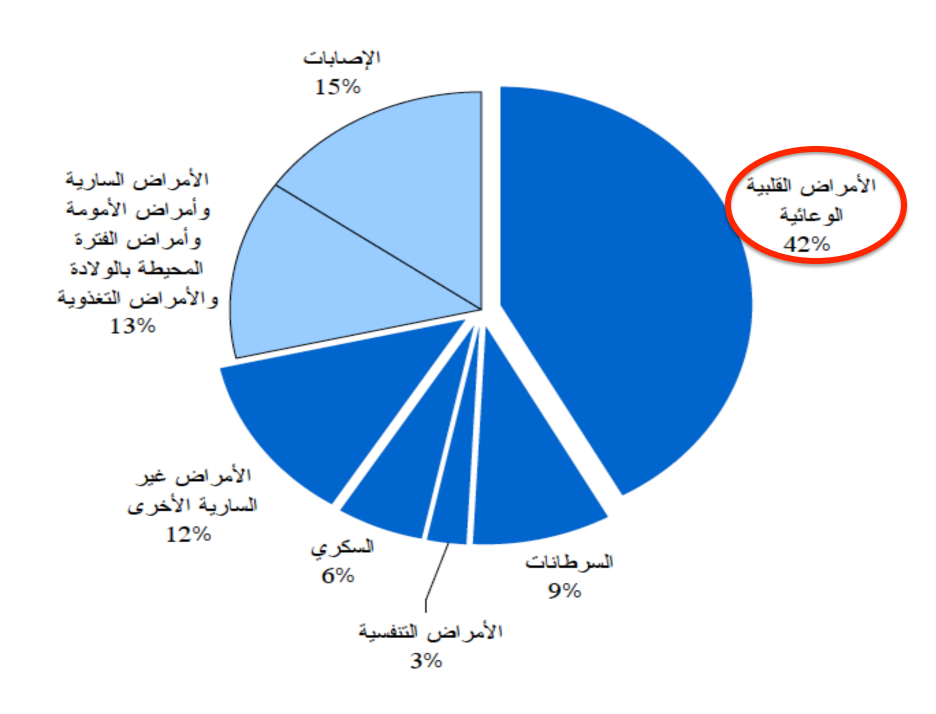
Acute Coronary Syndrome

Prof. Hussam F. Al-Faleh Cardiac Sciences Department

Why is Acute coronary syndrome important?

#1 Killer



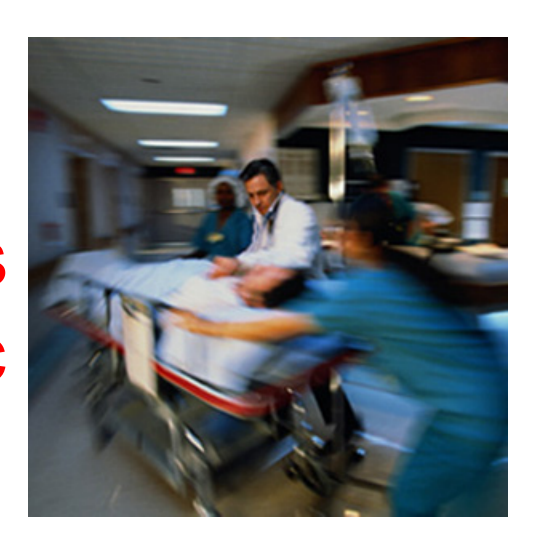
Morbidity and mortality rates of CAD in selected regions for 1990, estimated morbidity and mortality rates of CAD in selected regions for 2020, and projected increase in mortality from CAD from 1990 to 2020, for men and women

| | Men | | | Women | | |
|------------------------------|--------------------|--------------------|---|--------------------|--------------------|---|
| Region | 1990 (millions) | 2020 (millions) | Projected increase in mortality (%) | 1990 (millions) | 2020 (millions) | Projected increase in mortality (%) |
| Established market economies | 390 | 434 | 46 | 40.7 | 45.5 | 32 |
| India | 439 | 608 | 127 | 41.0 | 58.9 | 114 |
| China | 585 | 727 | 108 | 54.8 | 72.1 | 79 |
| Sub-Saharan Africa | 252 | 555 | 144 | 25.8 | 56.5 | 116 |
| Latin America | 222 | 331 | 144 | 22.3 | 33.6 | 141 |
| Middle East | 256 | 496 | 171 | 24.7 | 48.7 | 148 |





