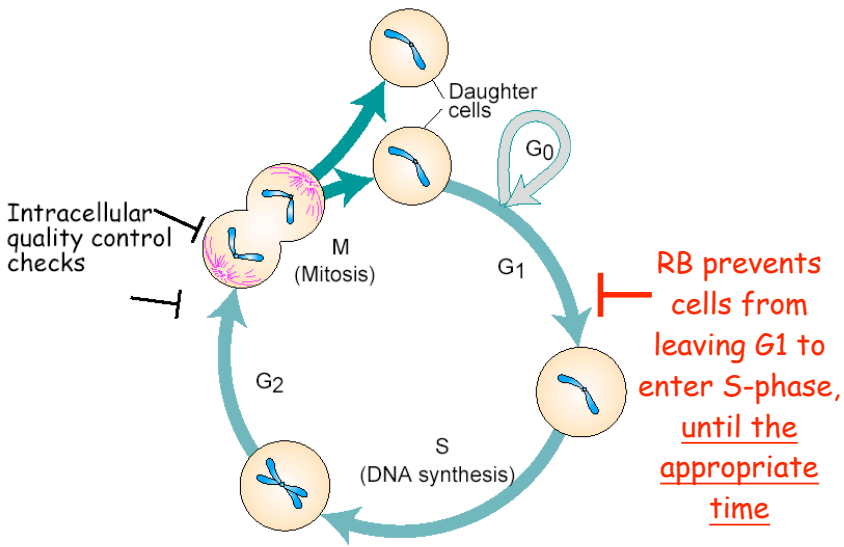
# Signal Transduction and Growth Regulation



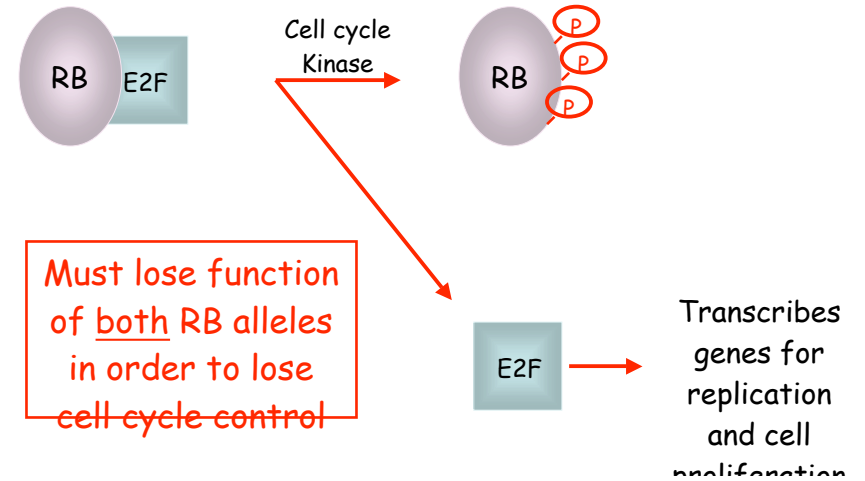
RB - the **Retinoblastoma** Gene - was the first example of a **Tumor Repressor Gene** (aka a Recessive Oncogene)



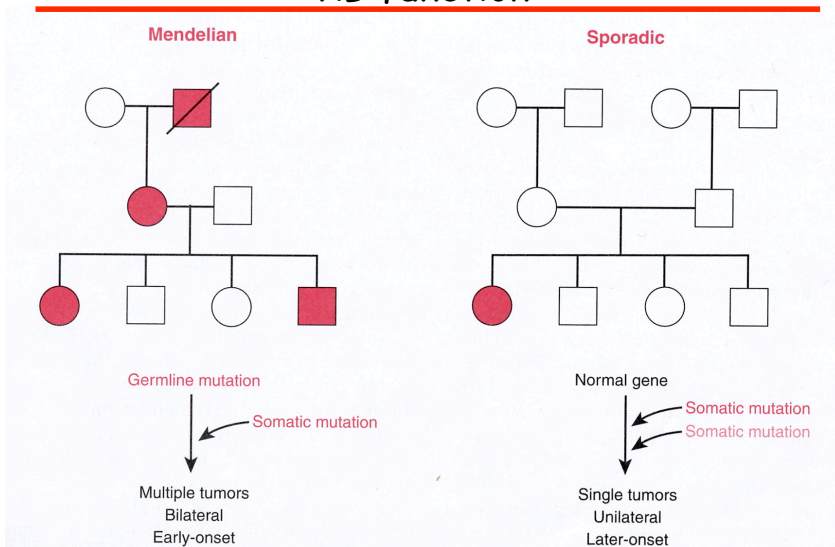
Loss of Function Mutations in both RB genes lead to malignant tumors of the retina during the first few years of life



