



Pathophysiology introduction

BY DR. SWATHI SWAROOPA B

▶ Pathology is literally the study (logos) of suffering (pathos).

▶ Pathology: scientific study of changes in the structure and function of the body in disease.



▶ General: concerned with basic reactions of cell and tissue to abnormal stimuli

▶ Systemic: concerned with specific organs and tissues to more or less well-defined stimuli



Pathophysiology: study of disordered function (physiological changes) and breakdown of homeostasis in diseases (biochemical changes)

Four aspects of a disease process that form the core of pathology

- ▶ Etiology
- ▶ Pathogenesis
- ▶ Morphological changes
- ▶ Clinical significance



2. Pathogenesis: sequence of events in response of cell/tissue to etiological agent
3. Morphologic changes: structural alterations in cells or tissues characteristic of disease.
4. Clinical manifestations: morphologic changes influence normal function and determine the clinical features (symptoms and signs)

Basic principles of cell injury and adaptation

Homeostasis

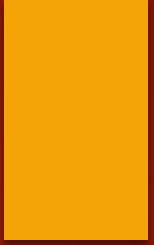
Homeostasis

- ▶ Homeostasis (ho⁻ me⁻ -o⁻ -STA⁻ -sis; homeo- **sameness**; -stasis standing still) is the condition of equilibrium (balance) in the body's internal environment due to the constant interaction of the body's many regulatory processes
- ▶ Homeostasis is **a dynamic** condition.
- ▶ In response to changing conditions, the body's equilibrium can shift among points in a **narrow range** that is compatible with maintaining life

- 
- ▶ An important aspect of homeostasis is **maintaining the volume and composition** of body fluids.

Control of Homeostasis

- ▶ Disruptions in homeostasis come from the external environment in the form of **physical insults**, from **internal environment** and from **psychological stresses**.
- ▶ Most cases the disruption of homeostasis is **mild and temporary**, and the responses of body cells quickly restore balance in the internal environment

- 
- ▶ Some cases the disruption of homeostasis may be **intense and prolonged**
 - ▶ The body has many **regulating systems** that can usually bring the internal environment back into balance
 - ▶ Mainly the **nervous system** and the **endocrine system**

Feedback Systems

A feedback system or feedback loop is a **cycle of events** in which the status of a body condition is

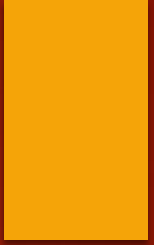
- ▶ Monitored,
 - ▶ Evaluated,
 - ▶ Changed,
 - ▶ Remonitored,
 - ▶ Reevaluated, and so on.
-
- ▶ Each **monitored variable**, such as body temperature, blood pressure, or blood glucose level, is termed a **controlled condition**.

- 
- ▶ Any **disruption** that changes a controlled condition is called a **stimulus**

Feedback system includes three basic components—

- ▶ Receptor,
- ▶ Control center, and
- ▶ Effector

- ▶ **Receptor**: is a body structure that **monitors changes** in a controlled condition and sends input to a control center
- ▶ **Input** is in the form of nerve impulses or chemical signals
- ▶ **Control center** (in body): **sets the range of values** within which a controlled condition should be maintained, evaluates the input it receives from receptors, and **generates output** commands when they are needed
- ▶ Output from the control center typically occurs as nerve impulses, or hormones or other chemical signals

- 
- ▶ **Effector**: is a body structure that **receives output** from the control center and **produces a response** or effect that changes the controlled condition.
 - ▶ Nearly every organ or tissue in the body can behave as an effector

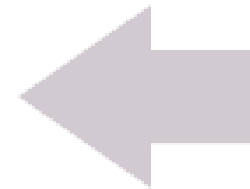
Some stimulus
disrupts homeostasis by

Increasing or
decreasing a

Controlled condition
that is monitored by

Receptors

that send



Receptors

that send

Input

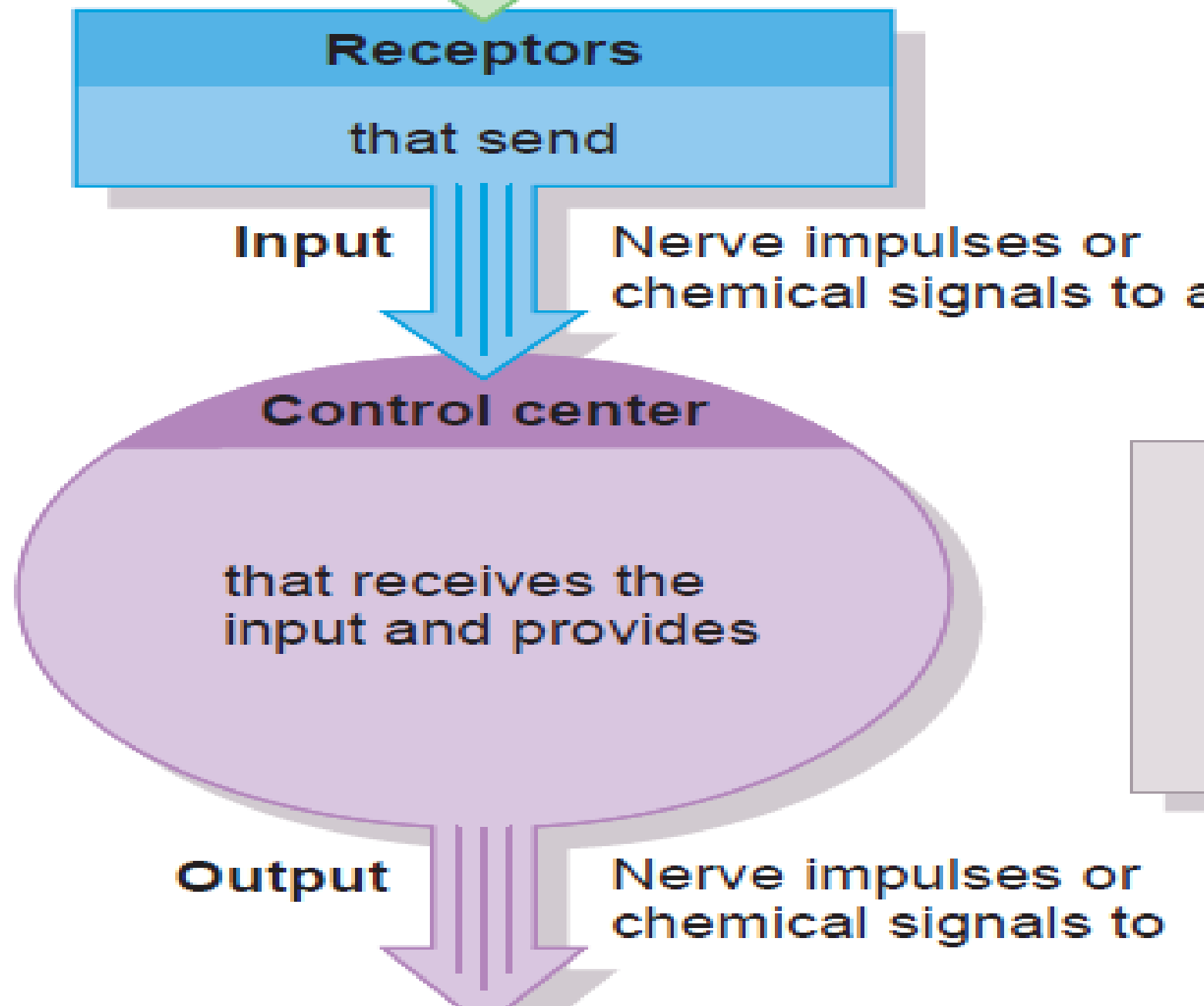
Nerve impulses or
chemical signals to a

Control center

that receives the
input and provides

Output

Nerve impulses or
chemical signals to



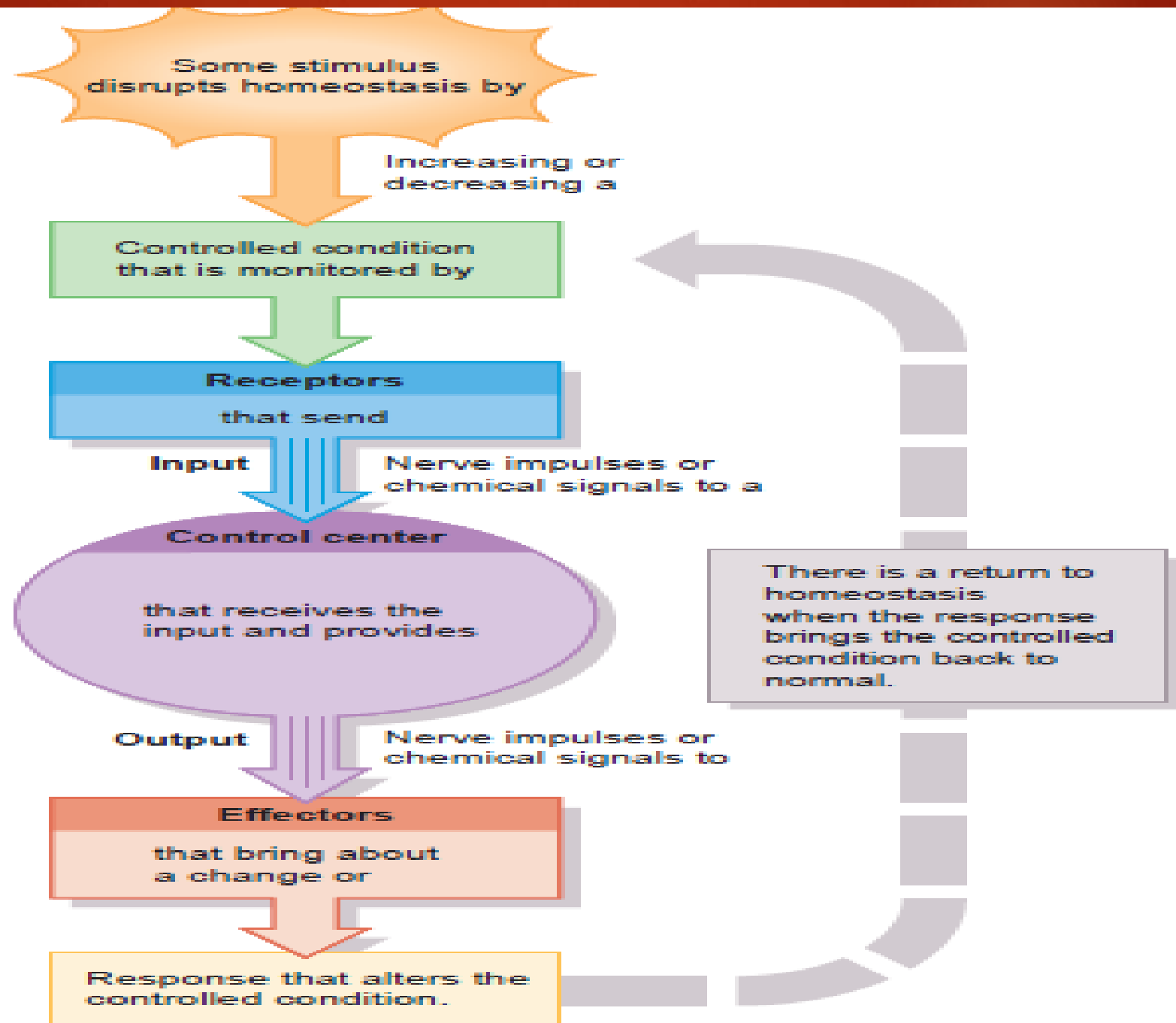
Output

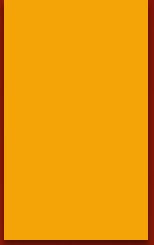
Nerve impulses or
chemical signals to

Effectors

that bring about
a change or

Response that alters the
controlled condition.