30% of ER admissions are cardiac



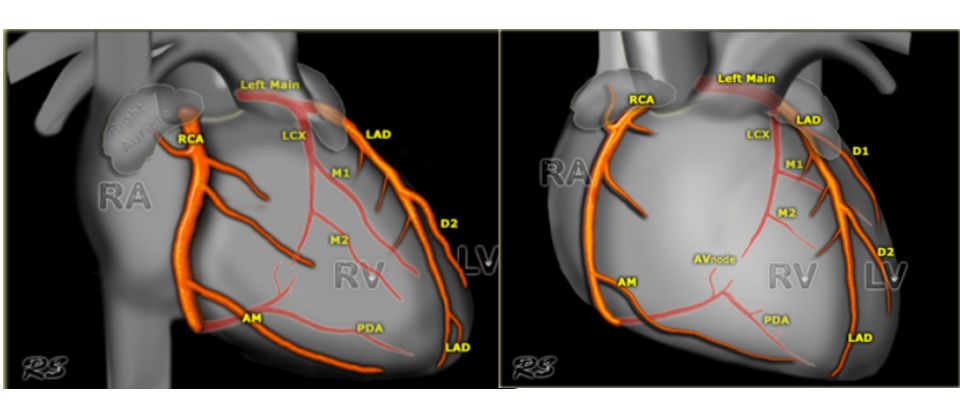
Objectives

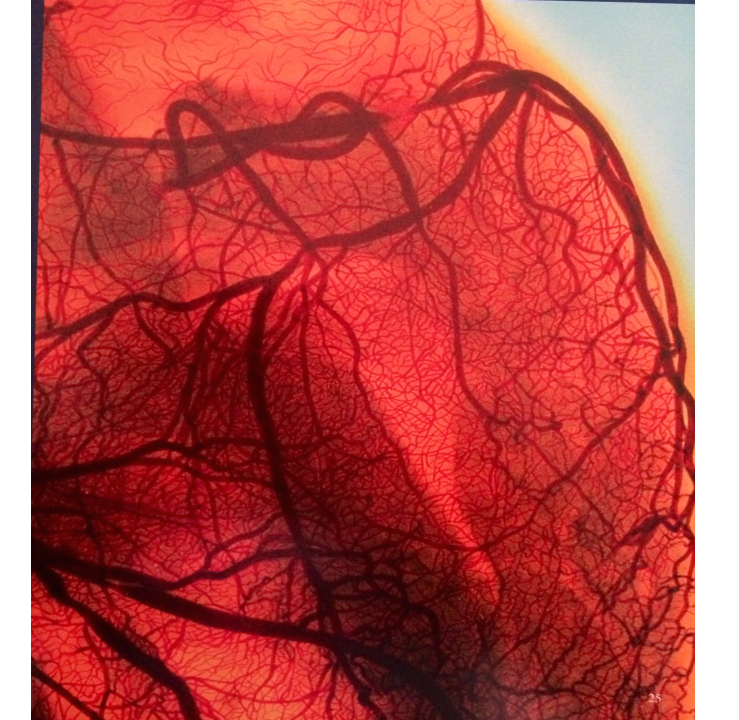
- Pathophysiology of ACS- How?
- Classification of ACS- How to label?
- Diagnostic workup- Recognize it?
- Initial management- Save a life?
- Common complications of ACS- The aftermath?

Resources

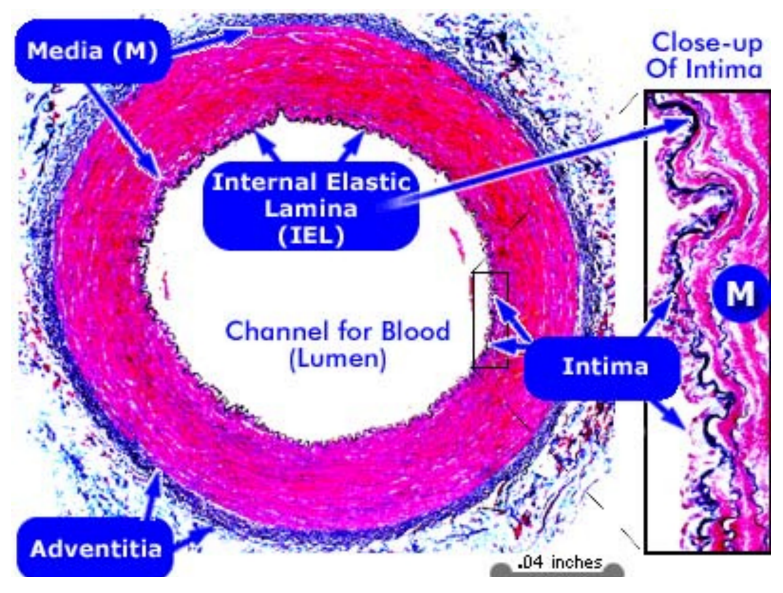
- Davidson or Kumar
- Lecture
- Optional reading (4 articles)

What are coronary arteries ??