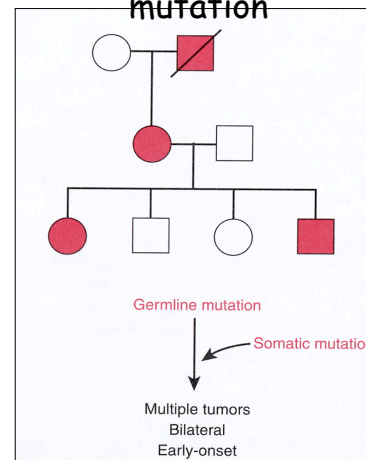
Phosphorylation of RB at the appropriate time in G<sub>1</sub> allows release of the E2F Transcription Factor



Two ways to get retinal tumors due to loss of RB function



The Retinoblastoma disease behaves as an autosomal dominant mutation



• In order to lose cell cycle control MUST lose function of both alleles

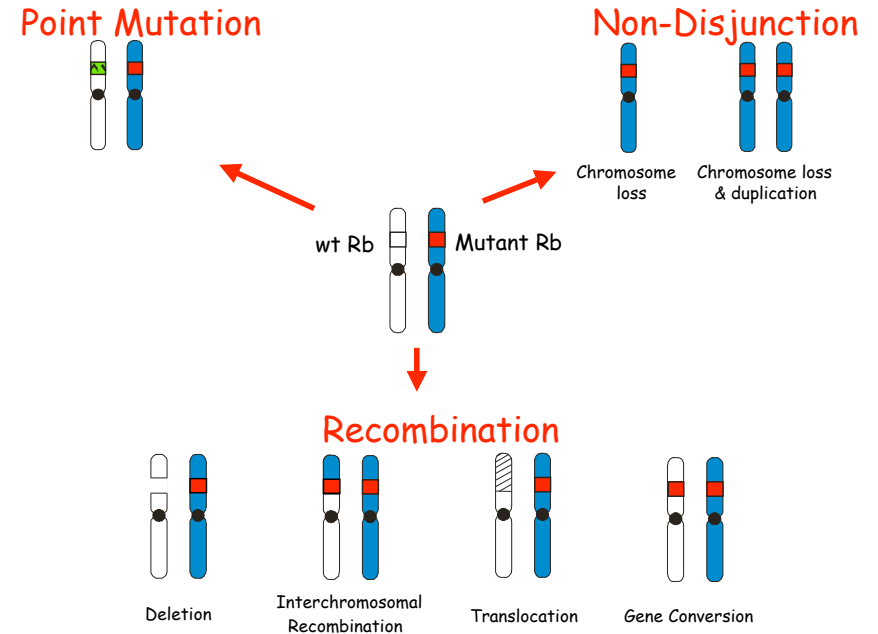
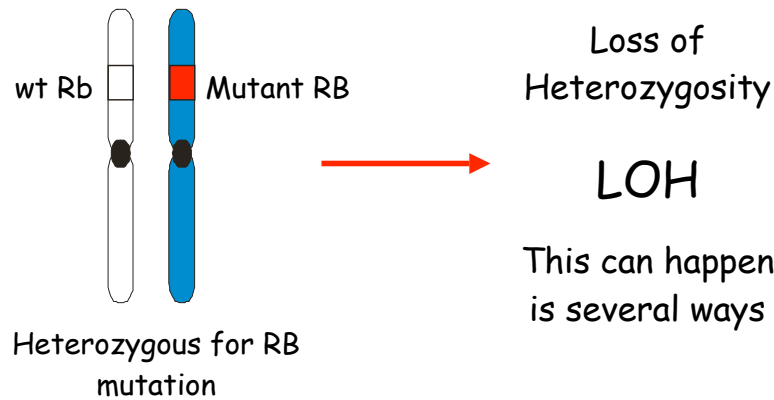
• But, for Mendelian inheritance of RB, children need only inherit only one non-functional allele