- 
- ▶ A group of **receptors and effectors** communicating with their control center **forms a feedback system** that can regulate a controlled condition in the body's internal environment.
 - ▶ In a feedback system, the response of the system **“feeds back”** information to change the controlled condition in some way, **either negating it (negative feedback) or enhancing it (positive feedback).**

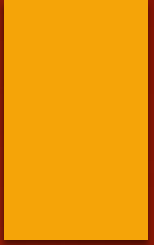
Types of feed back system

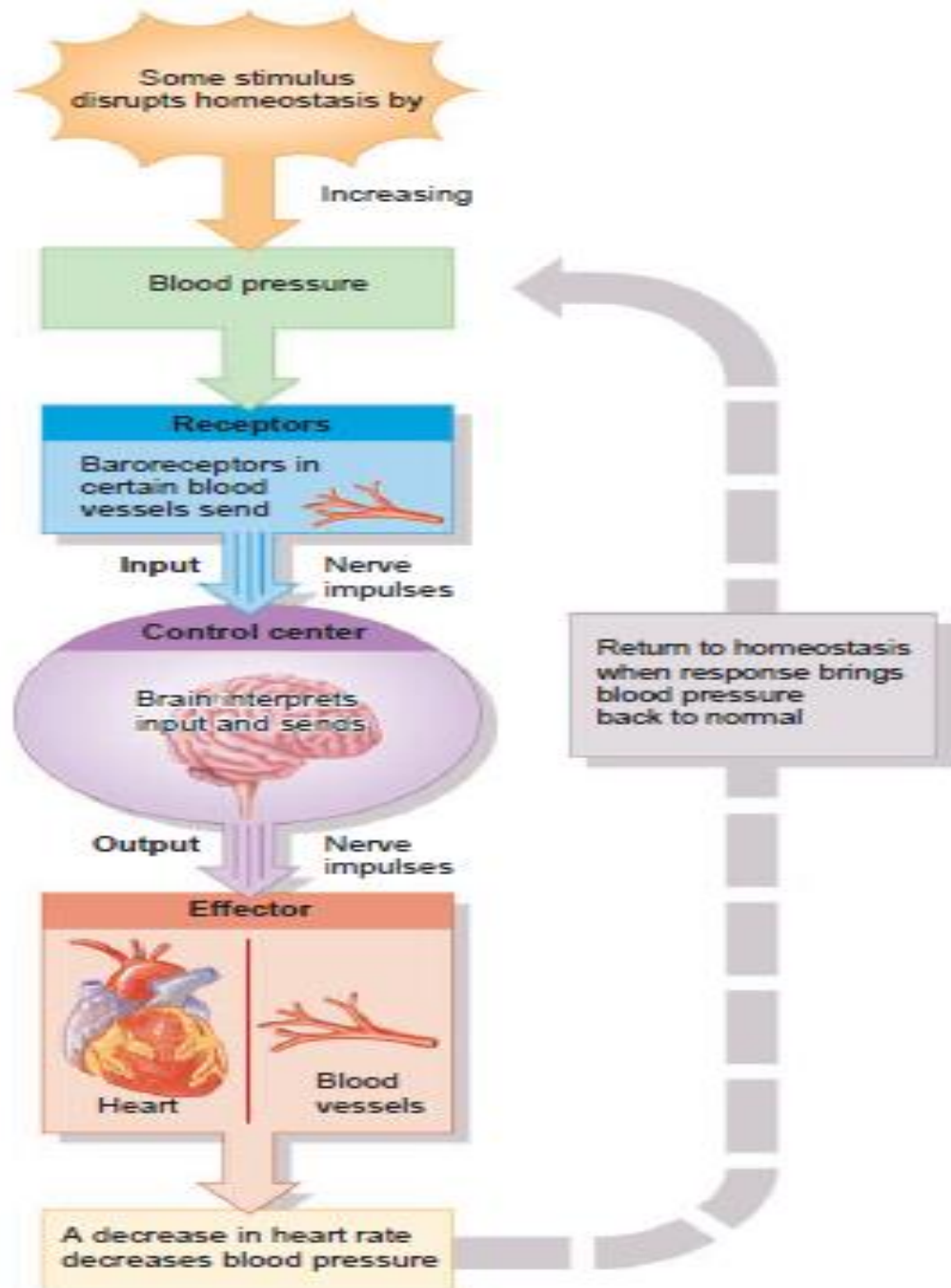
NEGATIVE FEEDBACK SYSTEMS:

- ▶ A negative feedback system **reverses a change** in a controlled condition.

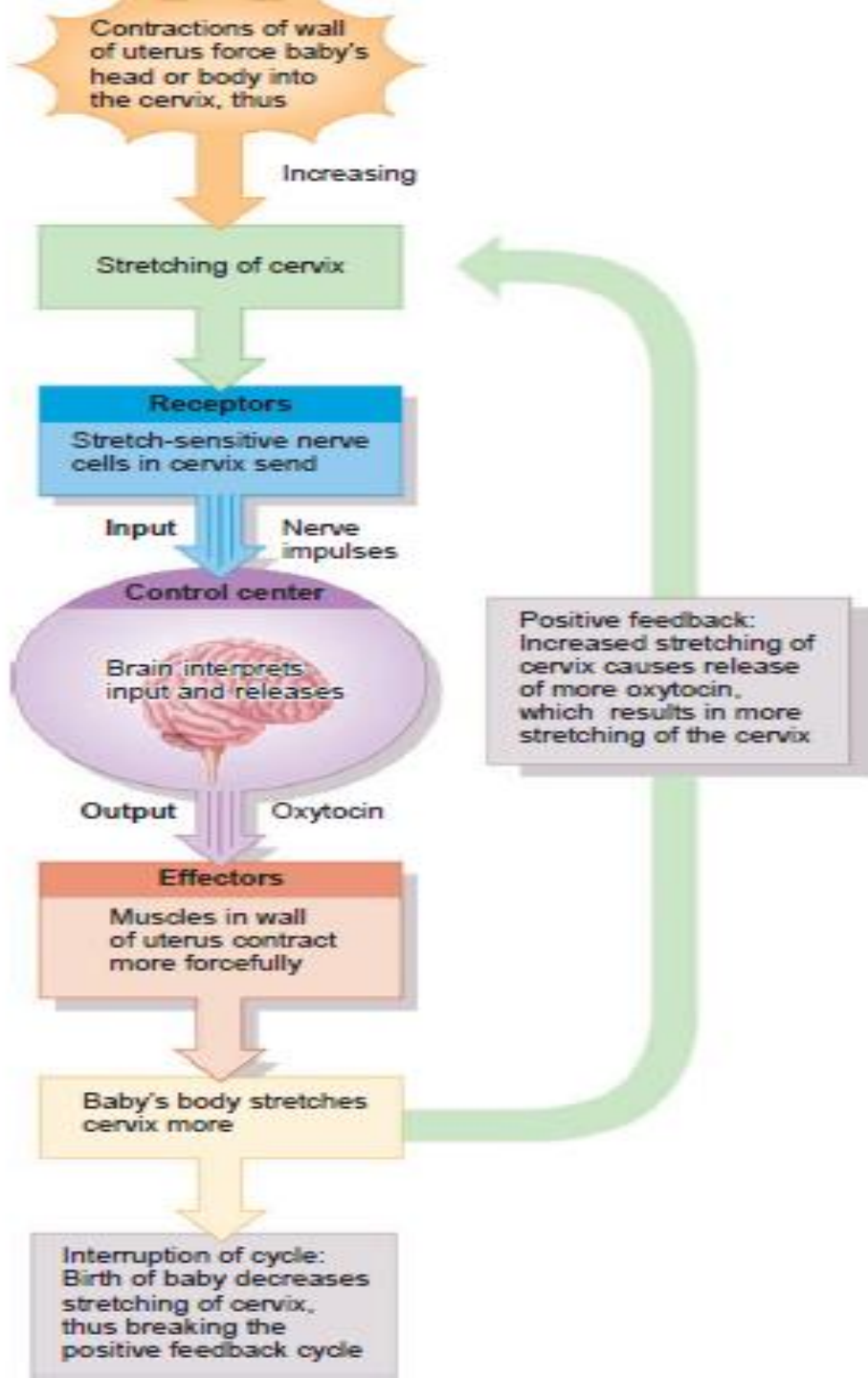
POSITIVE FEEDBACK SYSTEMS:

- ▶ The feedback is used to **increase the size of the input**. By nature, such systems are unstable, and they are most often associated with pathological conditions.
- ▶
- ▶ A positive feedback system tends to **strengthen or reinforce** a change in one of the body's **controlled conditions**.

- 
- ▶ A positive feedback system operates similarly to a negative feedback system, except for the way **the response affects the controlled condition**
 - ▶ The control center still provides commands to an effector, but this time the effector produces a physiological response that adds to or **reinforces the initial change in the controlled condition.**
 - ▶ The action of a positive feedback system **continues until it is interrupted by some mechanism**



Homeostatic regulation of blood pressure by a negative feedback system.



Positive feedback control of labor contractions during birth of a baby.

Difference between positive and negative feedback systems

POSITIVE

a positive feedback system continually reinforces a change in a controlled condition, some event outside the system must shut it off.

If the action of a positive feedback system is not stopped, it can “run away” and may even produce life-threatening conditions in the body.

Positive feedback systems reinforce conditions that do not happen very often.

NEGATIVE

The action of a negative feedback system, by contrast, slows and then stops as the controlled condition returns to its normal state.

Negative feedback systems regulate conditions in the body that remain fairly stable over long periods

Cell responses to stress

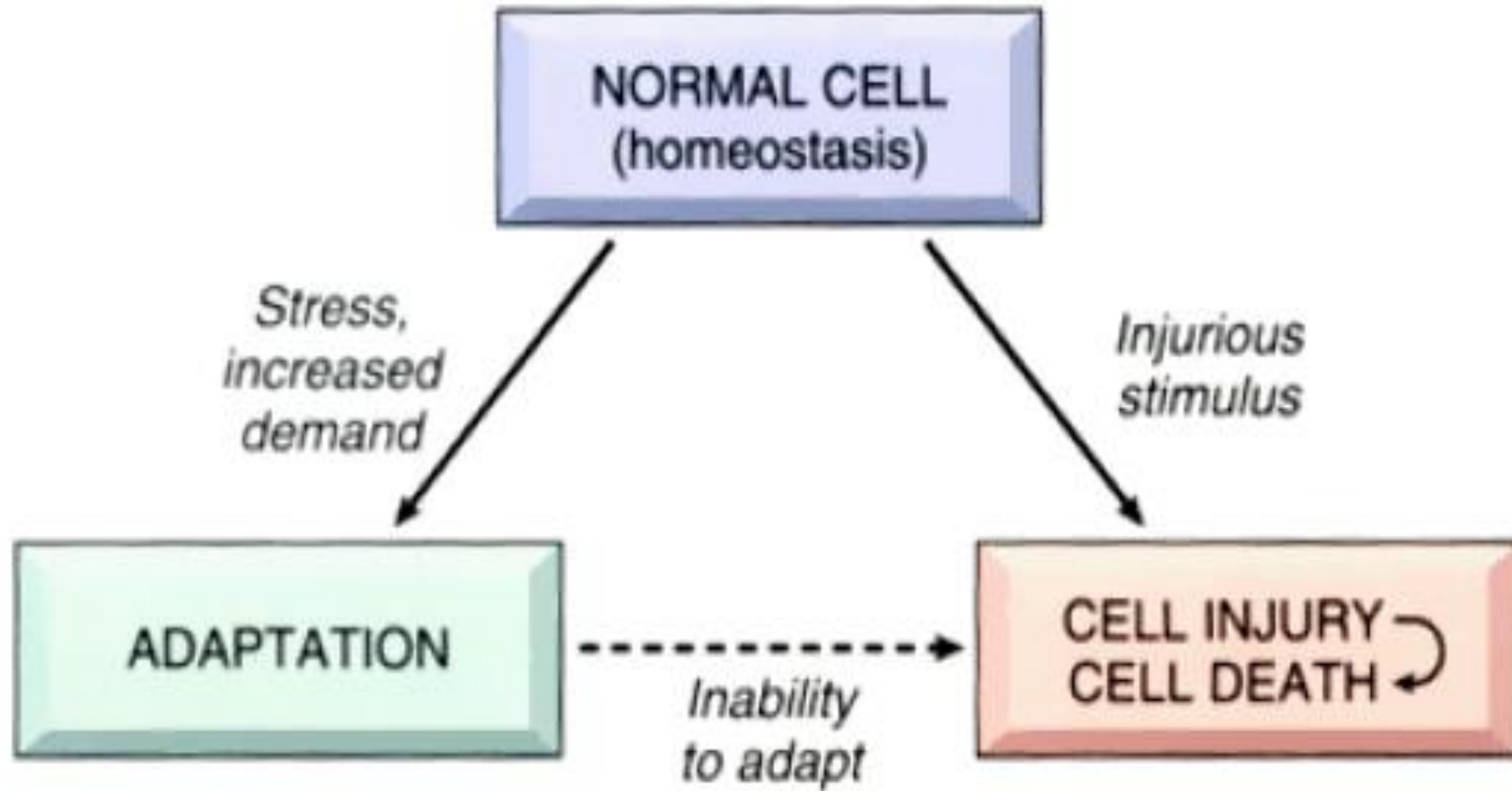
- ▶ Severe **physiologic stresses** and some **pathologic stimuli** may bring about a number of physiologic and morphologic cellular adaptations
- ▶ A new but **altered steady states are achieved**, preserving the viability of the cell and modulating its function as it responds to such stimuli
- ▶ In certain pathologic conditions, when cells are **damaged beyond repair**, and especially if the damage affects the **cell's nuclear DNA**