Artery histology



What are the risk factors for CAD?

Diabetes Mellitus

One of every 4 Saudis has DM

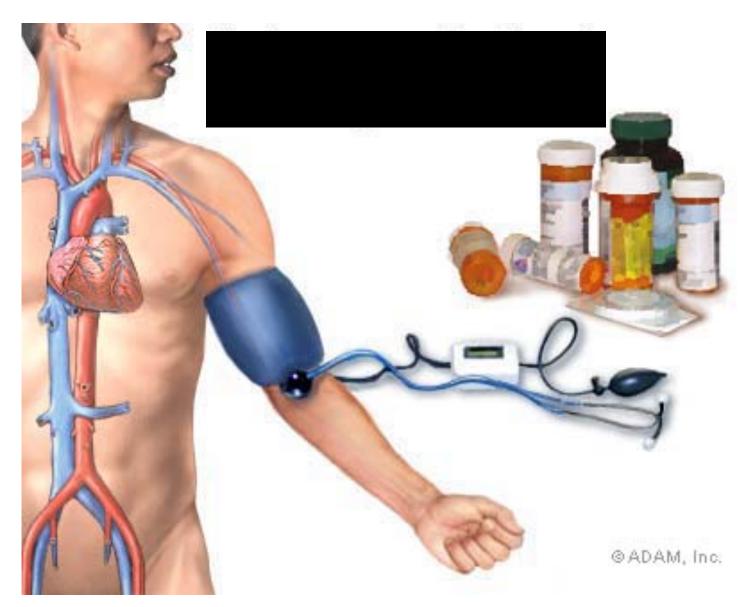


Smoking





Hypertension



Hyperlipidemia





Obesity





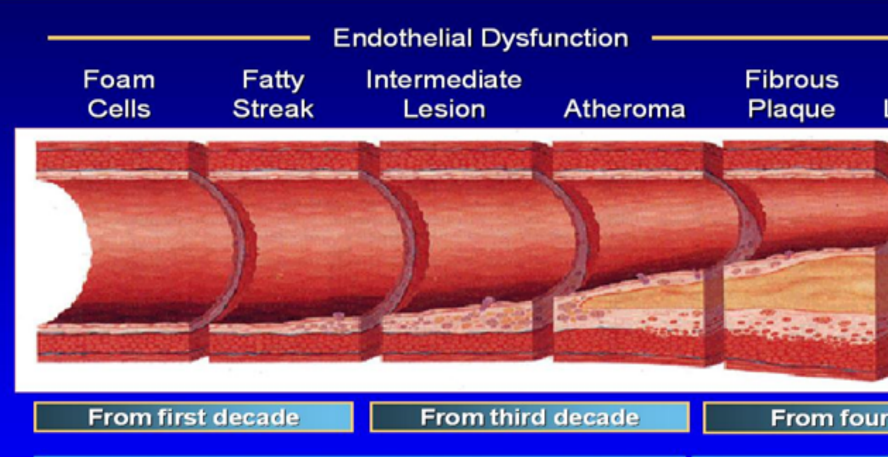
Other Risk factors

- Age : males ≥45, females ≥55
- Gender (Male gender)
- Family history of Premature CAD: males ≤55 females ≤65
- Stress ???

Role of genetics??

Known modifiable risk factors explain >90% of the occurrence of MI in populations around the world

Atherosclerosis Timeline



Growth mainly by lipid accumulation

Smooth muscle and collagen

Stary et al. Circulation. 1995;92:1355-1374.



Angina Pectoris

The remarkable facts, that the paroxysm, or indeed the disease itself, is excited more especially upon walking up hill, and after a meal; that thus excited, it is accompanied with a sensation, which threatens instant death if the motion is persisted in; and, that on stopping, the distress immediately abates, or altogether subsides; have . . . formed a constituent part of the character of Angina **Pectoris**