• To explain this the "TWO HIT" hypothesis was proposed

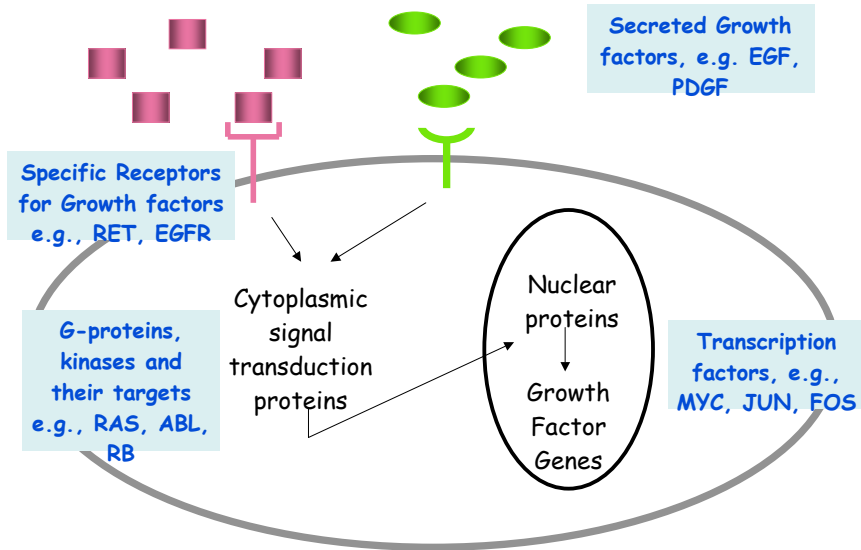
• During development of the retina a second mutation is almost certain to occur

