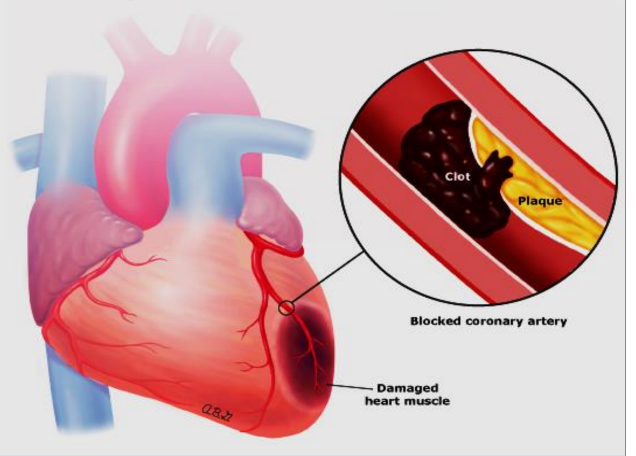


ISCHEMIC HEART DISEASE-2 Acute Myocardial Infarction

Dr. Nisreen Abu Shahin Associate Professor of Pathology Pathology Department University of Jordan

Acute Myocardial Infarction (MI)



- MI = *heart attack*.
- Necrosis of heart muscle due to ischemia.
- A significant cause of death worldwide.

Clinical Features of acute MI Severe, crushing substernal chest pain that radiates to neck, jaw, epigastrium, or left arm dyspnea (if pulmonary Dizziness; sweating congestion and edema) rapid and weak pulse cardiogenic shock (in massive MIs >40% of left nausea (in posterior MI) ventricle)

Sometimes: No typical symptoms (silent infarcts)

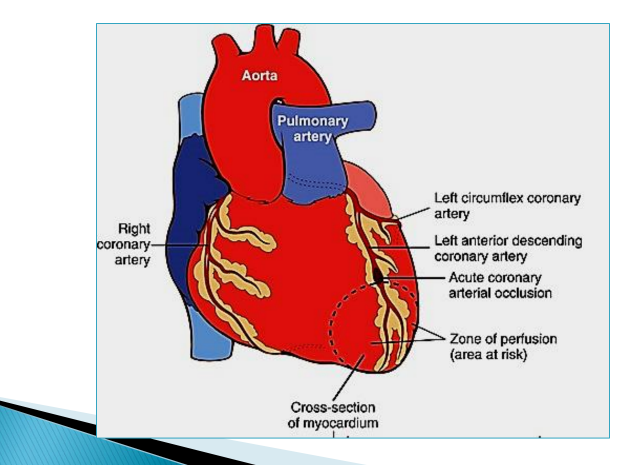
Silent infarcts:

- A variable percentage of MIs are <u>asymptomatic</u>
- Confirmed only on ECG and lab workup.
- particularly in:
 1- DM (peripheral neuropathies)
 2- the elderly

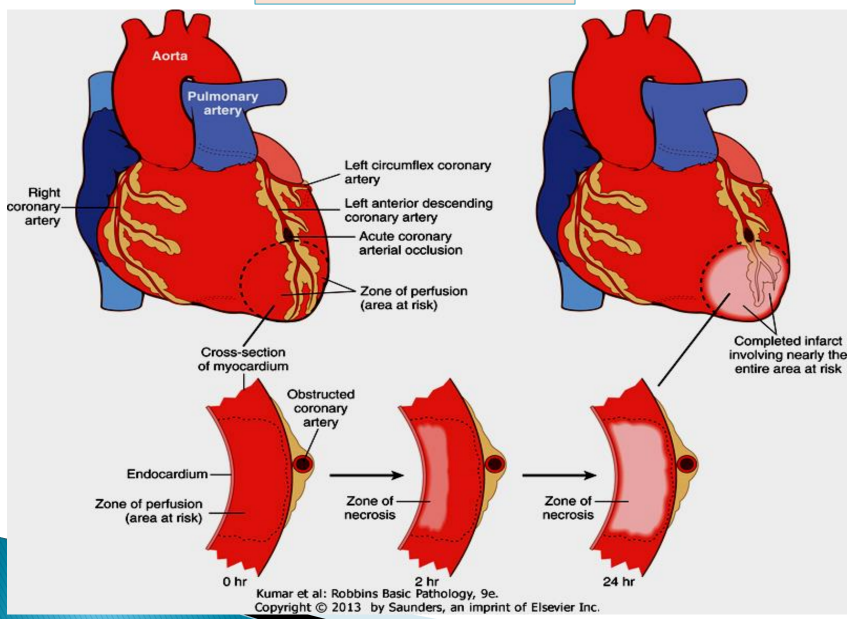


MI- Causes:

 Acute occlusion of the proximal left anterior descending (LAD) artery is the cause of <u>40% to 50% of all MI cases</u>



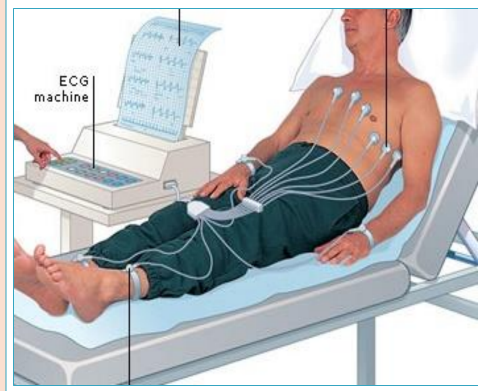
MI-Evolution



Evaluation of MI

- Clinical signs and symptoms
- Electrocardiographic (ECG) abnormalities

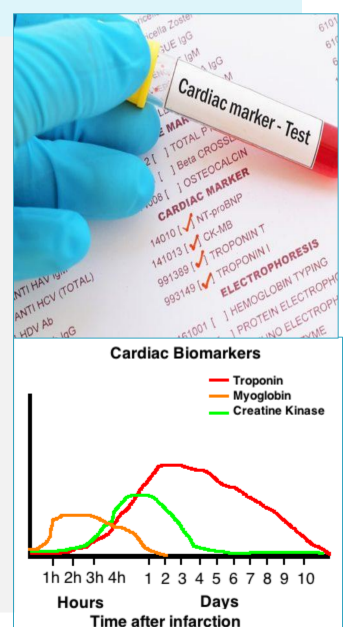
Laboratory evaluation: blood levels of intracellular macromolecules that leak out of injured myocardial cells through damaged cell membranes.



Cardiac enzymes in MI

 1-Myoglobin
 2-Cardiac Troponins T and I (TnT, TnI)
 3-Creatine kinase (CK); specifically the myocardial-specific isoform (CK-MB)
 4- Lactate dehydrogenase

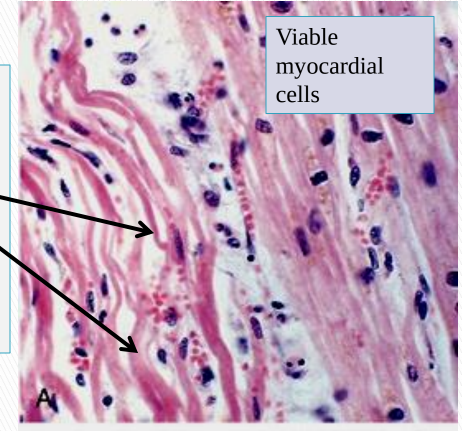
- Cardiac troponins T and I (TnT, TnI), are the best markers for acute MI.
- Creatine kinase CK-MB is the second best marker after the cardiac-specific troponins.



LearnTheHeart.com

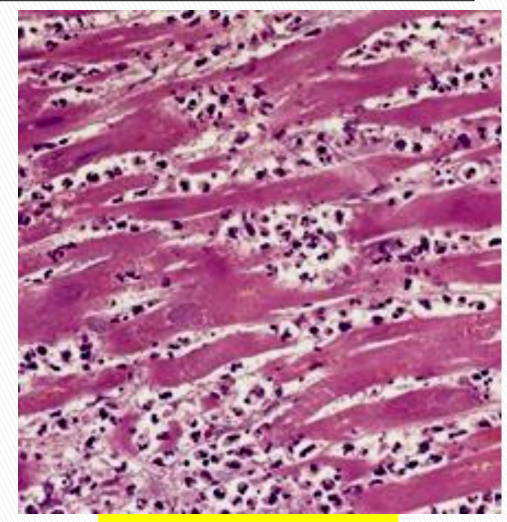
<24 hr:

coagulative **necrosis** and **wavy fibers** Necrotic cells are separated by edema fluid



Stain: Hematoxylin & Eosin (H&E)