John Warren, 1812

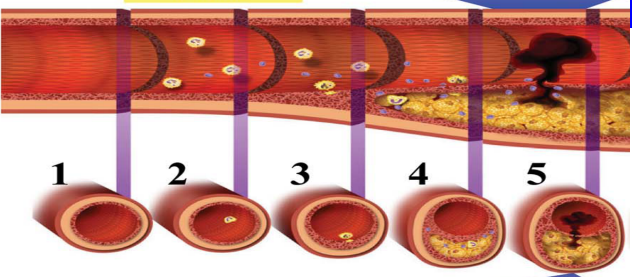
Atherosclerosis Timeline

Endothelial Dysfunction Intermediate **Fibrous** Complicated Foam Fatty Cells Streak Lesion Atheroma Plaque Lesion/Rupture From first decade From third decade From fourth decade Smooth muscle Thrombosis, Growth mainly by lipid accumulation

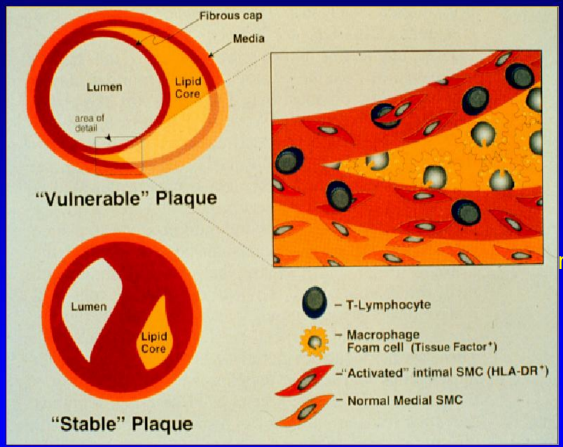
and collagen

hematoma

Stary et al. Circulation. 1995;92:1355-1374.



"Vulnerable" Plaque and "Stable" Plaque

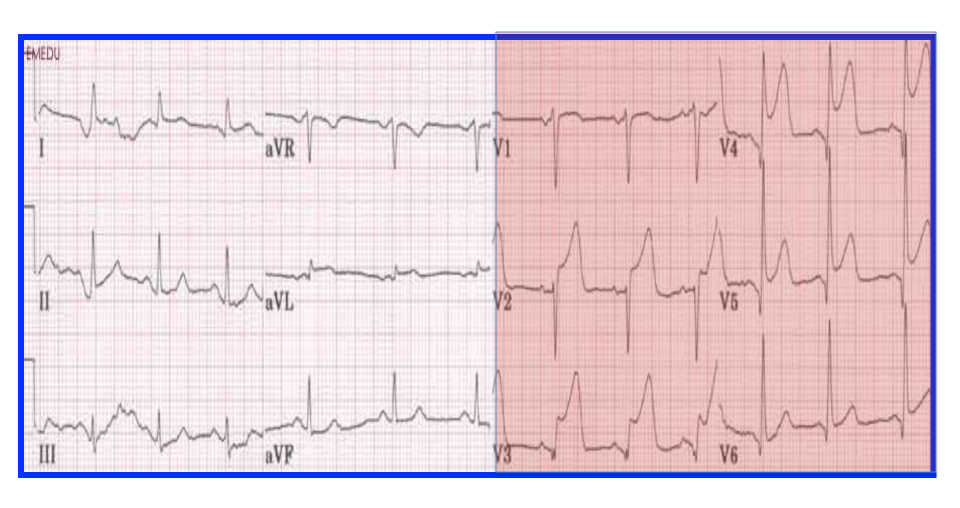


Secretion of Matrex metalloprotenases

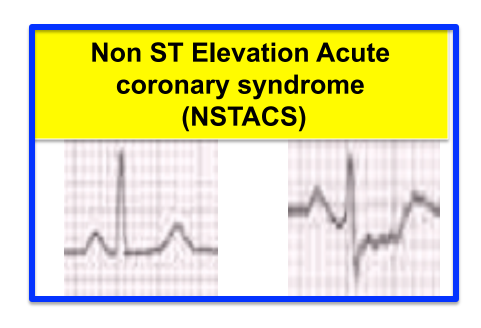
Libby. Circulation. 1995;91:2844-2850.