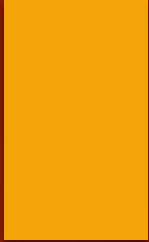
Cell responses to stress

- ▶ Exposed to **sublethal or chronic stimuli** may not be damaged but may show a variety of subcellular alterations.
- ▶ **Metabolic derangements** in cells may be associated with intracellular accumulations

Nature and Severity of Injurious Stimulus	Cellular Response
Altered physiologic stimuli:	Cellular adaptations:
• Increased demand, increased trophic stimulation (e.g. growth factors, hormones)	• Hyperplasia, hypertrophy
• Decreased nutrients, stimulation	• Atrophy
• Chronic irritation (chemical or physical)	• Metaplasia
Reduced oxygen supply; chemical injury; microbial infection	Cell injury:
• Acute and self-limited	• Acute reversible injury
• Progressive and severe (including DNA damage)	• Irreversible injury → cell death
	•••• Necrosis
	•••• Apoptosis
• Mild chronic injury	• Subcellular alterations in various organelles
Metabolic alterations, genetic or acquired	Intracellular accumulations; calcifications
Prolonged life span with cumulative sublethal injury	Cellular aging

Cell responses to stress



- 
- a) Adaptation
 - b) Cell injury
 - c) Subcellular alterations
 - d) Intracellular accumulations
 - e) Calcification
 - f) Cell aging

Adaptations induced by

1. Factors produced by the responding **cells themselves**
2. Factors produced by **other cells** in the environment
3. Activation of various **cell surface receptors** and downstream signaling pathways

Adaptations associated with

1. Induction of new protein synthesis by the target cells
2. The induction of cellular proliferation
3. Shifting cells from producing one type of proteins to another
4. Markedly overproducing one protein

Types of adaptations

- ▶ **Hyperplasia**: increase in number of cells in an organ or tissue (increase volume of organ)
- ▶ **Hypertrophy**: increase in size of cells, resulting in an increase in size of organ
- ▶ **Atrophy**: shrinkage in size of cells
- ▶ **Metaplasia**: reversible change of one adult cell type(epithelial /mesenchymal) is replaced by another cell type.
- ▶ **Dysplasia**: disordered cellular development (also called atypical hyperplasia)

Hyperplasia

► Hyperplasia

Physiologic

Pathologic

Hormonal

Compensatory

Mechanism-

- Increased local production of growth factors
- Increased levels of growth factor receptor

Activation of
particular
intracellular
signaling pathways

Production of
transcription
factors

Turn on many
cellular genes that
encode growth
factors

Cellular
proliferation

Hyperplasia classification

Physiologic

Hormonal

Occurs with influence of hormones

e.g. Hyperplasia of female breast at puberty, pregnancy and lactation

Hyperplasia of pregnant uterus

Compensatory

Hyperplasia occurs due to removal or damage of part of an organ

e.g. Regeneration of liver following partial hepatectomy

Regeneration of epidermis after skin abrasion

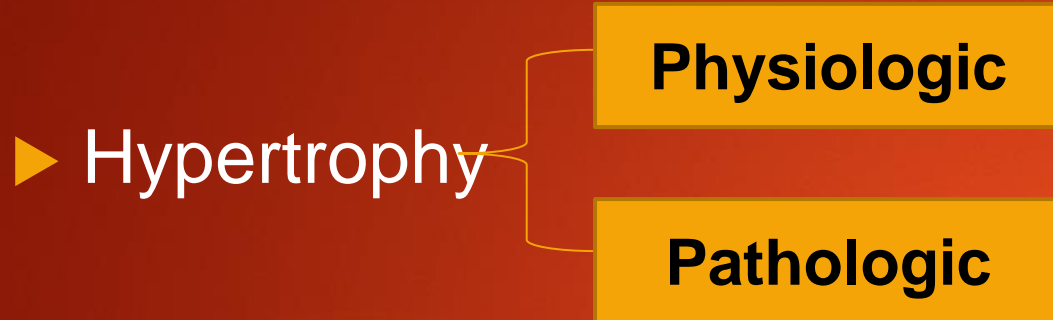
Pathologic

Due to excessive hormonal stimulation and growth factors acting on target cells

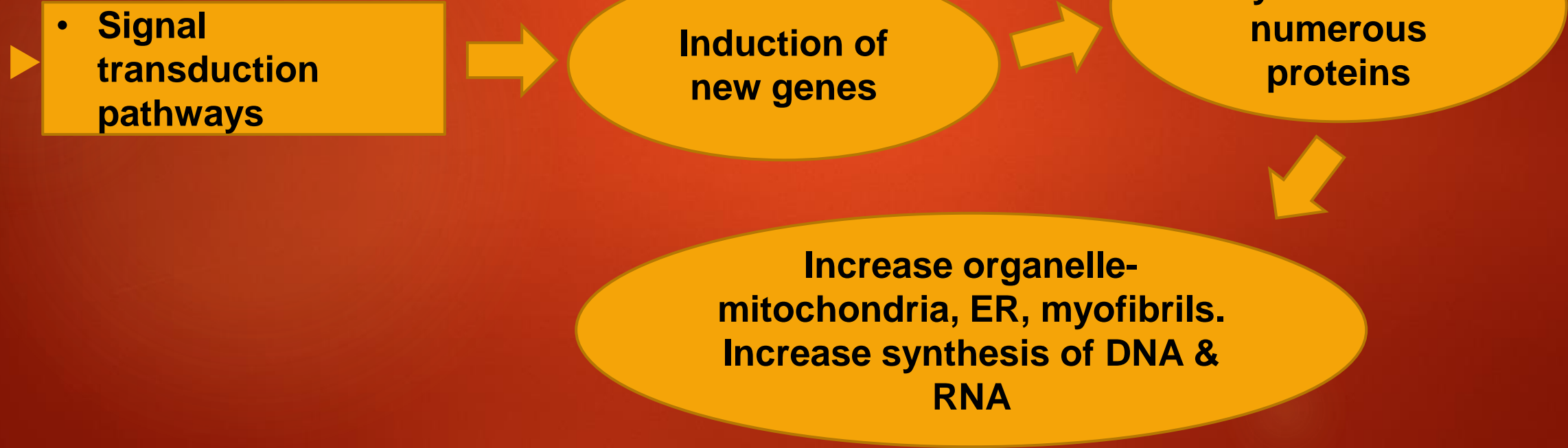
e.g. Endometrial hyperplasia following oestrogen excess

Formation of skin warts or lesions (hyperplasia of epidermis) due to papilloma virus

Hypertrophy



Mechanism-1.



2.

- **Early development genes re-expressed in hypertrophic cells**



Product of these genes participate in cellular response to stress

Genes that are induced during hypertrophy include-

1. Encoding transcription factors
2. Growth factors
3. Vasoactive substances

Hypertrophy classification

Physiologic	Pathologic
<p>Most common stimulus is work load.</p> <p>Work load is shared by greater mass of cellular components and each muscle fiber is spared excess work &so escapes injury</p> <p>e.g. Enlarged size of uterus in pregnancy(hormone induced hypertrophy)</p> <p>Hypertrophy of breast during lactation due to prolactin and estrogen.</p>	<p>Hypertrophy of cardiac muscle due to chronic hemodynamic overload, resulting from</p> <p>Hypertension</p> <p>Aortic valve disease (stenosis)</p>

Atrophy

► Atrophy

Physiologic

**Decrease of uterus size after
parturition**

Pathologic

Depends on underlying cause

Mechanism of atrophy

Results from

- ▶ Decreased protein synthesis
- ▶ Increased protein degradation

**Due to decreased
metabolic activity**

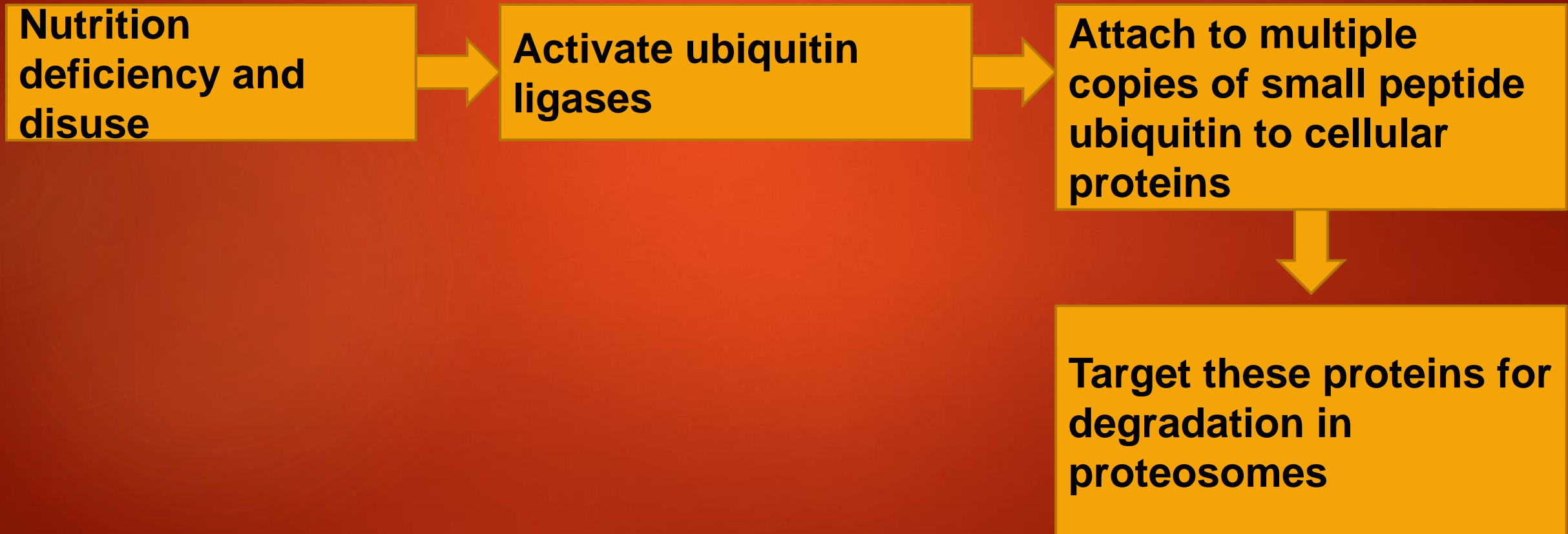


**Decreased protein
synthesis**

Mechanism of atrophy

Protein degradation

1. Ubiquitin-proteasome pathway



Mechanism of atrophy

Protein degradation

2. Increased autophagy (self eating)

Increased
autophagy



Increased autophagic
vacuoles

Autophagy: starved cells eat its own components to find nutrients and survive.