• RB is one of the very few cancers

# How is the second RB allele rendered non-functional?



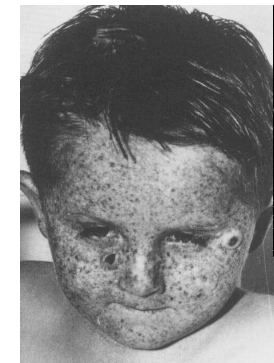
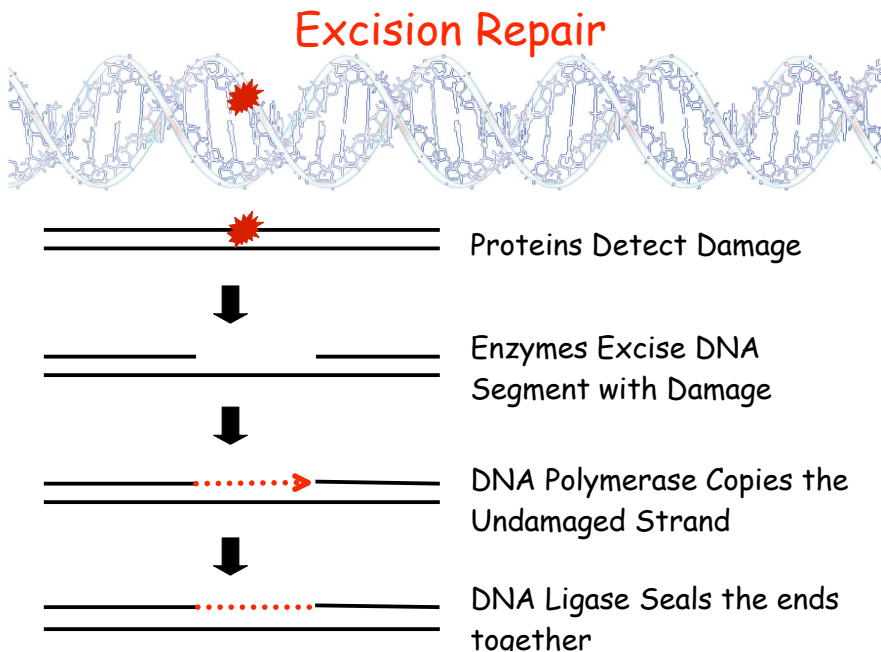
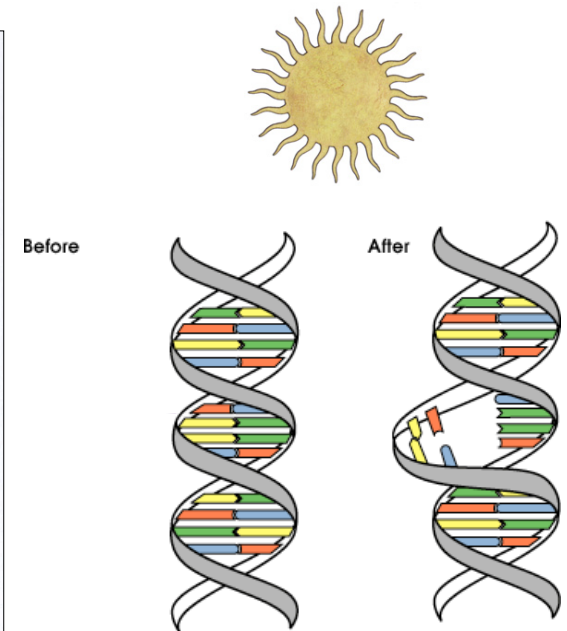
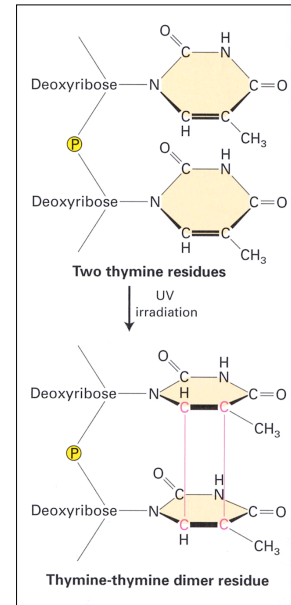
# Signal Transduction and Growth Regulation



# What chromosomal events convert proto-oncogenes to dominantly acting oncogenes and contribute to loss of tumor suppressor genes?

- Point mutations (e.g., RAS, Rb)
- Partial deletion mutations (e.g., RTKs)
- Chromosomal translocations that produce novel fusion proteins (e.g., Bcr-Abl)
- Chromosomal translocation to juxtapose a strong promoter upstream and the proto-oncogene such that it is inappropriately expressed (e.g., cMyc, Bcl2)
- Gene amplification resulting in overexpression (e.g., N-Myc)

All of these mutational events are induced by natural and man-made environmental agents