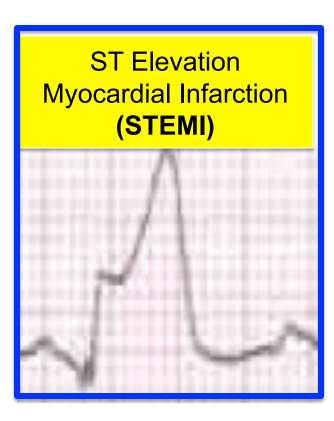
INVESTIGATIONS IN THE EMERGENCY ROOM

ST- Elevation Myocardial Infarction

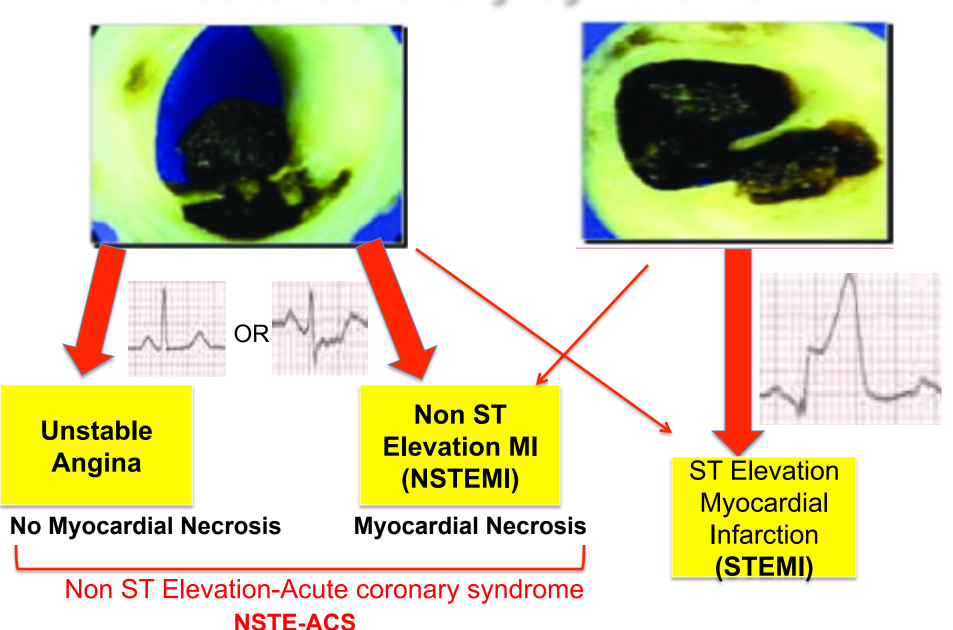


Acute Coronary Syndrome





Acute Coronary Syndrome

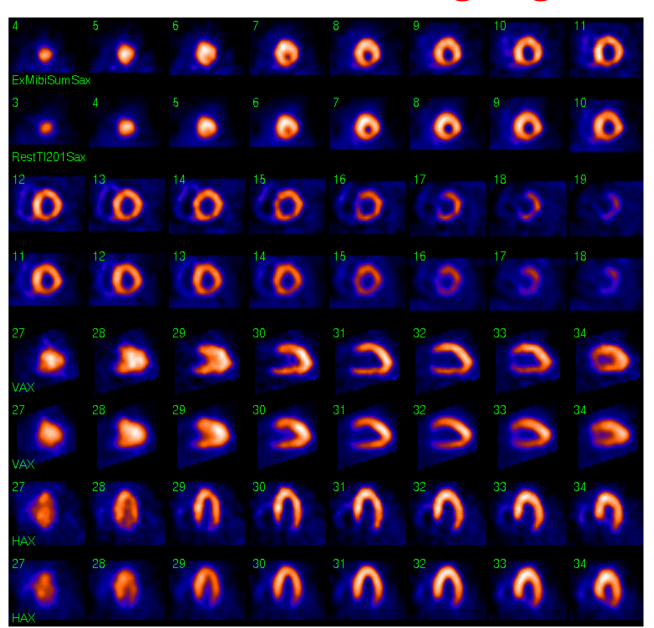


What is Myocardial Infarction?

Third Universal Definition of MI

- Typical rise in cardiac troponin T or I, CK-MB with at least one of the following:
 - 1. Ischemic symptoms
 - 2. Pathological Q wave on ECG
 - 3. Ischemic ECG changes (e.g. ST elevation or depression, new LBBB)
 - 4. Imaging evidence of new loss of viable myocardium or a new WMA
 - 5. Identification of an intracoronary thrombus by angiography or autopsy.

Nuclear imaging



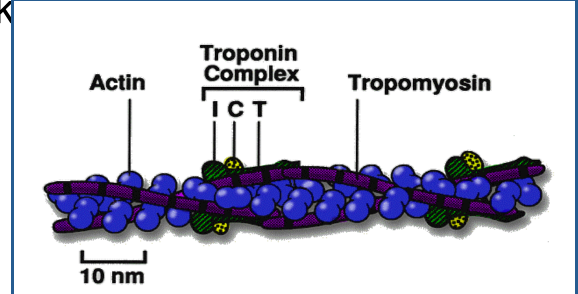
MARKERS FOR MYOCARDIAL NECROSIS

Creatin Kinase (CK)

