Biochemistry of Carcinogenesis

Lecture # 35

Alexander N. Koval
What is Cancer?

- The term "cancer" refers to a group of diseases in which cells grow and spread unrestrained throughout the body.
  - It is difficult to imagine anyone who has not heard about this disease. Most people have been affected because either a loved one, a friend, or even they themselves are cancer survivors.
  - It is therefore important for everyone to have a basic understanding about the nature, diagnosis, causes, prevention, and treatment of cancer.

- Learn about the following topics:
  - What is cancer?
  - How is cancer detected and diagnosed?
  - What causes cancer?
  - What is the link between genes and cancer?
  - What is cancer prevention?
Different Kinds of Cancer

- Cancer can originate almost anywhere in the body.

- **Carcinomas**, the most common types of cancer, arise from the cells that cover external and internal body surfaces.
  - Lung, breast, and colon are the most frequent cancers of this type.

- **Sarcomas** are cancers arising from cells found in the supporting tissues of the body such as bone, cartilage, fat, connective tissue, and muscle.

- **Lymphomas** are cancers that arise in the lymph nodes and tissues of the body's immune system.

- **Leukemias** are cancers of the immature blood cells that grow in the bone marrow and tend to accumulate in large numbers in the bloodstream.
2006 Estimated US Cancer Cases*

Men 720,280

Prostate 33%
Lung & bronchus 13%
Colon & rectum 10%
Urinary bladder 6%
Melanoma of skin 5%
Non-Hodgkin lymphoma 4%
Kidney 3%
Oral cavity 3%
Leukemia 3%
Pancreas 2%
All Other Sites 18%

Women 679,510

31% Breast
12% Lung & bronchus
11% Colon & rectum
6% Uterine corpus
4% Non-Hodgkin lymphoma
4% Melanoma of skin
3% Thyroid
3% Ovary
2% Urinary bladder
2% Pancreas
22% All Other Sites

*Excludes basal and squamous cell skin cancers and in situ carcinomas except urinary bladder.
Source: American Cancer Society, 2006.
## 2006 Estimated US Cancer Deaths*

<table>
<thead>
<tr>
<th>Tumor Site</th>
<th>Men</th>
<th>Women</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lung &amp; bronchus</td>
<td>31%</td>
<td>26%</td>
</tr>
<tr>
<td>Colon &amp; rectum</td>
<td>10%</td>
<td>15%</td>
</tr>
<tr>
<td>Prostate</td>
<td>9%</td>
<td>10%</td>
</tr>
<tr>
<td>Pancreas</td>
<td>6%</td>
<td>6%</td>
</tr>
<tr>
<td>Leukemia</td>
<td>4%</td>
<td>4%</td>
</tr>
<tr>
<td>Liver &amp; intrahepatic bile duct</td>
<td>4%</td>
<td>4%</td>
</tr>
<tr>
<td>Esophagus</td>
<td>4%</td>
<td>3%</td>
</tr>
<tr>
<td>Non-Hodgkin lymphoma</td>
<td>3%</td>
<td>3%</td>
</tr>
<tr>
<td>Urinary bladder</td>
<td>3%</td>
<td>2%</td>
</tr>
<tr>
<td>Kidney</td>
<td>3%</td>
<td>2%</td>
</tr>
<tr>
<td>All other sites</td>
<td>23%</td>
<td>23%</td>
</tr>
</tbody>
</table>

*ONS=Other nervous system.

Source: American Cancer Society, 2006.
Loss of Normal Growth Control

- Cancer arises from a loss of normal growth control.
- In normal tissues, the rates of new cell growth and old cell death are kept in balance.
- In cancer, this balance is disrupted.
  - This disruption can result from uncontrolled cell growth or loss of a cell's ability to undergo "apoptosis." Apoptosis, or "cell suicide," is the mechanism by which old or damaged cells normally self-destruct.
The Beginning of Cancerous Growth

- During the development of skin cancer, the normal balance between cell division and cell loss is disrupted.
- The basal cells now divide faster than is needed to replenish the cells being shed from the surface of the skin.
- Each time one of these basal cells divides, the two newly formed cells will often retain the capacity to divide, thereby leading to an increase in the total number of dividing cells.
Tumors (Neoplasms)

- This gradual increase in the number of dividing cells creates a growing mass of tissue called a "tumor" or "neoplasm."
- If the rate of cell division is relatively rapid, and no "suicide" signals are in place to trigger cell death, the tumor will grow quickly in size; if the cells divide more slowly, tumor growth will be slower.
- But regardless of the growth rate, tumors ultimately increase in size because new cells are being produced in greater numbers than needed.
- As more and more of these dividing cells accumulate, the normal organization of the tissue gradually becomes disrupted.
Cancers are capable of spreading through the body by two mechanisms: invasion and metastasis.

