

Nomenclature

Sarcomas	Mesenchymal tumors
Carcinomas	Epithelial tumors

Epidemiology (Acquired predisposing conditions)

1. Chronic inflammation
2. Immunodeficiency states
3. Precursor lesion

Clinical aspects

1. Local effects of tumor encroachment of tissues/ organs
- 2a. Functional activity e.g. Hormone synthesis
- 2b. Paraneoplastic syndromes -> Ectopic hormone secretion
3. Bleeding & infections when tumor ulcerates thru adjacent surface
4. Rupture/ infarction
5. Cachexia (Weakness, e.g. weight loss)

Molecular basis of cancer

Nonlethal genetic damage
Hallmark: Genetic alteration

Cancer genes (Target of genetic damage)

1. **Oncogene (Mutated gene)**
 - Mutation from proto-oncogenes
2. **Tumor suppressor genes**
 - Prevent uncontrolled growth
3. **Apoptosis-regulating gene**
 - Overexpressed in cancer cell-> Protect against apoptosis
4. **Regulate interactions between tumor and host cells**
 - Change recognition of tumor by host immune system

2. Insensitivity to Tumor suppressor signals

Retinoblastoma Gene

- Active hypophosphorylated state: Halts cell cycle
- Inactive hyperphosphorylated state
- **Heterozygosity: X Affect cell function**
- **Both to be inactivated to affect function**

p53

1. Cell cycle arrest
2. DNA repair

2. Insensitivity to Tumor suppressor signals (cont)

3. Apoptosis

4. Evasion of cell death

- Overexpression of BCL-2 protein -> Long life

8. Evasion of immune surveillance

Host defence against tumor -- Tumor immune

Tumor antigens

Antitumor effectors

- | | |
|---|--------------------|
| - Overexpressed cellular proteins, Oncogenic viral products, Differentiation antigens | - CD8+ |
| - Oncogenic viral products | - NK lymphocytes |
| - Differentiation antigens | - Macrophages |
| | - Humoral immunity |

Immune evasion

- | | |
|---------------------|-----------------------|
| - Immunosuppression | - Selective outgrowth |
| - Antigen masking | - X MHC expression |
| - Apoptosis of CD8+ | - X Costimulation |

10. Tumor-promoting inflammation

- Interaction between inflammatory cell& tumor
1. Proliferation-promoting factor release
 2. Growth suppressor removal
 3. Cell death resistance
 4. Angiogenesis
 5. Invasion & Metastasis
 6. Immune evasion

Benign Malignant differentiation

Benign Malignant

Differentiation & anaplasia

- | | |
|-------------------------------|---|
| 1. Well differentiated | 1. Well to undifferentiated (Anaplasia: Functional& structural differentiation loss) |
| | - Dysplasia (Disordered growth) |

Benign Malignant differentiation (cont)

- Carcinoma in situ (Non-invasive malignant tumor)

Rate of growth

- Correlates w./ level of differentiation

2. Progressive & slow **2. Erratic** (Unpredictable)

Local invasion

3. No, expansion w./ clear boundaries **4. Yes**, infiltrate & destroy

Metastasis

(1) Seeding of body cavities (2) Lymphatic spread (3) Hematogenous spread

4. Absent **4. Frequently present**

1. Self-sufficiency in growth signals

Proto-oncogenes - Normal genes, promote proliferation

Oncogenes - Mutant version, function anonymously w./o growth-promoting signals

Oncoproteins - Proteins encoded

Self-sufficient in:

- 1. Growth factors & receptors**
- 2. Signal transduction proteins**
- 3. Transcription factors**
- 4. Cyclins & CDKs**

3. Altered cellular metabolism

Warburg effect

- Aerobic situation: Distinct form of cellular metabolism
- **High levels of glucose uptake**
- **Increased conversion of glucose to lactose via glycolytic pathway**

5. Limitless replicative potential: Telomerase

- Telomerase shorten with each cell division
- Cancer cell have enzyme that regenerate telomerase

6. Sustained angiogenesis

- Controlled by balance between angiogenesis promoter (VEGF) and inhibitors (bFGF)

7. Invasion & Metastasis

- Invasion of extracellular matrix

a. Loosening of intracellular junctions

b. Degradation

c. Attachment

d. Migration

- Embolus: Evade WBC killing

9. Genetic instability

- Both copies of DNA repair proteins are lost

1. Hereditary Nonpolyposis Cancer Syndrome

2. BRCA-1 & BRCA-2 (80% familial breast cancer, not sporadic-associated)

Carcinogenic Agents

1. Chemical Carcinogenesis

Initiation

- Carcinogen exposure -> permanent DNA

Promotion

- Promoter induce tumor in initiated cell (Nontumorigenic)
- Promoting agent enhance proliferation & results in cancer

2. Radiation Carcinogenesis

- UV rays (UVB, 280-320nm)
- Ionizing radiation (X-ray, gamma ray, particles)

3. Oncogenic DNA viruses

- 1. Papillomaviruses (HPV)**
- 2. Epstein-Barr virus (EBV)**
- 3. Hep B virus (HBV)**
- 4. Kaposi sarcoma herpes virus (KSHV)**