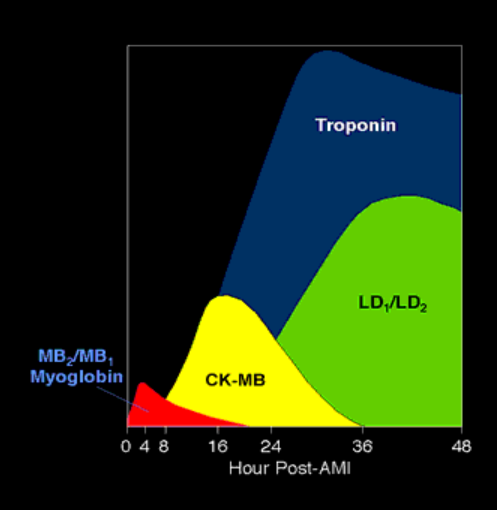
- MI causes release of certain enzymes and proteins into the blood stream.
- Creatin Kinase (CK) is released from multiple organs such as the myocardium, skeletal muscles, and the brain.
- The Iso-form CK-MB, is cardio-specific
- Starts to rise 4-6 hrs after onset of ischemia, then falls within 48-72hrs.

Troponin

- Cardiospecific proteins Troponin I, and T are the most sensitive & specific markers for myonecrosis.
- Released with 4-6hrs, but can last upto 2 week



Relationship between onset of MI and release of markers



BACK TO RASHED IN ER

Management

Aims of therapy

- Open Artery and Improve oxygen supply
 - 1. Supplemental O2
 - 2. Coronary vasodilators (Nitroglycerine)
 - 3. Antiplatelet agents
 - 4. Reperfusion therapy
 - a. Fibrinolytic therapy
 - b. Primary Percutanous coronary intervention (PCI)
 - 5. Antithrombotic agents

Aims of therapy

- Reduce O2 demand
 - 1. Beta blockers (Propranolol, Metoprolol)
 - 2. Analgesics (Morphine)
- Other medications
 - ACE inhibitors (Enalapril, Lisinopril)
 - Statin therapy

Reperfusion therapy

Fibrinolytics

- ONLY USED FOR STEMI (NOT NSTEMI)
- Reduces short and long term mortality
- Should be given during a 12hr window, and given ASAP.
- 2 types of fibrinolytics:
 - 1. Non Fibrin specific
 - (Streptokinase)
 - 2. Fibrin specific

Fibrin specific agents

| Characteristic | Alteplase (t-PA) | Reteplase (rPA) | Tenecteplase (TNK) | Lanoteplase (nPA) |
|-----------------------------------|--|---|---|--|
| Immunogenicity | No | No | No | ? |
| Plasminogen activation | Direct | Direct | Direct | Direct |
| Fibrin specificity | ++ | + | +++ | + |
| Plasma half-life | 4–6 min | 18 min | 20 min | 37 min |
| Dose | 15-mg bolus plus 90- min infusion up to 85 mg | 10+10-MU double bolus 30 min apart | ±0.5 mg/kg single bolus | 120 KU/kg single bolus |
| PAI-1 resistance | No | ? | Yes | ? |
| Genetic alteration to native t-PA | No | Yes | Yes | Yes |
| | Recombinant version | Finger, EGF, and kringle-1 regions deleted | 2 single amino acid substitutions in kringle-1 and substitution of 4 amino acids in catalytic domain | Finger, EGF regions deleted and glycosylation sites in kringle-1 domain modified |

Absolute contraindications

Any prior intracranial hemorrhage

Known structural cerebral vascular lesion

Known intracranial neoplasm

Ischemic stroke within the past 3 months (except for acute stroke within 3 hours)

Suspected aortic dissection

Active bleeding or bleeding diathesis (excluding menses)

Significant closed-head or facial trauma within 3 months

Relative contraindications

History of chronic, sever, poorly controlled hypertension

Systolic pressure >180 mm Hg or diastolic >110 mm Hg

History of prior ischemic stroke >3 months previously, dementia, or known intracranial pathology not covered in absolute contraindications

