ISCHEMIC HEART DISEASE

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25-9-2023

A 60 YEAR OLD MALE, SMOKER, DM PRESENTED C/O CHEST PAIN OF 6 MONTHS DURATTION

Retrosternal Heavy PPT by exertion Relieved by rest Last about 5 minutes

DIAGNOSIS:

Ischemic Heart Disease (IHD)

Basic: coronary circulation Myocardial oxygen supply

What is IHD Causes of IHD

Manifestations of IHD

Treatment

Cardiovascular Disease is the Leading Cause of Death Worldwide¹ **HIV/AIDS** 5 **Pulmonary disease** 7 Injuries 9 Cancer 13 Infectious and 19 parasitic diseases Cardiovascular disease* 29 10 5 15 20 25 30 0 Percentage of total deaths in 2002

*Ischemic heart disease, cerebrovascular disease, hypertensive heart disease, inflammatory heart disease and rheumatic heart disease

1. The World Health Report 2004. WHO Geneva, 2004. Available at: http://www.who.int/whr/2004/en/. Accessed January 2006.

Heart Anatomy

The heart is about the size of a fist and weighs 300-450 gm

The average beat per minute is 70

 The average adult heart pumps about 6000-7500 liters of blood per day.

Coronary Anatomy



Inferior II, III, aVF

Coronary Circulation physiology

1- Flow during basal cardiac circulation: 70-80 ml/min/100gm

2- Flow during maximal cardiac work: 300-400ml/min/100gm

3- High oxygen extraction: 65%-75% (fixed)

4-80% of coronary flow occurs in diastole

5- Collateral pathways is anatomically present but not functioning

6- Cardiac tissue hypoxia is the potent stimulus to open the collateral

Ischemic Heart Disease

Myocardial oxgen demand

- 1- Heart rate
- **2- Contractility**
- 3- Wall tension
- 4- Muscle mass (wall thickness)

Myocardial Oxygen supply

- 1- Coronary flow (patency of coronary artery)
- 2- Hemoglobuline level
- 3- Myocardial oxygen extraction
- 4- Arterial oxygen saturation

Physiology of coronary circulation <u>Myocardial ischemia: imbalance between oxygen</u> <u>supply and demand</u>

Myocardial oxygen demand:

- 1- Heart rate
- 2- Contractility
- 3- Wall tension
- 4- Muscle mass (wall thickness)

Myocardial Oxygen supply:

1- Coronary flow (patency of coronary artery)

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CAUSES OF Myocardial ischemia

Reduced Myocardial O2 Supply

1-Coronary artery disease (atherosclerosis and nonatherosclerosis)

2-decrease flow of oxygenated blood: Sever Anemia carboxyhemoglobulinemia Hypotension Increased Myocardial O2 Demand

1-Left Ventricular Hypertrophy: hypertension aortic stenosis hypertrophic cardiomyopathy

2- Increase cardiac output: Thyrotoxicosis Rapid Tachyarrhythmias

Causes of coronary artery disease

Atherosclerosis 95%	Nonatherosclerosis
	1-Arteritis
Risk factors	(SLE,RA,Takayasu ,)
	2-Embolism
	3-Coronary mural thickening (amyloidosis,radiation therapy,)
	4-Coronary luminal
	aortic dissection
	5-Congenital coronary artery anomalies

Risk Factors for Cardiovascular Disease

Modifiable

- Hypertension
- Smoking
- Hyperlipidaemia
 - Raised LDL-C
 - Low HDL-C
 - Raised triglycerides
- Diabetes mellitus
- Dietary factors
- Lack of exercise
- Obesity
- Homocysteinemia
- Lipoprotein a
- Gout
- Thrombogenic factors: fibrinogen, factors V,VII
- Excess alcohol consumption

Non-modifiable

- Personal history of CVD
- Family history of CVD
- Age: M>45, F>55
- Gender M>F (Premenopausal)
- Personality type A
- Genetic factors: ACE gene





Prevalence of Modifiable Conventional Risk Factors in Patients With CAD in Jordan



95% of Patients With CAD in Jordan Have at least one of the Modifiable Conventional Risk Factors

Number of conventional risk factors in individuals with CAD in Jordan



Hammoudeh et al. [JoHARTS 2] European Heart Journal, September 2005 International Journal of Cardiology, July 2006