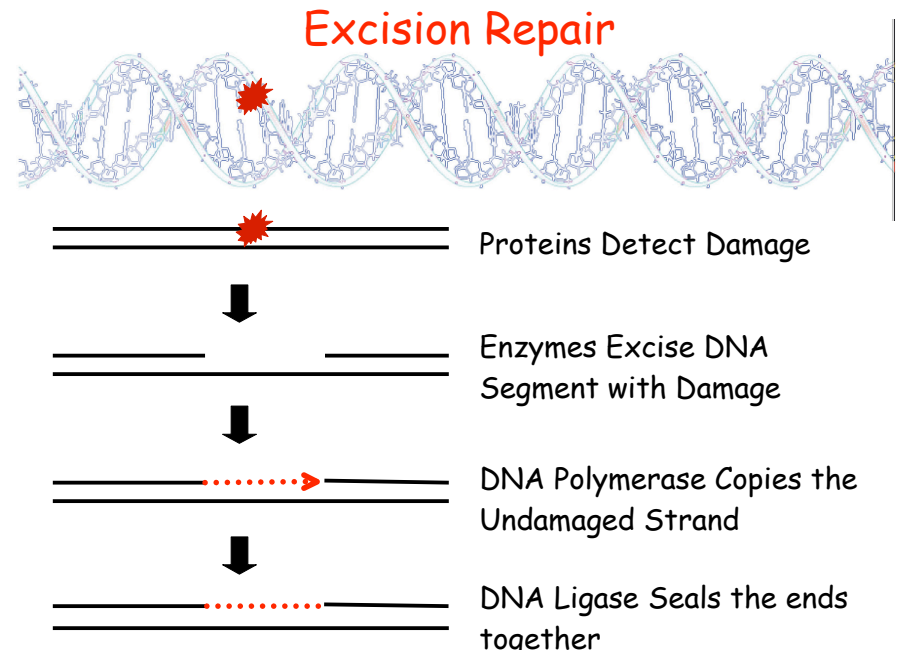
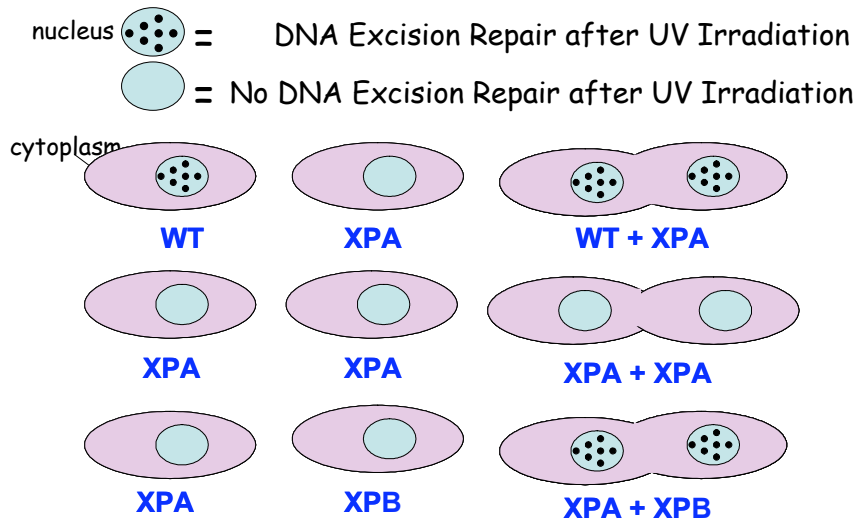


Xeroderma Pigmentosum An Autosomal Recessive Disease

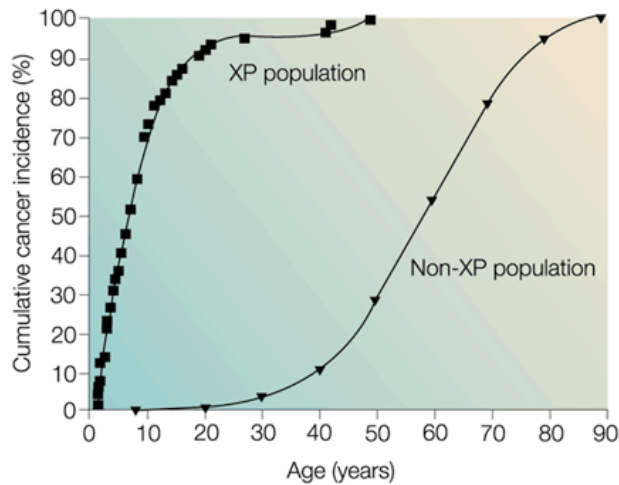
2000-fold increased risk of

..