Recent (within 2–4 weeks) internal bleeding

Noncompressible vascular punctures

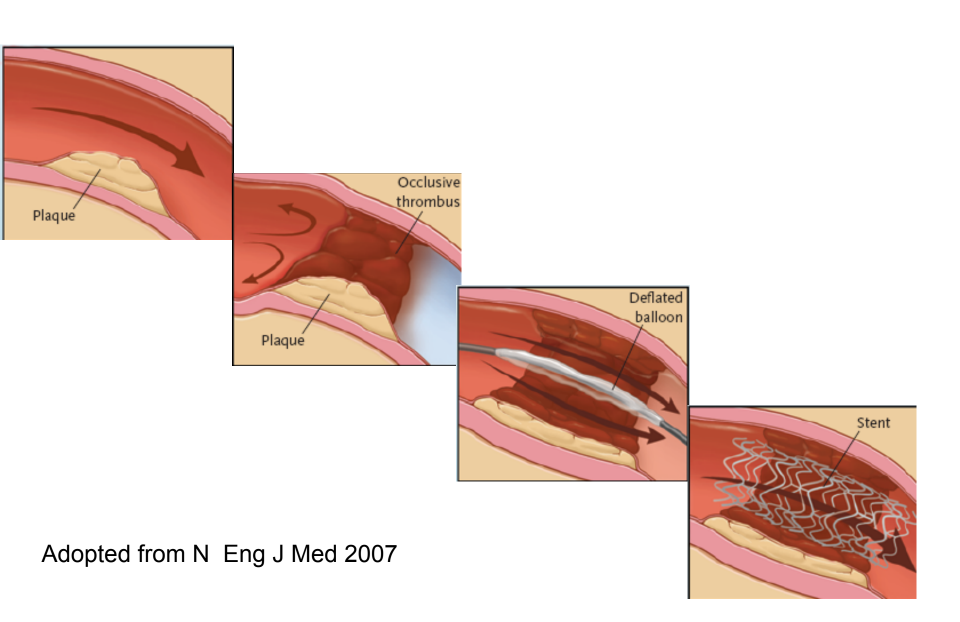
Pregnancy

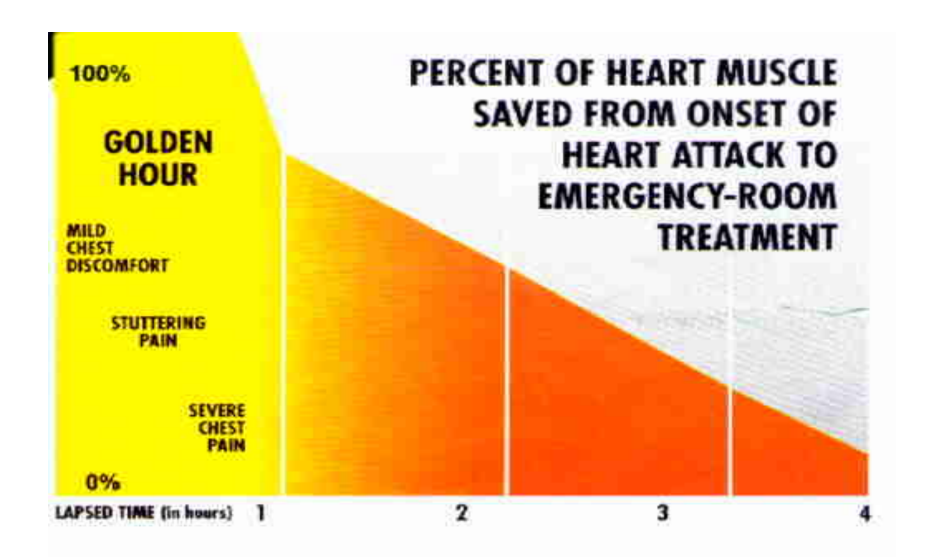
Active peptic ulcer

Current use of anticoagulants: the higher the INR, the higher the risk of bleeding

For streptokinase/anistreplase: prior exposure (more than 5 days previously) or prior allergic reaction to these agents