Causes of pathological atrophy

- ❑ **Disuse atrophy-wasting or muscle atrophy in complete bed rest**
- ❑ **Denervation atrophy-rapid atrophy of muscle fibers due to damage of nerve which innervated them**
- ❑ **Ischemic atrophy-atrophy due to decreased blood supply to tissue (atrophy of brain in cerebral atherosclerosis)**
- ❑ **Inadequate nutrition /starvation atrophy-muscle wasting due to malnutrition (seen in cancer and chronic inflammatory disease)**

Causes of pathological atrophy

- ❑ **Endocrine atrophy**-loss of endocrine stimulation leads to decreased metabolic activity of tissue (hypopituitarism may lead to atrophy of thyroid, adrenal & gonads)
- ❑ **Pressure atrophy**- Tissue compression for long time (enlarging benign tumor can cause atrophy in surrounding compressed tissues)
- ❑ **Idiopathic atrophy**- No obvious cause (e.g. myopathies, testicular atrophy)

Metaplasia

► Metaplasia

**Epithelial
metaplasia**

**Columnar to
squamous**

**Squamous to
columnar**

**Connective
tissue
metaplasia**

**Osseous
metaplasia**

**Cartilaginous
metaplasia**

Mechanism of Metaplasia

**Signals generated
by cytokines,
growth factors,
extracellular matrix
components**



**Reprogramming
of stem cells**



Metaplasia

Epithelial metaplasia

Columnar to squamous- In cigarette smoking ciliated epithelial cells of trachea & bronchi replaced by stratified squamous epithelial cells

In cholelithiasis-replacement of normal secretory columnar epithelium by non functioning stratified squamous epithelium.

Squamous to columnar

In barett esophagus squamous is replaced by intestine like columnar cells under the influence of refluxed gastric acid

Connective tissue metaplasia

Osseous metaplasia: Formation of cartilage, bone or adipose tissue (mesenchymal tissue) in tissues that generally don't contain them. Formation of bone in muscle in myositis ossificans.

Cartilaginous metaplasia: occurs where there is less mobility during the healing of fractures

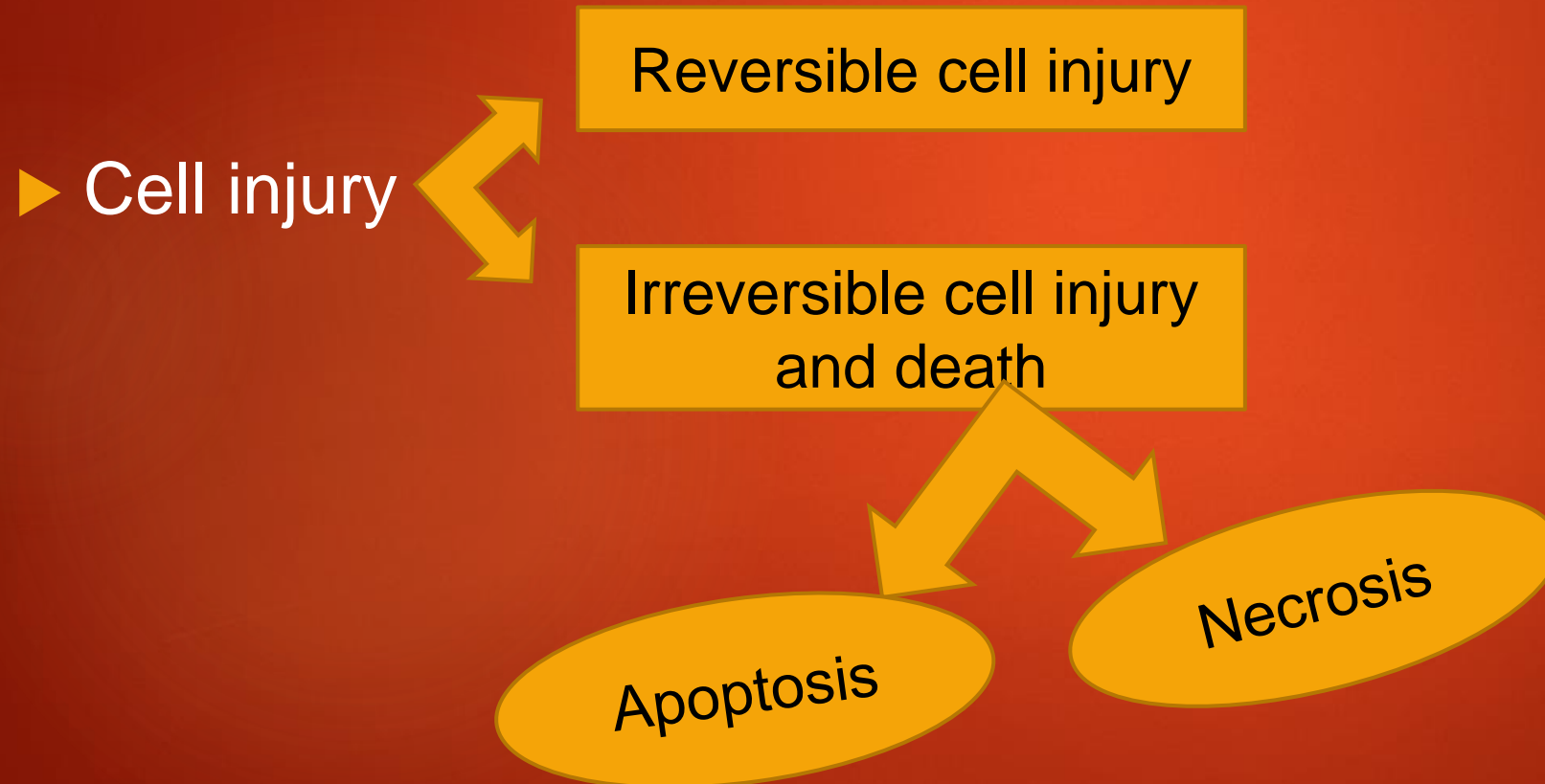
In synovial chondromatosis, cells of the synovial membrane undergo metaplasia to become cartilage-producing chondrocytes. .

Dysplasia

- ▶ Disordered cellular development also called as atypical hyperplasia
- ▶ Both metaplasia and hyperplasia are seen
- ▶ Caused by chronic irritation and prolonged inflammation.
- ▶ Epithelial dysplasia of the cervix (cervical intraepithelial neoplasia – a disorder commonly detected by an abnormal pap smear)

Cell injury and death

- ▶ Results when cells are **stressed so severely that they are no longer able to adapt** or when cells are exposed to inherently damaging agents.



Causes of cell injury

1. Oxygen deprivation



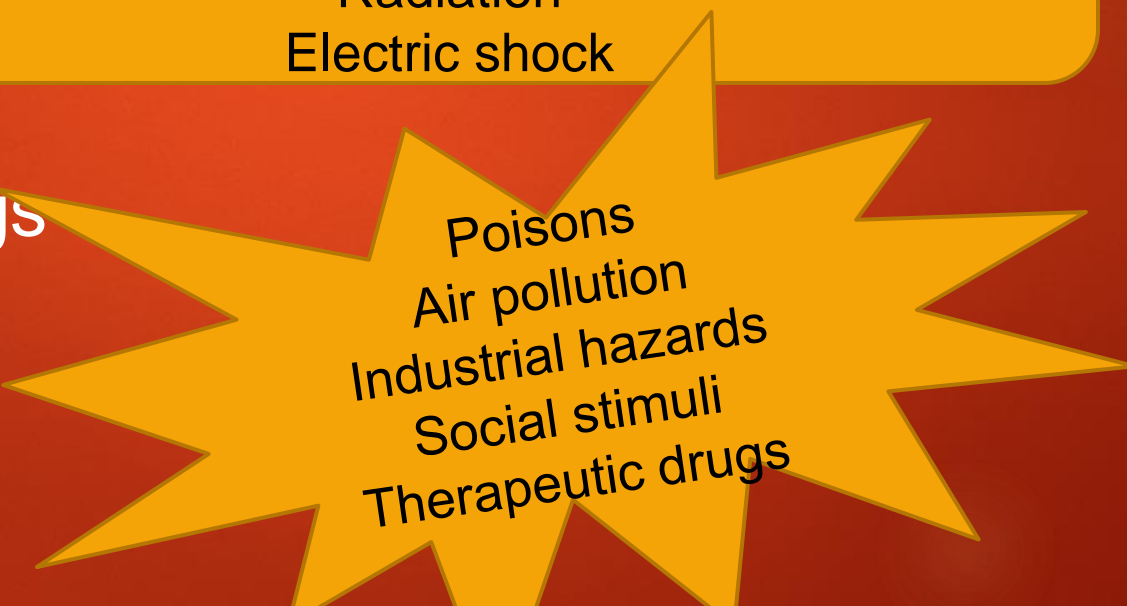
Hypoxia

Ischemia

2. Physical agents-

Mechanical trauma
Temperature extremes
Change in atmosphere pressure
Radiation
Electric shock

3. Chemical agents & drugs



Poisons
Air pollution
Industrial hazards
Social stimuli
Therapeutic drugs

4. Infectious agents & immunologic rxn's

Virus, fungi, bacteria
parasites and anaphylactic
rxn's

5. Genetic derangement

Enzymatic abnormalities
Variations in genetic make
up

6. Nutritional imbalances

Nutritional excess
Nutritional deficiency

Mechanism of cell injury

Cell response to injury depends on

- ▶ Type of injury
- ▶ Its duration
- ▶ Severity

Consequences of cell injury depends on

- ▶ Type
- ▶ State
- ▶ Adaptability of injured cell

- 
- ▶ Cell injury results from **functional & biochemical abnormalities** in one or more of several essential **cellular components**

