### patho unit3 inflammation and repair Cheat Sheet by damn via cheatography.com/195477/cs/40963/

#### Acute inflamation

#### Vascular response

1. increase blood flow by vasodilation (histamine)+ vascular congestion

- -> redness, heat
- 2. increase permeability of vessel by retraction and injury
- -> edema

3. Lymph flow increase to drain extravascular fluid + secondary inflammation

#### Terminations

- 1. Mediator bursts rapidly due to short half lives e.g. neutrophil
- 2. Trigger stop signals
- proinflammatory leukotriene to anti-inflammatory lipoxins
- release anti-inflammatory cytokines

#### **Chemical mediators**

Cell-derived	Plasma-derived
1. Vasoactive amines	1.Complement system
- Histamine by mast cell	- Inflammation
- Serotonin by platelet aggreg- ation	- Opsonisation & Phagocytosis
	- Cell lysis
2. Arachidonic acid metabolites	
Both by leukocyte in lipoxy- genagse pathway	2.Clotting system
Leukotrienes	- Clotting system: induce thrombin formation

- LTC4,D4,E4: Vasoconstriction, increase vascular permeability	- Kinin system: vasoactive
- Inhibit by LT receptor antagonist	- Complement system
Lipoxins	- Fibrinolytic system
- Suppress inflammation	
Prostacyclin	3.Kinins
- PGI2, PDI2, PEI2: Vasodilation	- form bradykinin
	a. increase vascular permeability
3.Cytokines& chemokines	b. non-vascular smooth muscle contraction
Cytokines	c. pain
- Tumor necrosis factor (TNF) & interl- eukin-1 (IL-1)	4. rapidly inactivated
a. Increase endothelial cell adhesion mo	lecule expression
b. Activation and aggregation of PMN	
c. Systemic acute-phase response:Feve	r
Chemokines	
- Attract WBC	
Causes	
1. Prolonged inflammation	
2. Prolonged toxic substance exposure	
3. Autoimmune disease	

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#### Morphological change

1. Mononuclear cell infiltration (e.g. macrophage, lymphocyte, plasma cell)

- 2. Cell destruction by inflammatory cells
- 3. Repair attempts by fibrosis & angiogenesis

#### Types of inflammation

Granulomatus inflammation	Defective inflammation
- Produce granuloma containing an difficult offending agent	-delayed wound healing
- squamous>epitheloid	Excessive inflammation
- fuse> multinuclear giant cell	- Abnormal reaction of body e.g. allergy
	- Fibrosis & tissue injuries

#### Scar formation steps(Connective tissue deposition)

- 1. Angiogenesis
- 2. Granulation tissue formation
- 3. Connective tissue remodelling'

#### Cutaneous wound healing(1)

- Clean wound, only epithelial layer

#### Inflammatory phase

- 1. Formation of blood clot
- Neutrophil appears after 24hrs
- Proteolytic enzyme to clean out debris and invading bacteria

#### Proliferative phase

- 1. Formation of granulation tissue
- Induction of fibroblast and endothelial cell proliferation
- Composed of newly formed thin capillaries & loose ECM also
- Peak at day5

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#### Cutaneous wound healing(1) (cont)

- To cover the wound

#### 2. Angiogenesis

- i. VEGF >Vasodilation& I+permeability
- ii. Proteolytic degradation of parent vessel BM>> capillary sprout
- iii. Migration of endothelial cells toward angiogenetic stimulus

 iv. Proliferation of endothelial cell behind leading edge of migrating cells

v.Maturation of end. cells into capillary tubes

vi. Recruitment of periendothelial cells for mature vessel

3.Cell proliferation and collagen deposition

 Macrophage replace neutrophils after 48hrs (key cellular constituents> main resource for chemokines & GF)

- Migration and proliferation of fibroblast at injury site > secrete and deposit collagen
- Epithelial cells proliferate to centre of wound

#### Remodeling phase

- 4. Scar formation
- Granulation tissue>Scar
- Composed of inactive spindle-shaped fibroblasts, dense collagen, fragments of elastic tissue, ECM
- Pale, avascular
- 5. Connective tissue remodeling
- -Balance between ECM synthesis & degradation
- Degradation of collagen & MMPs > smaller & softer scar

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Defective	Morphologic patterns and systemic effects	
1.Defective scar formation	Morphologic	Cytokine-induced systemic reaction aka Acute
- Ulceration	patterns	phase responses
2.Excessive sf (keloid)	1.Serous	1.Fever by pyrogens
3.Contracture	2. Fibrinous	2. Leukocytosis
	- Increase cell	3.Phase proteins
Cellular response	fibrin	
important leukocyte; neutrophil and macrophages	- risk of scar	- CRP, Fibrinogen,SAA
1. Adhesion to endothelium		
a. Margination	3.Purulent	
- stasis of blood >settle out the central flow and marginate along	-Pus, leukocyte and debris	
endothelium surface	4. Ulcer	
b. Rolling	- Open lesion	
<ul> <li>complementary surface adhesion molecules sticks and release &gt; rolling along</li> </ul>	Possible outcomes	
- mediated by selectins, regulated by cytokines	1.Complete resolution	
c. Adhesion	2. Fibrosis/scarring	
- mediated by intergrins	3.Chronic inflammation	
	Colls and modiat	
2.Migration thru endo		
-secrete collagenase thru basement membrane	Macrophage	
- migrate toward chemotactic gradient	- dominant, from monocyte	
	Activated by:	
3.Chemotaxi	1. Classical path	way (microbicidal action)
-neutrophil>monocyte>macrophage	2. Alternative pathway	
	Functions	
4 Phagoestosis	1. Phagocytosis	and destruction
a Recognition by receptors to sol signals	2. Initiate tissue r	epair & scar formation and fibrosis involvement
	3. Secrete inflam	mation mediators (e.g. cytokines, clotting factors)
b. Activation by cytosolic Ca2+ and enzymes	4. Processing an	d presentation of Ag to immune system
d anti-inflammatory effects and wound repair		

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Tissue repair Regeneration (Cell proliferation)	Secondary intention (cont)
Depend on:	- Excessive collagen deposit
1. Cell types (Ability to repair)	- Pathologic process by persistent stimuli
2. Degree of injury	- Associated with loss of tissue
Proliferative potential	- Long-lasting
1. Labile (continuo)	
- e.g. epithelial cell, xxx tract	Healing Factors
2. Stable	Systemic
- e.g. salivary gland	1. Overall nutrition e.g. VitC
3. Permanent	2. Metabolic status > Vascular supply
- e.g. neuron, myocardium	3. Circulatory status
	4. Hormones > Cortico :(
Regulation mechanism	5. Age
1. Growth factors (+population,size,mitosis,survival)	Local
-VEGF	1. Infection
2. ECM	2. Movement
Cell-matrix interactions	3. Type, size, location
總之講緊growth同 differentiation要用at least 2 types of signal 一個就 用soluble(growth factor) 另一個就用insoluble(ECM)	4. Foreign bodies
Secondary intention	
- Cell loss more extensive	
Features	
1. More intense inflammatory tissue	
2. Abundant granulation tissue	
3. ECM accumulation	
4. Formation of large scar	
- Destroyed appendage are permanently lost	
5. Wound contraction	
- Reduce gap between dermal edge and wound area to close wound	

- Myofibroblast for mediator
- 6. Fibrosis



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