PRIMARY PCI





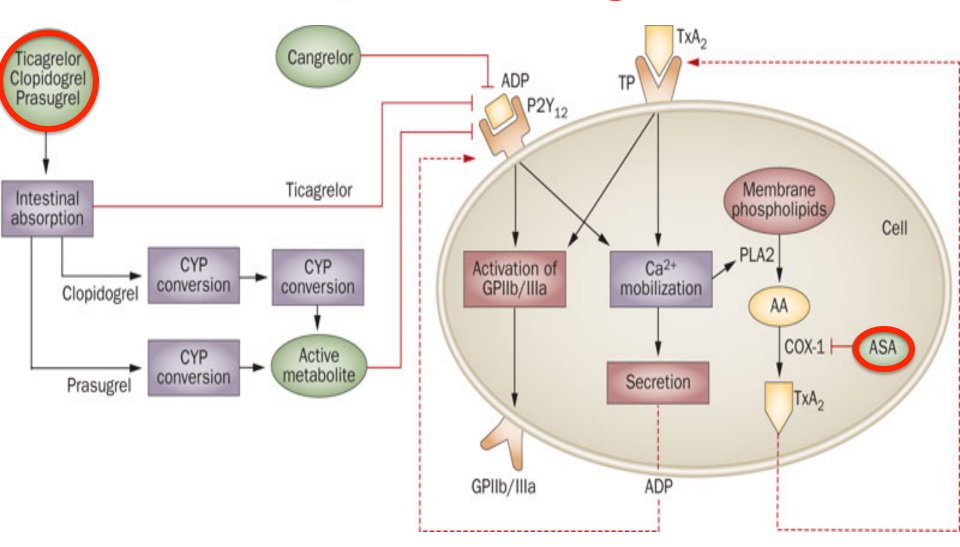






Door to needle time <30min
Door to balloon time <90min

Antiplatelet Agents



Aspirin (ASA)

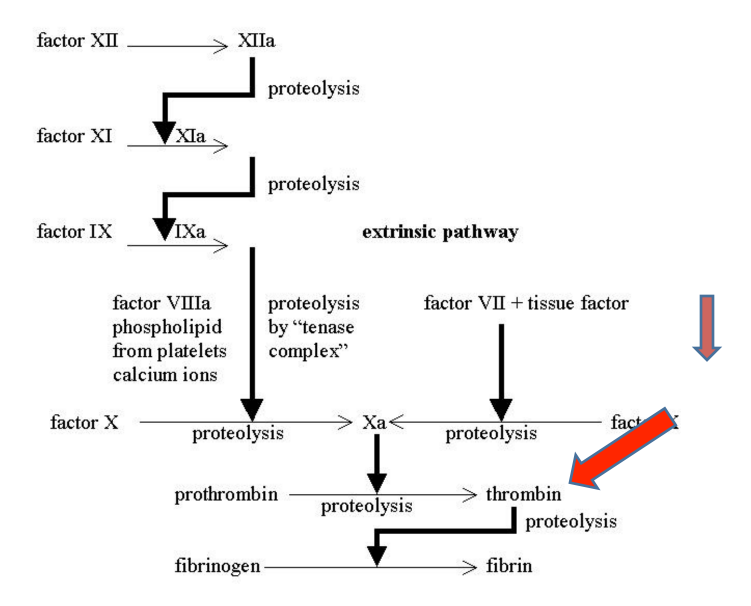
 ASA is Chewable 160 to 325 mg at presentation, then 75 to 325 mg daily.

P2Y₁₂ inhibitors

- More potent than ASA and is combined with ASA
- Both agents are powerful adjuncts to reperfusion therapy
- Examples:
 - Clopidogrel
 - 2. Ticagrelor
 - 3. Prasugrel

ANTITHROMBOTICS

intrinsic pathway

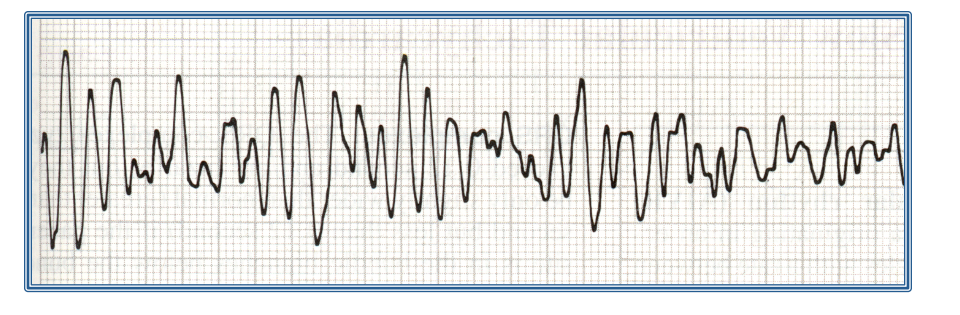


Antithrombotics

- Heparin
 - Unfractionated
 - Low molecular
- Prevents further thrombosis and aids in insuring patency of the occluded artery.

Back to Rashed in ER

- Was given chewable ASA
- 3 Sublingual Nitro tablets
- Just prior to arrival to cardiac cath lab for primary PCI, he lost consciousness



Ventricular Fibrillation

Complications of MI

- Electrical (Arrhythmia)
- Heart failure (Pulmonary Edema)
- Cardiogenic Shock
- Mechanical complications (usually occurs late after MI ...days to weeks)

Summery

- Plaque vulnerability is affected by an inflammatory process
- Acute coronary syndromes is a spectrum and is classified according to markers of ST changes and Myocardial necrosis
- In STEMI, time to reperfusion is critical in myocardial salvage (time is muscle)