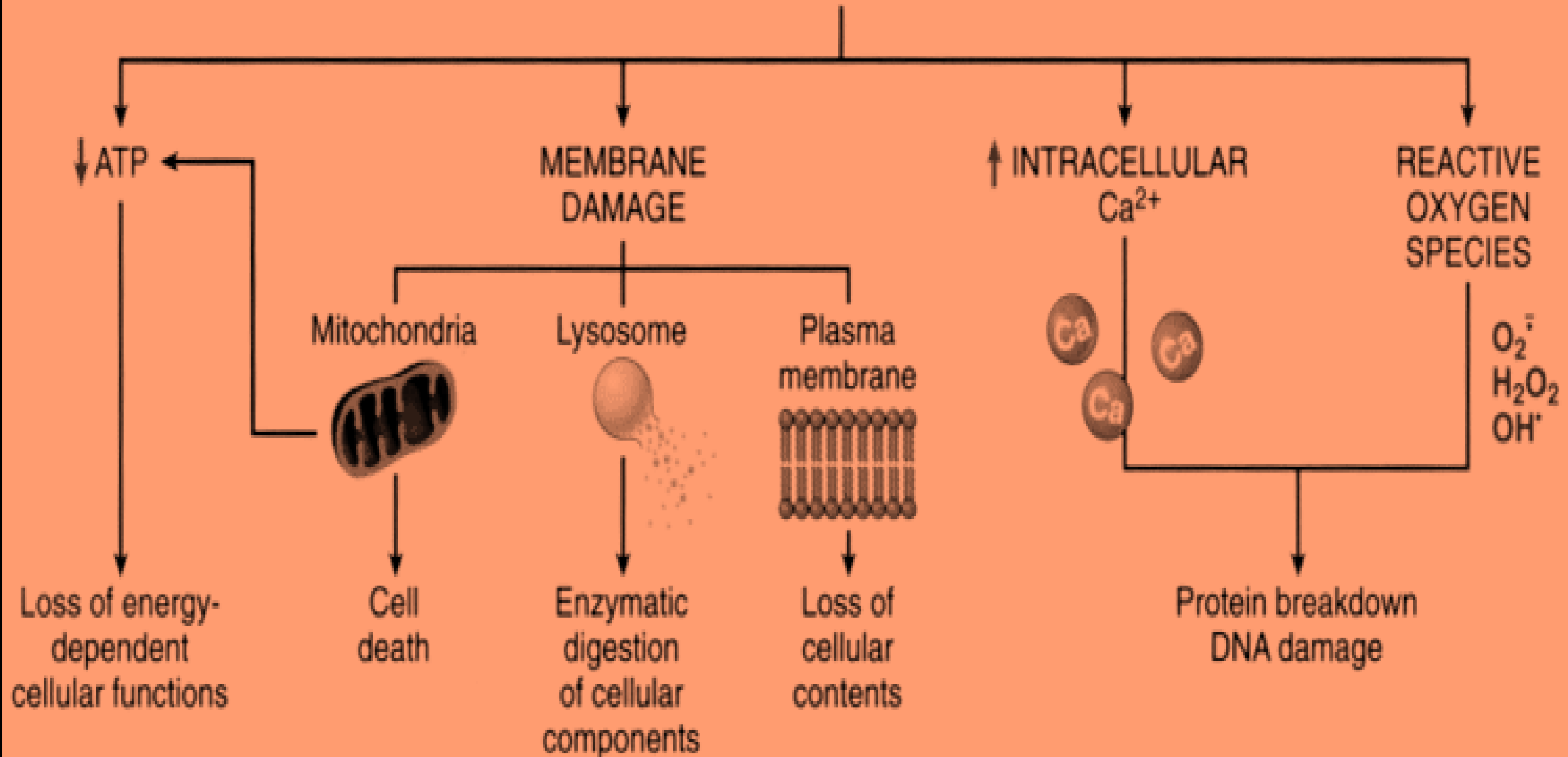
Important targets of injurious stimuli

- ▶ Aerobic respiration (oxidative phosphorylation in mitochondria)
- ▶ Cell membrane integrity
- ▶ Protein synthesis
- ▶ Cytoskeleton
- ▶ Genetic apparatus integrity

Biochemical mechanisms responsible for cell injury

- a) Depletion of ATP
- b) Mitochondrial damage
- c) Influx of intracellular Ca^{+2} & loss of Ca^{+2} homeostasis
- d) Oxidative stress
- e) Defects in membrane permeability.

INJURIOUS STIMULUS



Depletion of ATP

- ▶ ATP produced in 2 ways



Oxidative phosphorylation
Glycolytic pathway

- ▶ Depletion of ATP is associated with both hypoxia and chemical stimuli
- ▶ ATP required for many synthetic & degradation process within the cell
- ▶ It involves in –membrane transport, protein synthesis, lipogenesis, deacylation & reacylation rxns, phospholipid turnover.

Ischemia



↓ Oxidative phosphorylation



↓ ATP

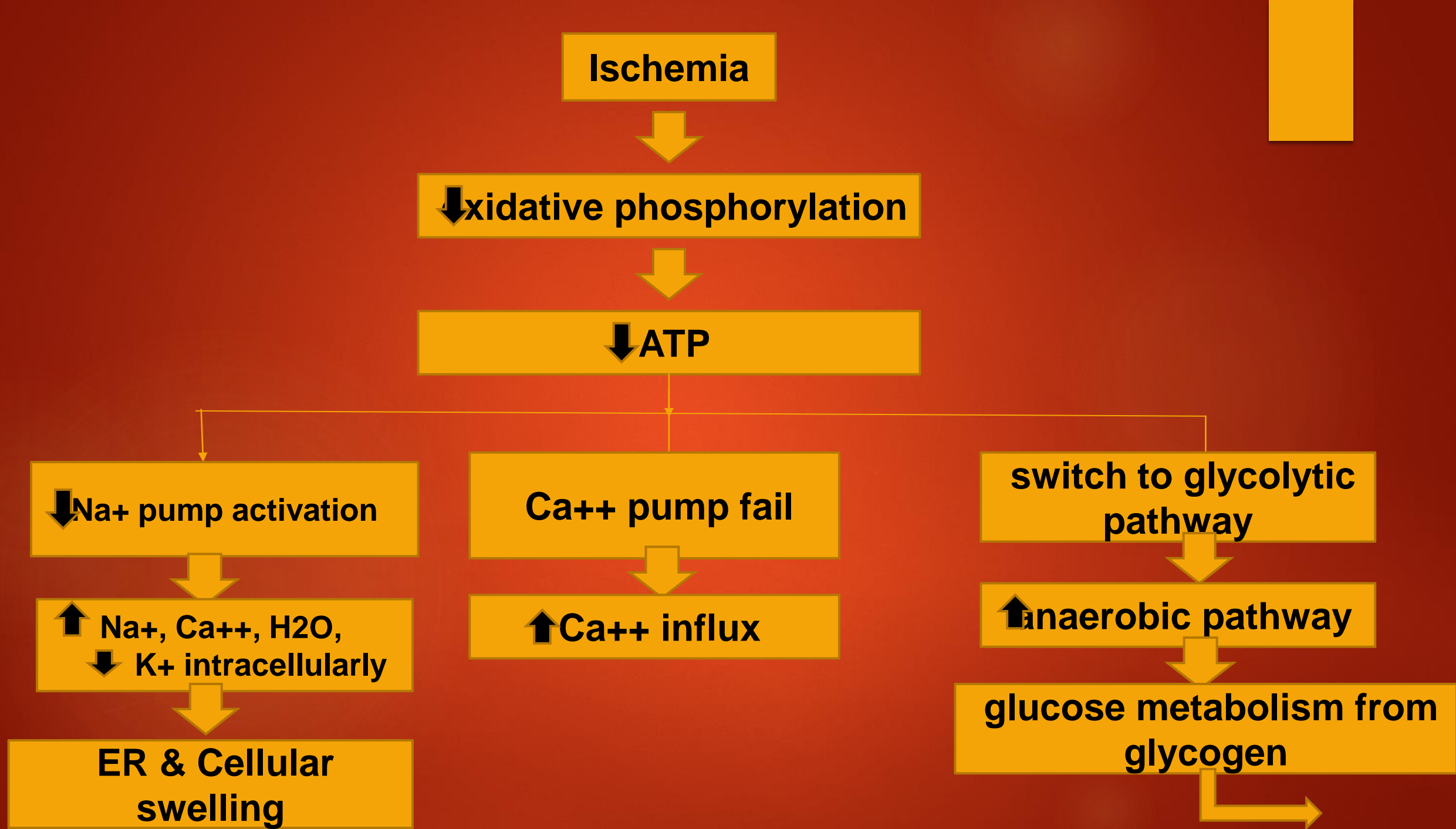
**↓ Na+ pump
activation**

**Ca++ pump
fail**

**switch to
glycolytic
pathway**

**disruption of
protein
synthesis**

**proteins
misfolded**



glucose metabolism from
glycogen



↓ glycogen(
glycolysis)