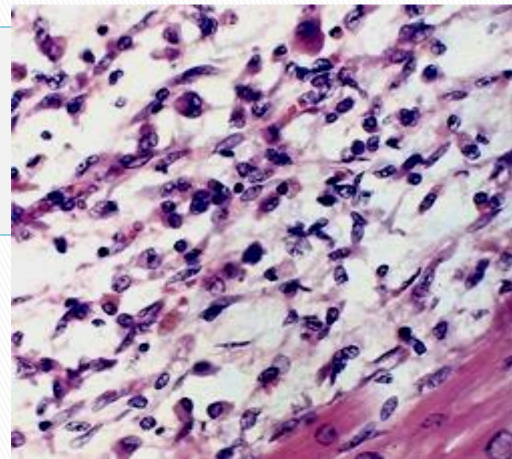
2 - 3 days:Dense neutrophilinfiltrate



Stain: Hematoxylin & Eosin (H&E)

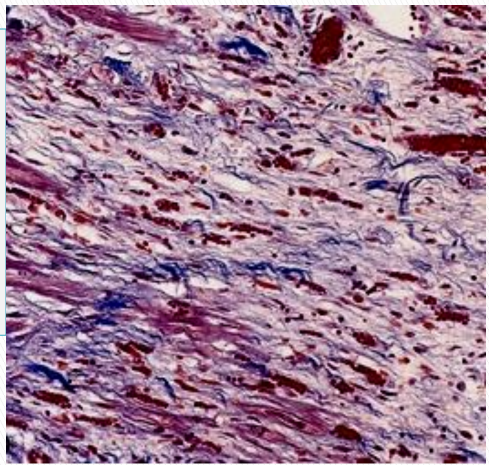
7 to 10 days:

complete removal of necrotic myocytes by **macrophages**



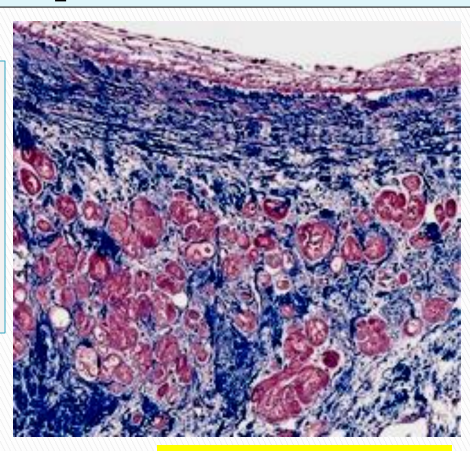
Stain: Hematoxylin & Eosin (H&E)

up to 14 days: Granulation tissue [loose connective tissue (blue) and abundant capillaries (red)]



Stain: Masson Trichrome (MT)

several weeks: Healed infarct consisting of a dense collagenous scar (blue)

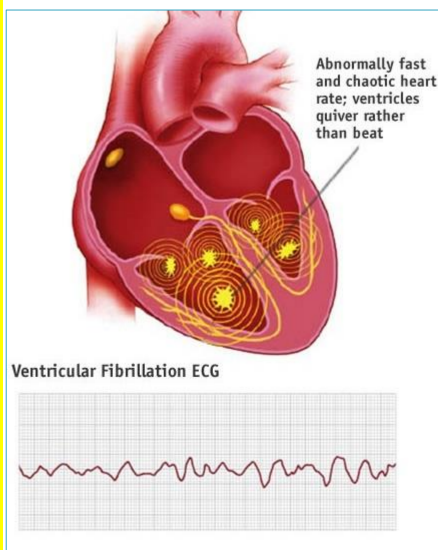


Stain: Masson Trichrome (MT)

Consequences & Complications of MI

1- <u>Death</u>:

- <u>50% occur before reaching</u> <u>hospital (within 1 hour of symptom</u> onset-usually as a result of lethal arrhythmias (<u>Sudden Cardiac</u> <u>Death</u>)
- Arrhythmias are caused by electrical abnormalities of the ischemic myocardium and conduction system
- With current medical care, patient outcome is better (*in-hospital death rate* has declined).