- Invasion refers to the direct migration and penetration by cancer cells into neighboring tissues.
- Metastasis refers to the ability of cancer cells to penetrate into lymphatic and blood vessels, circulate through the bloodstream, and then invade normal tissues elsewhere in the body.
Malignant versus Benign Tumors

- Depending on whether or not they can spread by invasion and metastasis, tumors are classified as being either benign or malignant.
- Benign tumors are tumors that cannot spread by invasion or metastasis; hence they only grow locally.
- Malignant tumors are tumors that are capable of spreading by invasion and metastasis.
- By definition, the term "cancer" applies only to malignant tumors.
Cell Cycle

M phase
- Mitosis
- Chromosome separation
- Cell division

G₂ phase
- Preparation for mitosis

S phase
- DNA replication
- Histone synthesis
- Centrosome formed
- Chromosome duplication

G₁ phase
- RNA and protein synthesis
- Cell growth

G₀ phase
- No cell division
- Restriction point

周期
- 0h
- 4h
- 12h
- 8h
Cell Cycle Regulation

- Cyclin-dependent protein kinases
  - CDK 1 - 6
- Cyclin
  - Cyclins A - E

**Interphase**
- Histone H1
- Laminin
- Protein kinases
- Transcription factors
- Other proteins

**Mitosis**
- Spindle formation
- Chromosome condensation
- Disappearance of nuclear membrane
- Transcription stop
- Cyclin degradation

**MPF**
- Active protein kinase

**Cyclin-dependent protein kinase (CDK1)**
- Catalytic subunit
- Regulatory subunit
- Cyclin B
- Cyclin fragments
- Proteolysis

**Phosphoprotein phosphatase**
- Protein kinase

**1**
- Phosphoprotein phosphatase
- Protein kinase

**2**
- Phosphoprotein phosphatase
- Protein kinase
Cell Cycle

- **G0**
  - 6-8 h
  - DNA, RNA, Protein

- **G1**
  - 6-12 h
  - RNA, Protein

- **S**
  - 3-4 h
  - RNA, Protein

- **G2**
  - 1 h
  - RNA, Protein

- **M**
  - 1 h
  - RNA, Protein

- **Mitosis, Cytokinesis**

**Key Proteins and Cyclins**

- **Cyc B/A**
  - CDK1

- **Cyc A**
  - CDK2

- **Cyc E**
  - CDK2

- **Cyc D’s**
  - CDK4,6

- **p53**
- **pRb**

- **Lamin H1 Abl**

- **Eric Niederhoffer SIU-SOM**
Variation in Cell Cycle Cyclins

Cell cycle phases

Cyclin-dependent kinases

Cdk4

Cdk2

Cdk1

cyclins

M

G₁

S

G₂

M

G₁

Start

D

E

A

B(A)
Cell Cycle Regulation

1. CDK phosphorylation
2. C degradation
3. C & CDK synthesis
4. CDK inhibition

DNA damage → Active p53

CDK2 → p21 → pRb

pRb → Enzymes for DNA synthesis → Passage from G1 to S

E2F → pRb

p21 → CDK inhibition
Oncogene products - biochemical functions

Diagram showing the relationship between tumor formation and initiation, transformation, and normal growth and differentiation.

Key elements:
- Tumor virus
- v-Oncogene
- Reverse transcription incorporation
- Oncogene
- Proto-oncogene
- Tumor suppressor gene
- Control protein (e.g., p53 protein, Rb protein)
- Defective control
- Altered proteins
- Defective suppressor gene
- Mutation, deletion, amplification, altered control
- Normal growth and differentiation

The diagram illustrates how oncogene products play a role in tumor formation and initiation, highlighting the impact on cell growth and differentiation.
The body’s cells are normally subject to strict "social" control. They only divide until they come into contact with neighboring cells; cell division then ceases due to contact inhibition.

Exceptions to this rule include embryonic cells, cells of the intestinal epithelium (where the cells are constantly being replaced), cells in the bone marrow (where formation of blood cells takes place), and tumor cells.