# Complementation in fused cells reveals 7 genes that cause Xeroderma Pigmentosum

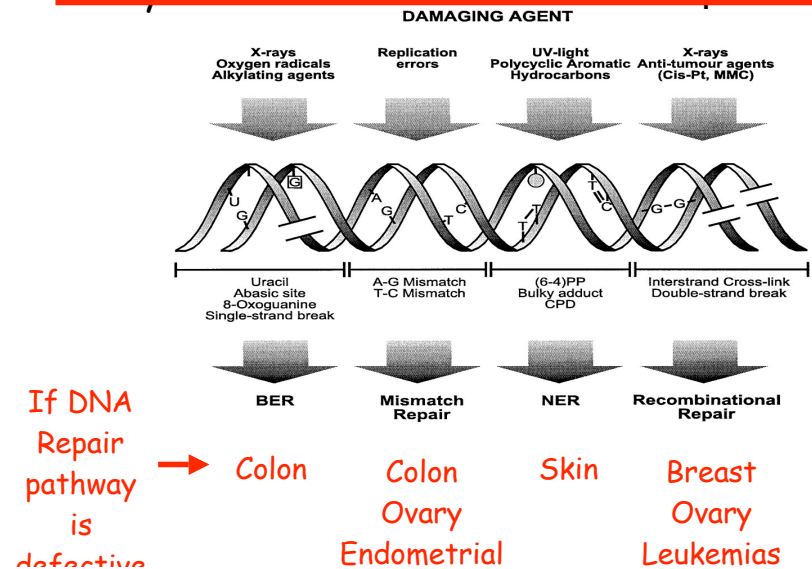


# Age at First Skin Cancer

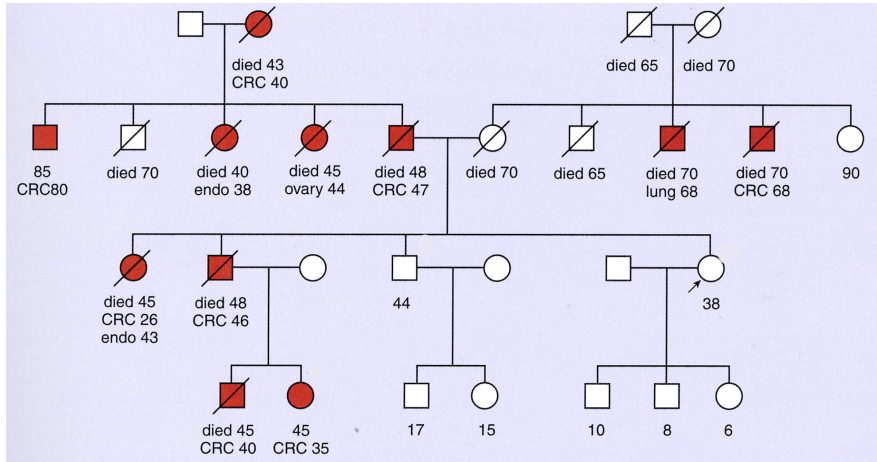


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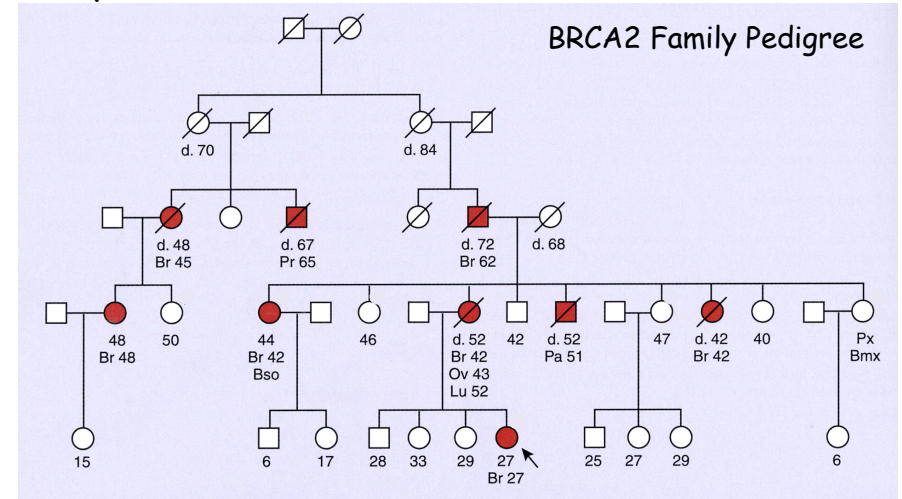
# There are Many Other Human Cancer Prone Syndromes Deficient in DNA Repair