Consequences & Complications of MI

• <u>2- Cardiogenic shock.</u>

- 15% In large infarcts (>40% of Left ventricle).
- 70% mortality rate important cause of in-hospital deaths.
- <u>3-Myocardial rupture</u>
- <u>4-Pericarditis</u>
- 5-Infarct expansion
- <u>6- Mural thrombus</u>
- <u>7- Ventricular aneurysm</u>

8-Progressive late heart failure

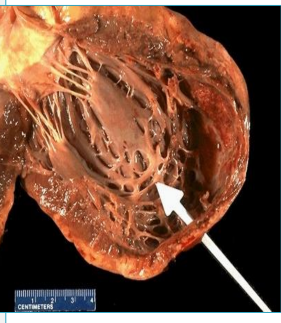
Complications of Myocardial Rupture Include:

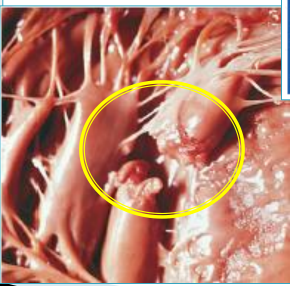
(1) rupture of the
ventricular free wall:
hemopericardium
and cardiac
tamponade (usually
fatal)

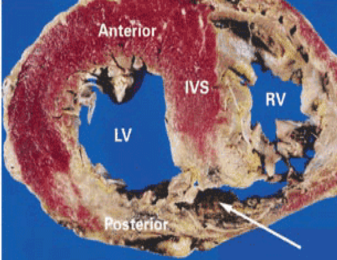
(2) rupture of the ventricular septum:VSD and left-to-right shunt

(3) papillary muscle rupture:

severe mitral or tricuspid regurgitation







4-Pericarditis.

- 2 to 3 days post a transmural MI
- spontaneously resolves (immunologic mechanism)

5-Infarct expansion.

disproportionate stretching, thinning, and dilation of the infarct region (especially with anteroseptal infarcts)

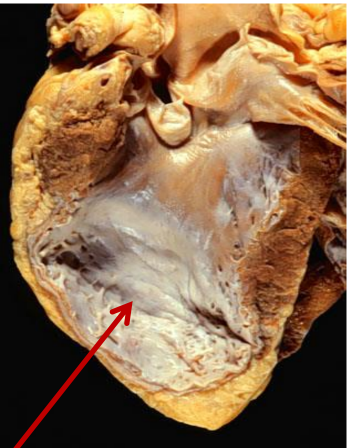
6-Mural thrombus.

loss of contractility (causing stasis) + endocardial damage \rightarrow *thromboembolism*

7-Ventricular aneurysm.

- A late complication
- most commonly result from a large transmural anteroseptal infarct that heals with the formation of thin scar tissue

Complications of ventricular aneurysms include: 1-mural thrombus 2-arrhythmias 3-heart failure

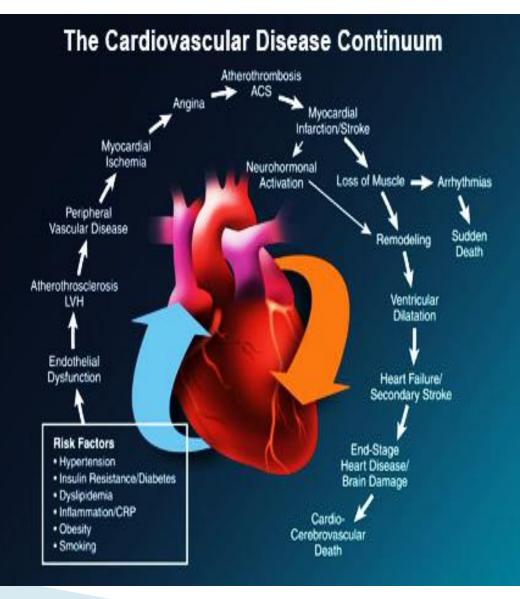


Long-term prognosis after MI

- depends on many factors: e.g. left
 ventricular function; severity of
 atherosclerosis in viable myocardium; etc...
- 1st year mortality $\approx 30\%$.
- Thereafter, the annual mortality rate $\approx 3\%$

Chronic Ischemic Heart Disease

- results from postinfarction cardiac decompensation that follows exhaustion of hypertrophic viable myocardium.
- progressive heart failure
- sometimes punctuated by episodes of angina or MI
- Arrhythmias are common



Sudden Cardiac Death (SCD)

Unexpected death from cardiac causes either without symptoms or < 24 hours of symptom onset</p>

CAD (atherosclerosis) is the most common underlying <u>cause</u>

Lethal arrythmias (v. fibrillation) is the most common <u>direct mechanism</u> of death

• With younger victims, other <u>non-atherosclerotic</u> causes are more common:

Non-atherosclerotic causes of SCD

- Congenital coronary arterial abnormalities
- Aortic valve stenosis
- Mitral valve prolapse
- Myocarditis
- Dilated/ hypertrophic cardiomyopathy
- Pulmonary hypertension
- Hereditary/ acquired abnormalities of cardiac conduction system