accumulation of lactic
acid & inorganic
phosphates



↓ PH



↓ activity of many cellular
enzymes & clumping of
nuclear chromatin

↓ ATP

**disruption of protein
synthesis**

**Detachment of
ribosomes from
rough ER &
dissociation of
polysomes to
monosomes**

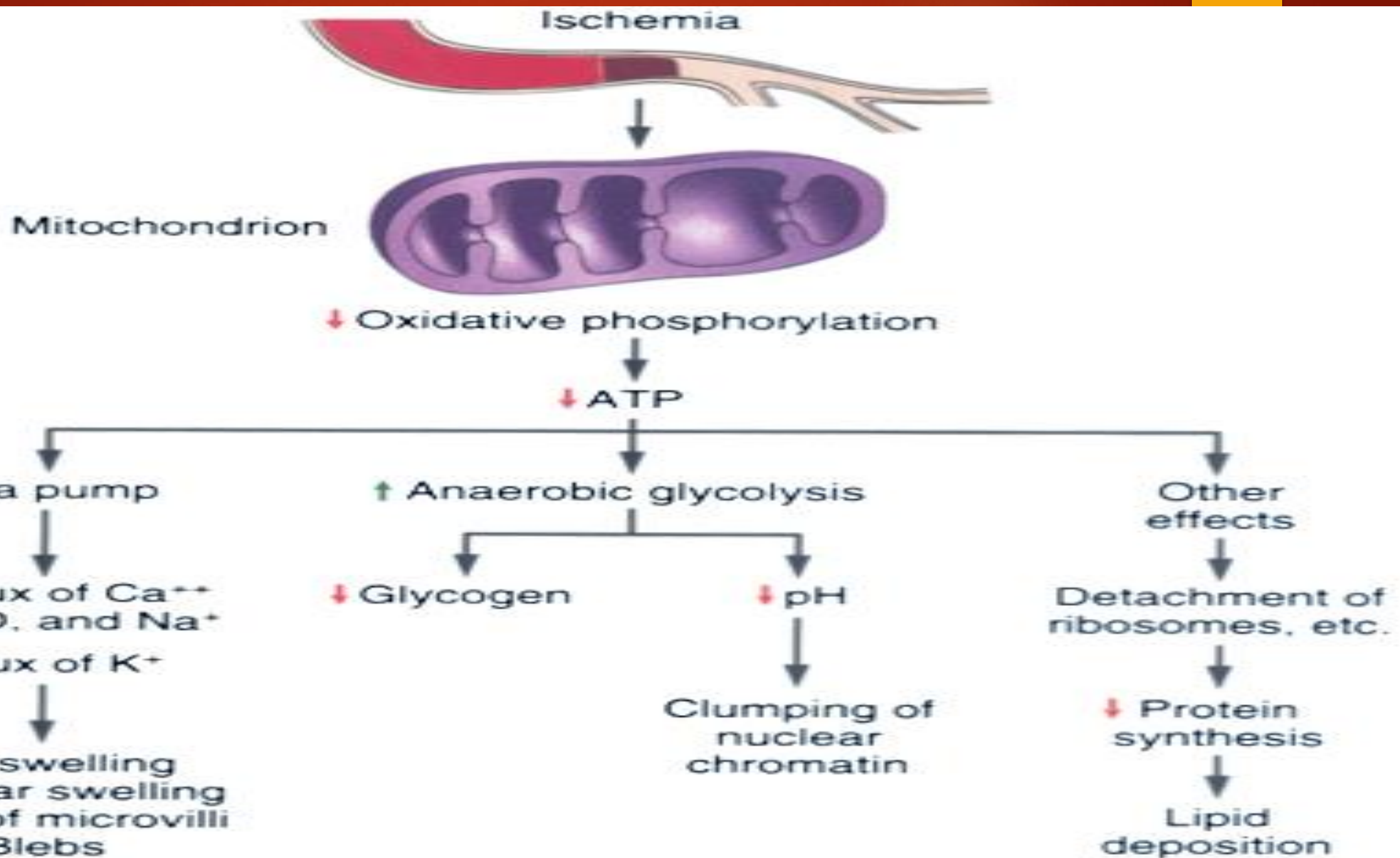
↓ Protein synthesis

**Necrosis (irreversible
damage)**

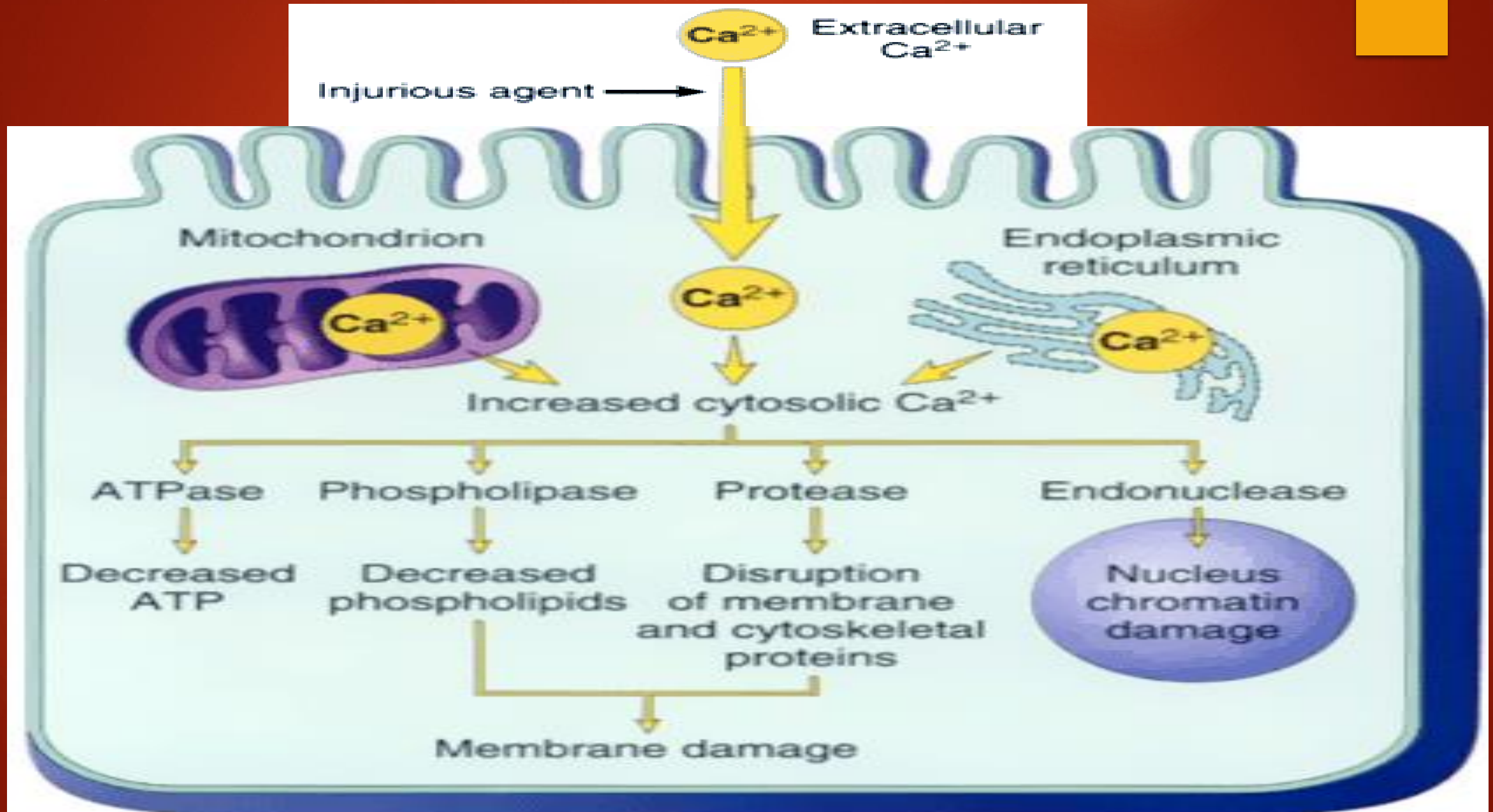
proteins misfolded

**trigger a cell
reaction-unfolded
protein response**

**Cell injury &
death**



Influx of intracellular calcium



**Hypoxia or toxins
increased cytosolic Ca²⁺
, oxidative stress, or
phospholipid breakdown**



**Formation of a high conductance
channel (mitochondrial permeability
transition pore)**



**Leaks protons and dissipates the
electromotive potential (which drives
oxidative phosphorylation)**



Damaged mitochondria



**Leak
cytochrome c**



Apoptosis

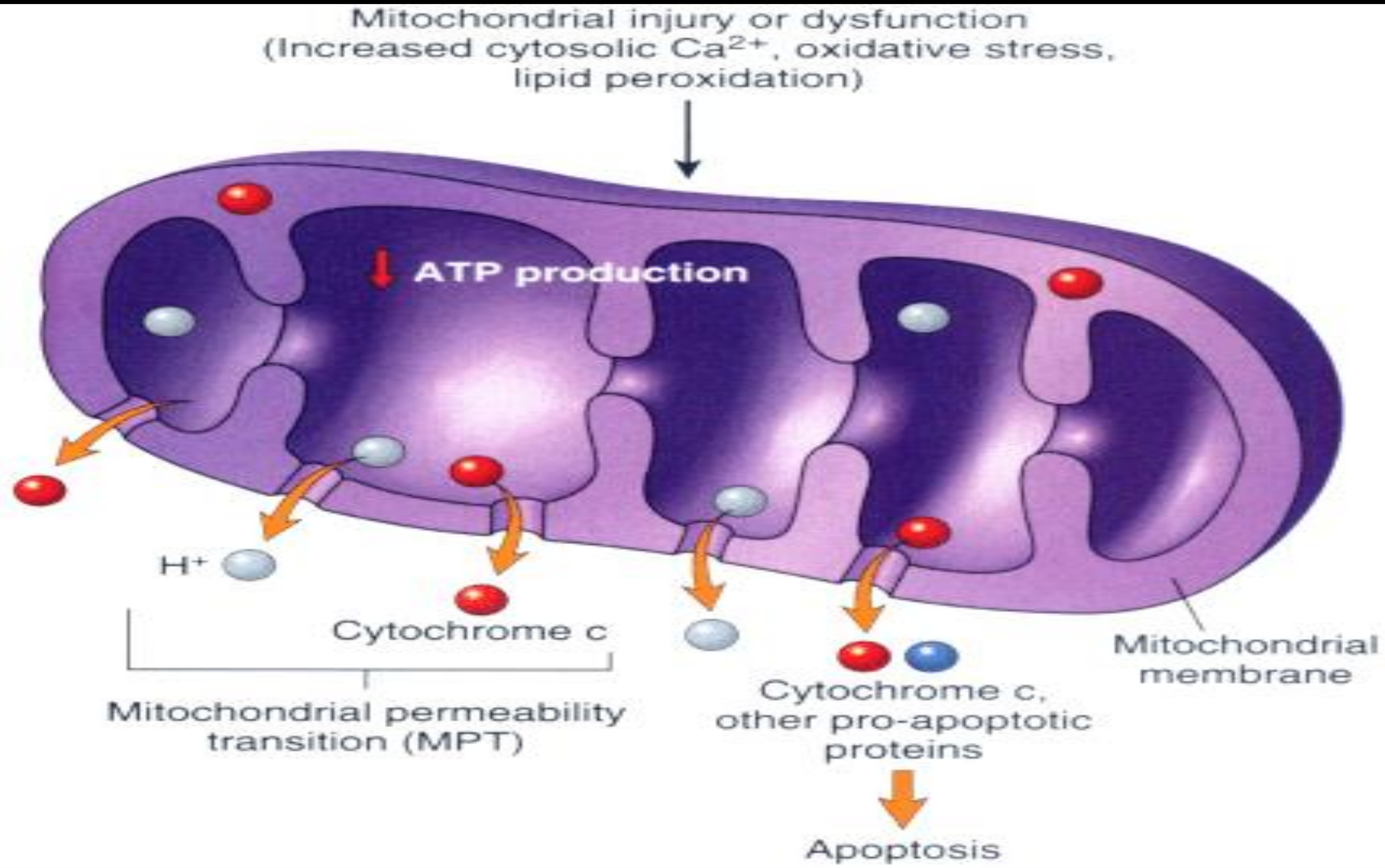
Irreversible mitochondrial transition



Death of cell

**Mitochondrial
damage**

Mitochondrial dysfunction

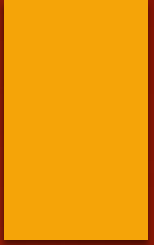


Oxidative stress

- ▶ Cell **generate energy** by reducing **molecular oxygen** to water
- ▶ During this process, small amounts of **partially reduced reactive oxygen forms** are produced as an unavoidable by product of mitochondrial respiration.

Some of these forms can damage

- ▶ Lipids
- ▶ Proteins
- ▶ Nucleic acids

- 
- ▶ Imbalance between free radical generating and radical scavenging system results in oxidative stress.
 - ▶ Free radicals initiate auto catalytic rxn's, where by molecules with which they react are themselves converted into free radicals to propagate the chain of damage.

Free radicals may be initiated within cells in several ways

- ▶ Absorption of **radiant energy**
- ▶ Enzymatic metabolism of **exogenous chemicals or drugs**
- ▶ The **reduction-oxidation reactions** that occur during normal metabolic processes
- ▶ Transition metals such as **iron and copper donate or accept free electrons** during intracellular reactions and catalyze free radical formation

Three reactions of reactive species are particularly relevant to cell injury

- ▶ Lipid peroxidation of membranes.
- ▶ Oxidative modification of proteins
- ▶ Lesions in DNA

Lipid peroxidation of membranes.

Double bonds in unsaturated fatty acids of membrane lipids are attacked by oxygen-derived free radicals



Lipid-free radical interactions yield peroxides



Peroxides
unstable and reactive

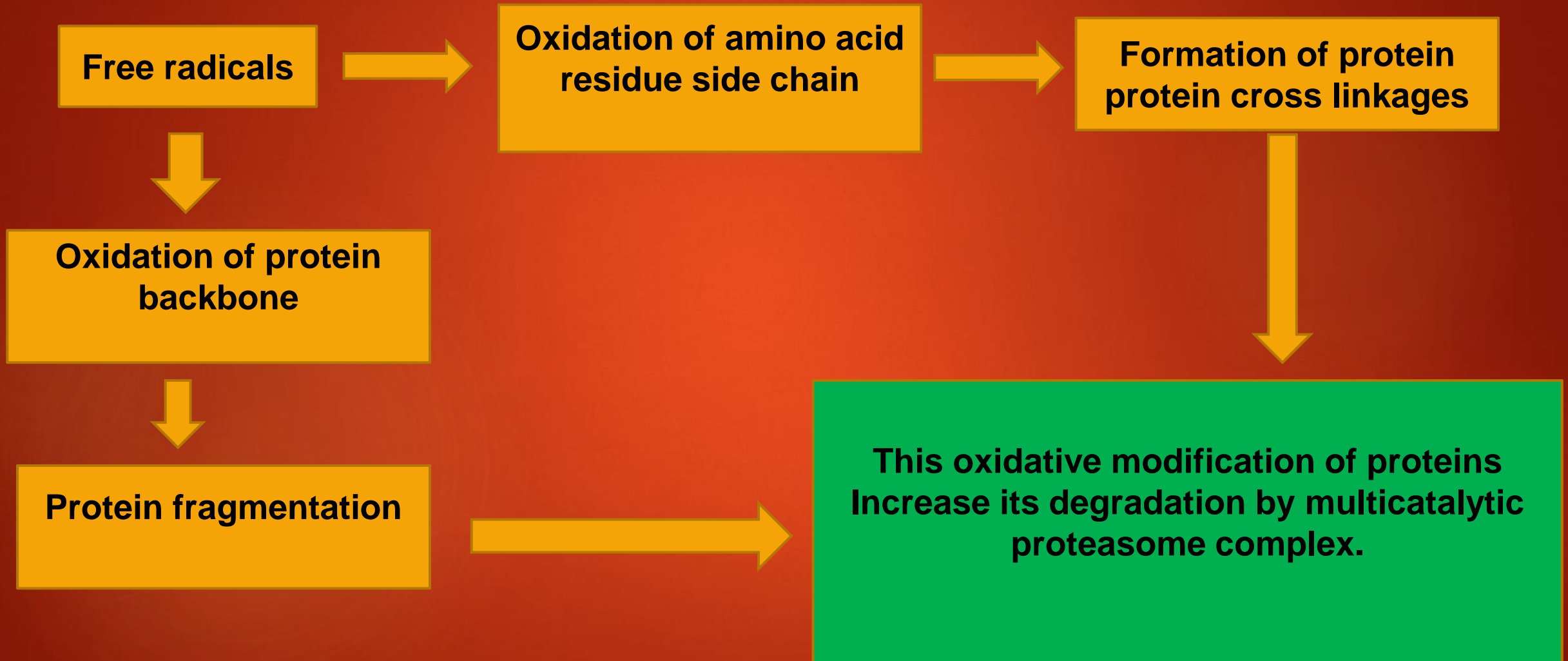


Autocatalytic chain reaction ensues (called *propagation*)