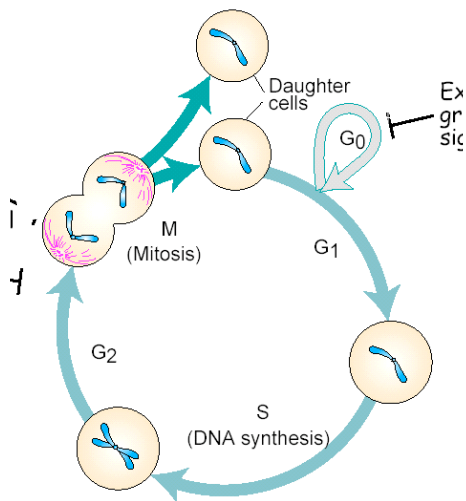
## Hereditary Nonpolyposis Colon Cancer DNA Mismatch Repair Defect Syndrome inherited as Autosomal **Dominant**



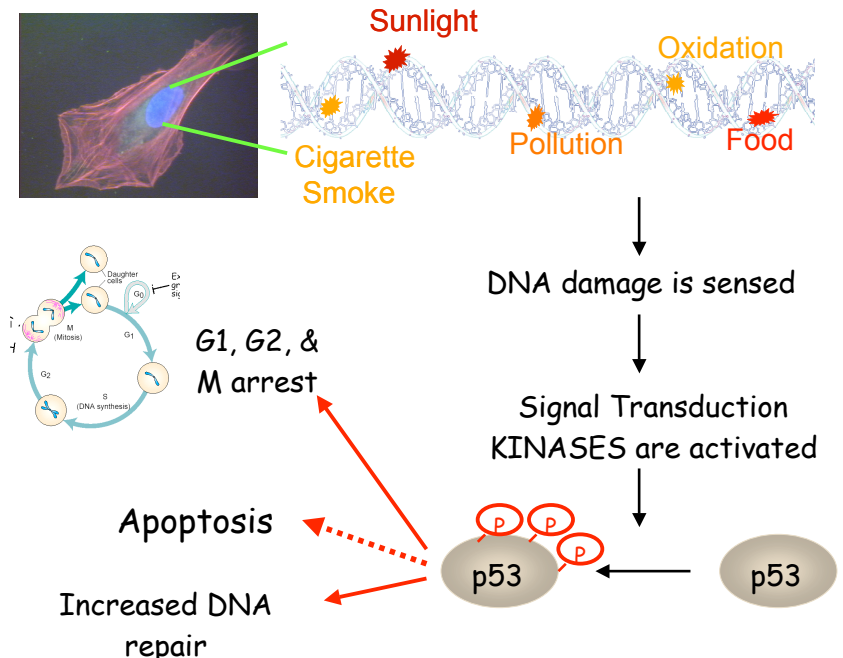
## Hereditary Breast Cancer Susceptibility DNA Recombination Repair Defect Syndrome inherited as Autosomal **Dominant**



## Cells need time to repair DNA: DNA Damage induces Cell Cycle Checkpoints



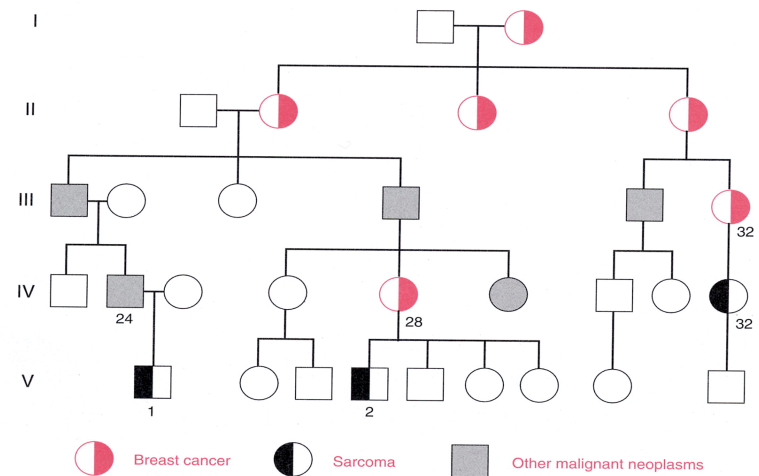
- DNA damage signals cell cycle check points
- If the damage is too great to fix by repair a signal is sent for the cell to undergo suicide *i.e.*,



Loss of p53 function occurs in more than 50% of human cancers!!