Uncontrolled cell proliferation is an important indicator of the presence of a tumor.

While normal cells in cell culture only divide 20–60 times, tumor cells are potentially immortal and are not subject to contact inhibition. In medicine, a distinction is made between.
Transformation

Tumor initiators
- Viruses
- Carcinogenic chemicals
- Physical processes (UV, radioactivity)

Tumor promoters
- e.g. Esters of phorbol
- Hormones

Tumor markers (examples)
- Tumor-associated antigens:
  - CEA: Carcinoembryonic antigen
  - AFP: α1-Fetoprotein
  - Hormones: Calcitonin, ACTH
  - Enzymes: Acid phosphatase

Tumor initiation: Genetic damage
- Indicators:
  - Differentiated
  - Non-dividing
  - Defined form

Tumor progression: Preferential propagation
- Indicators:
  - De-differentiated
  - Uncontrolled cell division
  - Altered cell surface
  - Altered cytoskeleton and nucleus

Tumor progression: Acquisition of malignancy
- Indicators:
  - De-differentiated
  - Uncontrolled cell division
  - Altered cell surface
  - Altered cytoskeleton and nucleus
Molecular Biology of Cancer

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Protein Data Bank, SDSC
The development and metastasis of human colorectal cancer and its genetic basis.

- A mutation in the APC tumor-suppressor gene in a single epithelial cell causes the cell to divide, although surrounding cells do not, forming a mass of localized benign tumor cells, or polyp.
- Subsequent mutations leading to expression of a constitutively active Ras protein and loss of two tumor-suppressor genes—a unidentified gene in the vicinity of DCC and p53—generate a malignant cell carrying all four mutations. This cell continues to divide, and the progeny invade the basal lamina that surrounds the tissue.
- Some tumor cells spread into blood vessels that will distribute them to other sites in the body.
- Additional mutations permit the tumor cells to exit from the blood vessels and proliferate at distant sites; a patient with such a tumor is said to have cancer.

[Adapted from B. Vogelstein and K. Kinzler, 1993, Trends Genet. 9:101.]
The APC Gene:
Representation of APC protein domains with respect to mutational analysis results

- The relative positions of various APC domains:
  - involved in homo-oligomerization of APC (N-terminus).
  - series of repeats of unknown function (similarity to the Drosophila armadillo protein),
  - mediate binding to β-catenin and its downregulation,
  - basic domain in the C-terminal third of the protein – facilitate complexing with microtubules (MT),
  - sequences near the C-terminus of APC – interact with the EB1 and human homolog of the Drosophila disc large (hDlg) protein.

Somatic mutations in the APC gene in colorectal cancer appear to cluster in a region termed the "mutation cluster region," and mutations at codons 1309 and 1450 are most common.

The Function of the APC, Axin, and GSK3β Proteins in the Regulation of β-catenin
Possible fates for carcinogen-damaged DNA

- Carcinogen damage leading to altered DNA
  - Efficient repair
  - Incorrect repair/ altered primary sequence
  - Apoptosis
    - Normal cell
    - DNA replication & cell division: fixed mutations
    - Cell death
      - Transcription & translation giving aberrant proteins
      - Carcinogenesis if critical targets are mutated: oncogenes, tumour-suppressor genes
Alkylating agents, anthracyclines

- Cross-linking of DNA components
- "Bending" of the DNA double helix

- Cyclophosphamide
- Adriamycin
- Cisplatin
Antimetabolites-2

1. Hypoxanthine phosphoribosyltransferase 2.4.2.8
2. Thiopurine methyltransferase 2.1.1.67
3. Amidophosphoribosyltransferase 2.4.2.14
4. Ribonucleoside diphosphate reductase 1.17.4.1

5. 5-fluorodeoxyuridine monophosphate
6. Methotrexate (amethopterin)

5-fluorouracil
5-fluorodeoxyuridine

Precursors

N5,N10-methylene-THF

Dihydrofolate

DNA

dTMP

dTTP

Thymidylate synthase 2.1.1.45
Dihydrofolate reductase 1.5.1.3

Hydroxyurea
Some Links for Cancerogene Researches

- www.pubmed.org
- http://atlasgeneticsoncology.org/index.html
- http://www.genecards.org/index.shtml
- http://www.ebi.ac.uk/msd
Thank you for your attention!