Extensive membrane, organelle, and cellular damage

Oxidative modification of proteins

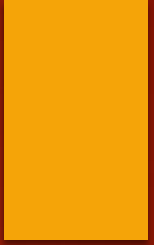


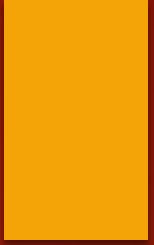
Lesions in DNA

**Free Radicals React with
thiamine in nuclear &
mitochondrial DNA**

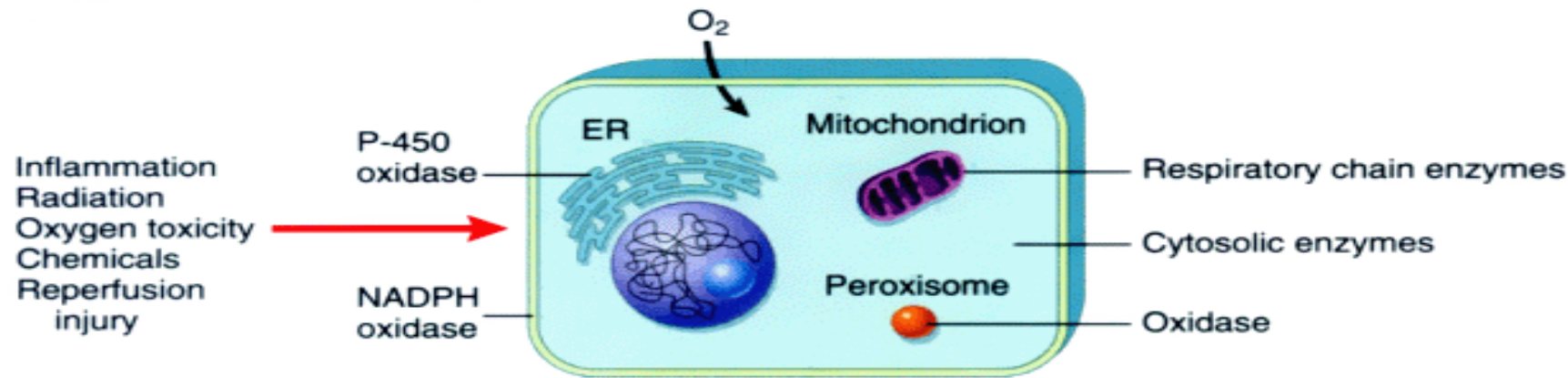


**Produce single
stranded breaks in
DNA**

- 
- ▶ Cells have developed multiple mechanisms to remove free radicals and thereby minimize injury.
 - ▶ Free radicals are inherently unstable and generally decay spontaneously
 - ▶ Several non enzymatic and enzymatic systems that contribute to inactivation of free radical reactions

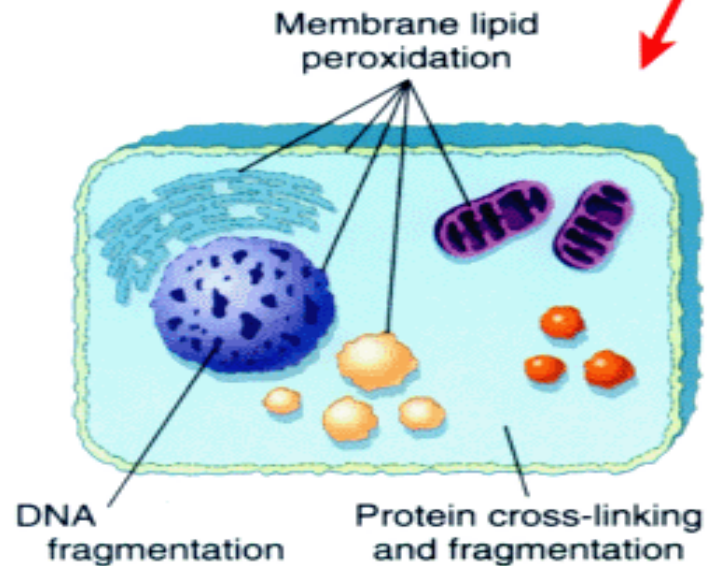
- 
- ▶ Antioxidants (non enzymatic)-Vit E & A, ascorbic acid, glutathione.
 - ▶ Enzymes-catalase, superoxide dismutase, glutathione peroxidase.
 - ▶ In many pathologic processes, the final effects induced by free radicals depend on the **net balance between free radical formation and termination**

A. FREE RADICAL GENERATION



Reactive oxygen species:
 $O_2^{\cdot-}$, H_2O_2 , OH^{\cdot}

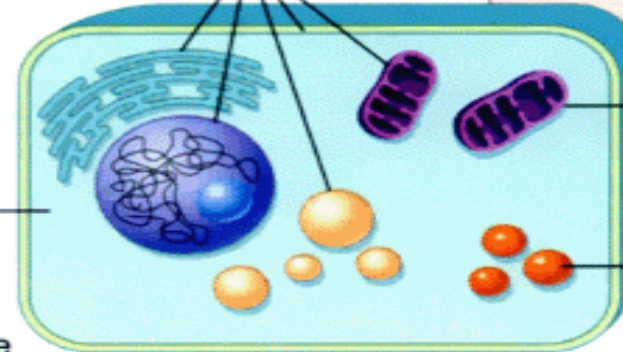
Reactive oxygen species:
 $O_2^{\cdot-}$, H_2O_2 , OH^{\cdot}



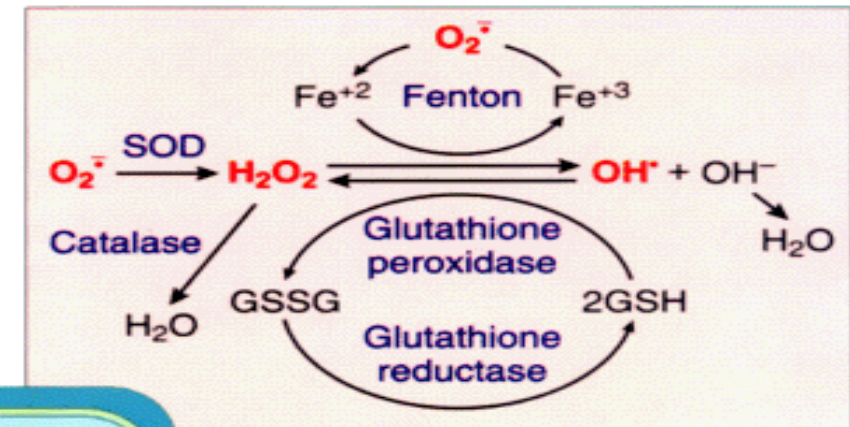
B. CELL INJURY BY FREE RADICALS

All membranes
• Vitamins E and A
• β -carotene

Cytosol
• SOD
• Vitamin C
• Glutathione peroxidase
• Ferritin
• Ceruloplasmin



C. NEUTRALIZATION OF FREE RADICALS – NO CELL INJURY



Membrane permeability defects

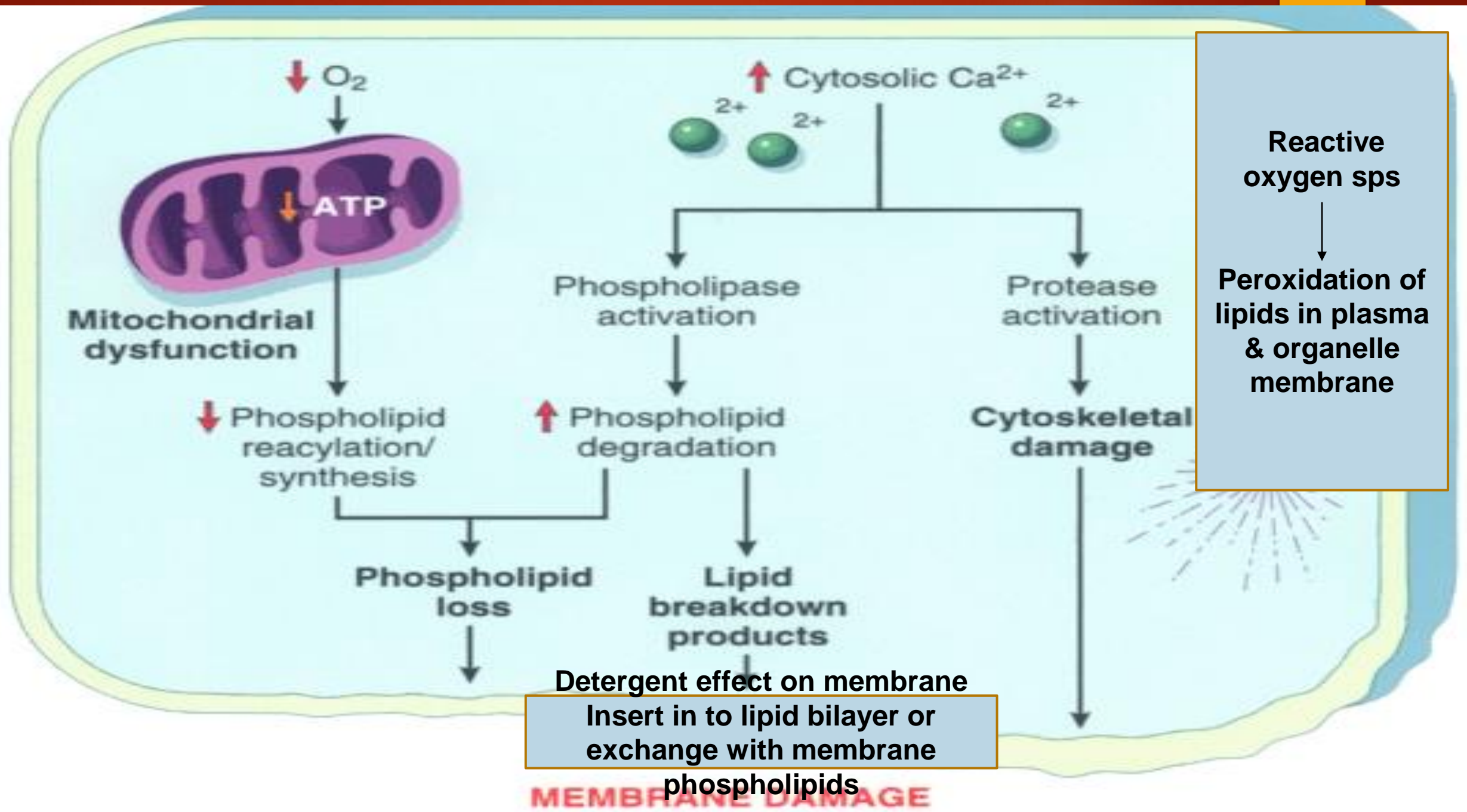
Bacterial toxins
Viral proteins
Lytic complement components
Physical & chemical agents

Directly

Damage plasma membrane

Others include

- 1. Mitochondrial dysfunction**
- 2. Loss of membrane phospholipids**
- 3. Cytoskeleton abnormalities**
- 4. Reactive O₂ species**
- 5. Lipid break down products**



Consequences of loss of membranes

- ▶ Plasma membrane damage results in **loss of osmotic balance** and **influx of fluids and ions**, as well as **loss of proteins, enzymes, coenzymes, and ribonucleic acids**.
- ▶ The cells may also **leak metabolites**, which are vital for the reconstitution of ATP, thus further **depleting net intracellular high-energy phosphates**.