- These cancer cells are genetically unstable because they are **less able** to do the following:
- Stop the cell cycling to allow time for DNA repair
- Carry out efficient DNA repair

## Li-Fraumeni Syndrome - Inheritance of one p53 null allele



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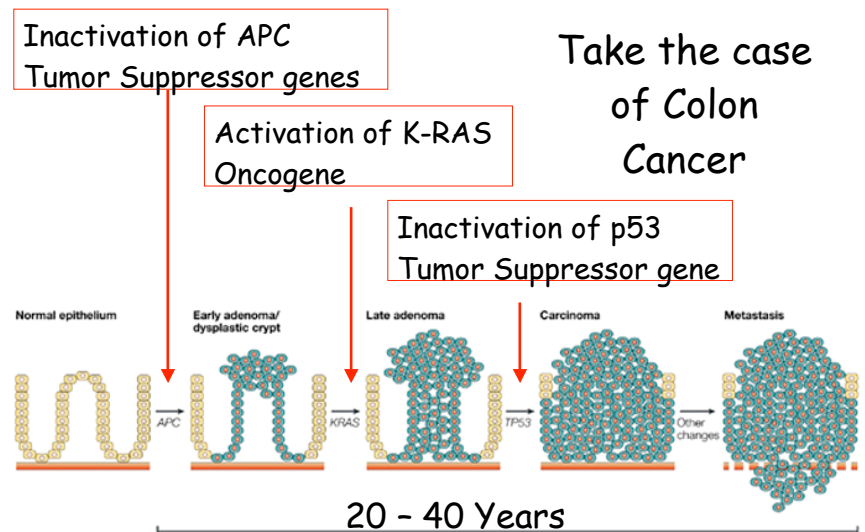
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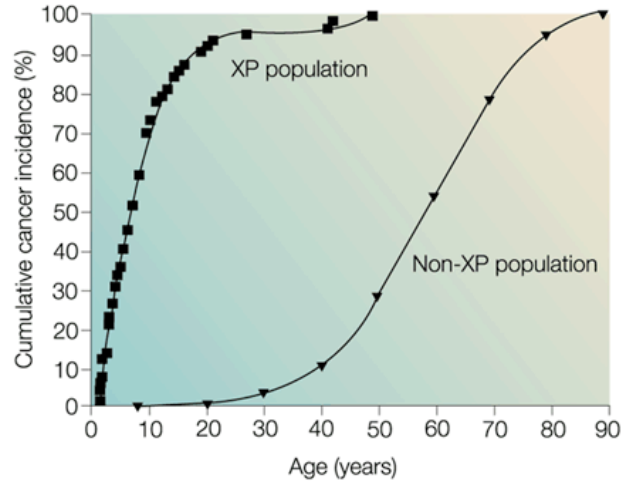
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Usually recessive, loss-of-function mutations that increase spontaneous and environmentally

Most fully blown cancers require inactivation of tumor suppressor genes and activation of oncogenes

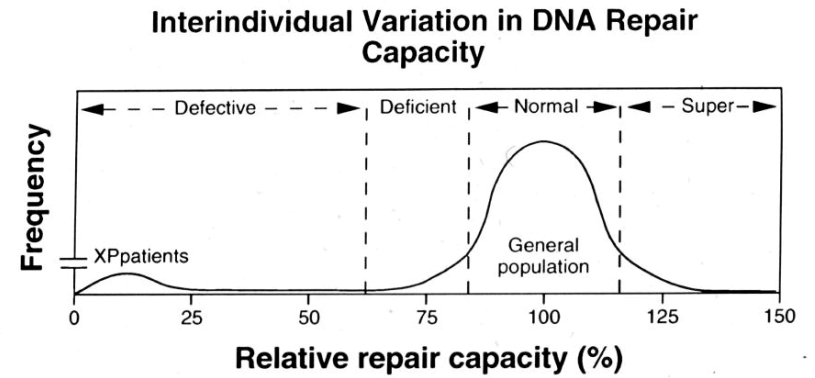


# Age at First Skin Cancer



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# Xeroderma Pigmentosum ~ 1/250,000



Wei *et al.*, Clinical Chemistry, Vol. 41, No. 12, 1995