Consequences of loss of membranes

- ▶ Injury to **lysosomal membranes** results in **leakage of their enzymes** into the cytoplasm and **activation** of these enzymes.
- ▶ Lysosomes contain **RNases, DNases, proteases, phosphatases, glucosidases, and cathepsins**.
- ▶ Activation of these enzymes leads to **enzymatic digestion of cell components**, resulting in loss of ribonucleoprotein, deoxyribonucleoprotein, and glycogen, and the **cells die by necrosis**.

Reversible and Irreversible Cell Injury

- ▶ **Persistent or excessive injury**, however, causes cells to pass the threshold into **irreversible injury**

Two phenomena consistently characterize irreversibility.

- ▶ Inability to reverse **mitochondrial dysfunction** (lack of oxidative phosphorylation and ATP generation)
- ▶ Profound disturbances in **membrane function**.



Whatever the mechanism(s) of membrane damage, the end result is

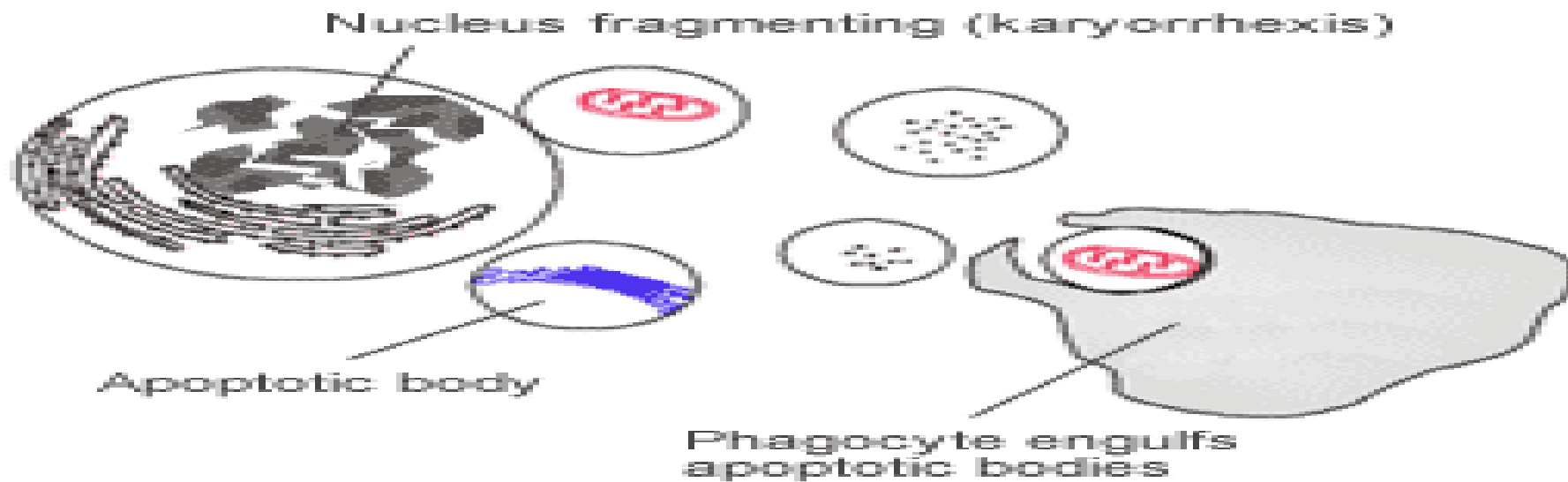
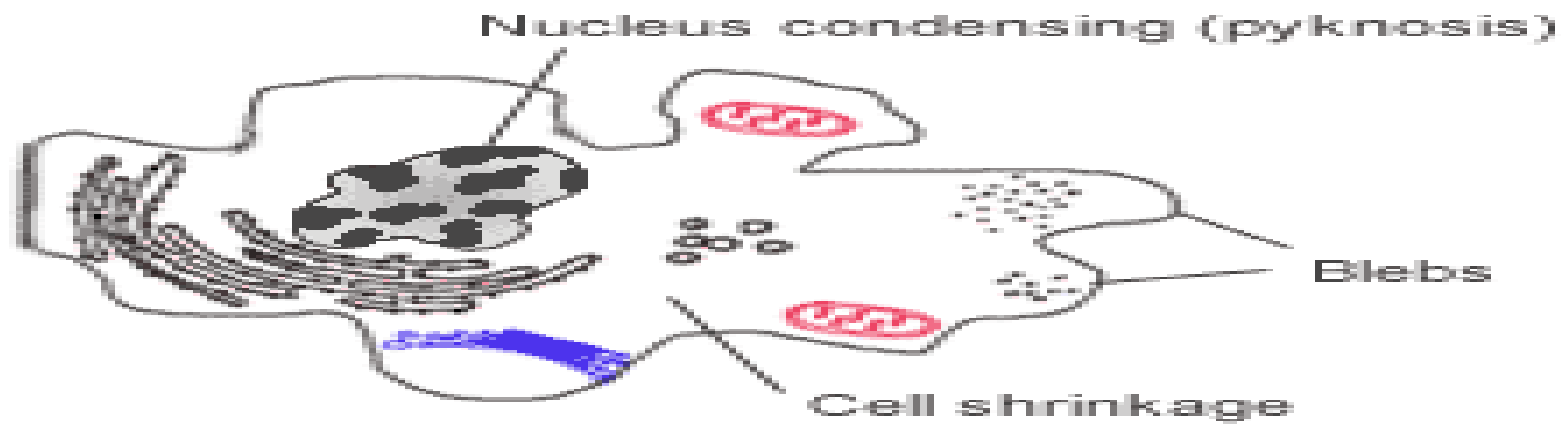
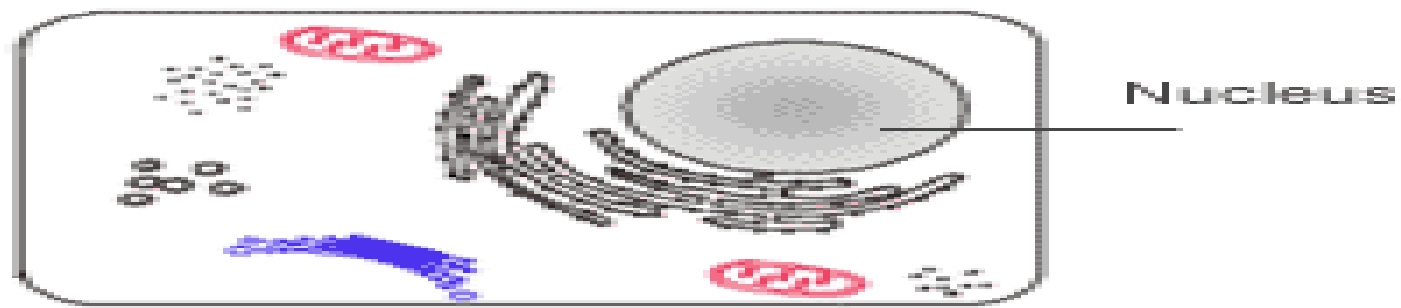
- ▶ Massive leak of intracellular materials and
- ▶ A massive influx of calcium

Morphology of Reversible injury

Two patterns of reversible cell injury (seen under the light microscope)

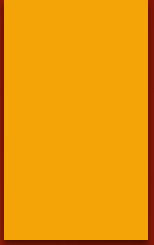
- ▶ Cellular swelling and
- ▶ Fatty change

- ▶ **Plasma membrane alterations**, such as blebbing, blunting, and distortion of microvilli; creation of myelin figures; and loosening of intercellular attachments
- ▶ **Mitochondrial changes**, including swelling, rarefaction, and the appearance of small phospholipid-rich amorphous densities
- ▶ **Dilation of the endoplasmic reticulum**, with detachment and disaggregation of polysomes
- ▶ **Nuclear alterations**, with disaggregation of granular and fibrillar elements.



Necrosis

- ▶ Necrosis refers to a spectrum of morphologic changes that follow cell death in living tissue, largely resulting from the progressive degradative action of enzymes on the lethally injured cell.
- ▶ Necrotic cells are unable to maintain membrane integrity and their contents often leak out
- ▶ The morphologic appearance of necrosis is the result of denaturation of intracellular proteins and enzymatic digestion of the cell.

- 
- ▶ Ultimately, in the living patient, most **necrotic cells and their debris disappear** by a combined process of **enzymatic digestion and fragmentation**, followed by **phagocytosis** of the particulate debris by leukocytes
 - ▶ If necrotic cells and cellular debris are **not promptly destroyed** and reabsorbed, they tend to **attract calcium salts** and other minerals and to become calcified. This phenomenon, called **dystrophic calcification**

Morphology of necrosis

Necrotic cells show

- ▶ Increased eosinophilia
- ▶ Glassy homogenous appearance (due to glycogen loss)
- ▶ Moth-eaten appearance of cytoplasm (digested cytoplasm)
- ▶ Calcification
- ▶ Replacement by phospholipid masses (myelin figures)
- ▶ Dilation of mitochondria with large amorphous densities
- ▶ Amorphous osmiophilic debris
- ▶ Discontinuities in plasma and organelle membranes
- ▶ Aggregates of fluffy material probably representing denatured protein

Nuclear change

- ▶ Karyolysis (basophilia of chromatin fade)
- ▶ Pyknosis (nuclear shrinkage & increased basophilia)
- ▶ Karyorrhexis (pyknotic nucleus undergo fragmentation)

Types of necrosis

- ▶ Coagulative necrosis
- ▶ Liquefactive necrosis
- ▶ Caseous necrosis
- ▶ Fat necrosis

Examples of cell injury and necrosis

▶ ??????????????????

Apoptosis

- ▶ Programmed cell death
- ▶ Pathway of cell death that is induced by a tightly regulated intracellular program in which cells destined to die by activate enzymes that degrade the cells' own nuclear DNA and nuclear and cytoplasmic proteins
- ▶ In apoptosis dead cells rapidly cleared before its contents leaked out
- ▶ Apoptosis occurs when cells are damaged beyond repair (esp when DNA affected)



It occurs in

1. Pathological conditions

- ▶ Cell death produced by a variety of injurious stimuli
- ▶ Cell death in tumors

2. Physiological conditions

- ▶ Cell death induced by cytotoxic T cells, a defense mechanism
- ▶ Hormone-dependent involution in the adult, such as endometrial cell breakdown during the menstrual cycle

Mechanism of apoptosis

Divided into two phases

- ▶ Initiation phase- Capases become catalytically active
- ▶ Execution phase-Capases act to cause cell death.

Initiation phase

Acts by 2 pathways

- ▶ Extrinsic or receptor mediated pathway
- ▶ Intrinsic or mitochondrial pathway

Execution phase

- ▶ Procaspases get activated (executor caspase) by **caspase 8 Or 10** in extrinsic pathway or by **caspase 9** in intrinsic pathway
- ▶ **Executor caspase** formed will further give **+ve feed back** to produce massive of active caspases
- ▶ This leads to **activation of death program** in which the activated caspases act on many cellular components (**cleave cytoskeleton ,nuclear protein, breakdown of nuclear matrix**) resulting in cell death.

Removal of dead cells

- ▶ Dying cells **secrete soluble factors** which recruit phagocytes
- ▶ **Apoptotic cells** have **marker molecules** on surface which facilitate early recognition by adjacent cells/phagocytes.
- ▶ **Macrophages secrete** substance which specifically binds to dead cells

Morphology of cell undergoing apoptosis

- ▶ Cell shrinkage
- ▶ Chromatin condensation, nucleus break down
- ▶ Cytoplasmic blebs
- ▶ Apoptotic bodies
- ▶ Phagocytosis of apoptotic cells or cell bodies by macrophages

Biochemical features

- ▶ Protein cleavage
- ▶ DNA breakdown
- ▶ Phagocytic recognition