Levels of Risk Associated with Smoking, Hypertension and Hypercholesterolaemia. Exponential effect



Serum cholesterol level (8.5 mmol/L, 330 mg/dL)

Pathogenesis of Atherosclerotic Plagues (mechanical shear stresses, biochemical, immunological, inflammation, genetics abnormalities) Endothelial damage (Dysfunction) Protective response results in production of cellular adhesion molecules (Cytokines, Chemokines, Growth factors) Monocytes and T lymphocytes attach to 'sticky' surface of endothelial cells Migrate through arterial wall to subendothelial space Macrophages take up oxidised LDL-C Lipid-rich foam cells Fatty streak and plaque

The 'Activated' Endothelium

`activated' endothelium

cytokines (eg. IL-1, TNF-\alpha)

chemokines (eg.MCP-1, IL-8)

growth factors (eg. PDGF, FGF)

CELLULAR ADHESION MOLECULES

attracts monocytes and T lymphocytes which adhere to endothelial cells induces cell proliferation and a prothrombic state

Endothelial Dysfunction in Atherosclerosis

4



Fatty Streak Formation in Atherosclerosis



Formation of the Complicated Atherosclerotic Plaque



Characteristics of Unstable and Stable Plaque



Cardiovascular risk factors and the stages of atherosclerotic plaque development

Grade I	Grade II	Grade III	Grade IV	Grade V	Grade VI
	209				
Foam cells	Fatty streak	Extracellular fatty streak	Lipid core	Atherosclerotic plaque lipid core embedded in fibrosis	Complicated atherosclerotic plaque (plaque rupture, thrombosis, hemorrhage)
 Intra- and extracellular accumulation of lipids Formation of lipid core 				Development of fibrosis surrounding lipid core • Plaque growth • Atherothrombosi • Plaque rupture	
Asymptomatic				─────────────────────────────────────	
AHA-Classification					
TIMI 28				COmmunication Control	



Major Manifestations of Atherothrombosis



Cerebrovascular disease

Coronary artery disease

Renal artery stenosis

Visceral arterial disease

Peripheral arterial disease (PAD)

The Spectrum of presentations Myocardial Ischemia



Adapted from Cannon CP. *Contemporary Diagnosis and Management of Acute Coronary Syndromes*. 2nd ed. Newtown, PA: Handbooks in Health Care Co.; 2008.

Clinical Manifestations of Atherosclerosis

Coronary heart disease

Asymptomatic, Angina pectoris, variant angina, unstable angina, myocardial infarction, congestive heart failure (CHF), arrhythmias, and sudden cardiac death.

Asympt



Cerebrovascular disease

Transient ischaemic attack, stroke

Peripheral vascular disease

 Intermittent claudication, gangrene, cold feet, painful feet, impotence

IHD-clinicopathological correlation

1- stable angina: stenosis > 70% luminal narrowing

2-variant angina: increase coronay tone 30% normal coronaries

3-unstable angina: rupture plaque subocclusive thrombus (incomplete occlusion) progress to myocardial infarction 15-30%

4-myocardial infarction: rupture plaque occlusive thrombus (complete occlusion)

Stable angina

Commonest form of angina Causes: imbalance between demand and supply Symptom: chest pain Location: central chest (others) Radiation: arm(s), neck, jaw Character : squeezing, pressure, heaviness,... **Duration: 2-10 minutes** Precipitating factors: exertion, emotional upset, heavy meal, sexual intercourse, cold weather **Relieving factors: nitrate, rest** Associated symptoms: dyspnea, diaphoresis, nausea Classes of angina: 1-4

Physical Examination: normal, sign of risk factors, peripheral vascular disease





Angina Chest Pain:

Clinical Diagnosis



CAUSES OF ANGINA

Reduced Myocardial O2 Supply 1-Coronary artery disease

Increased Myocardial O2 Demand

1-Left Ventricular Hypertrophy: hypertension aortic stenosis hypertrophic cardiomyopathy

2-Sever Anemia < 9 gm/dl

2- Rapid Tachyarrhythmias

NYHA Grading of cardiac symptoms (angina / dyspnea)

Grade 1:

Cardiac disease without resulting limitation of physical activity. Ordinary physical activity does not cause chest pain (dyspnea).

Grade 2: Slight limitation of physical activity. Comfortable at rest. Ordinary physical activity result in chest pain (dyspea).

Grade 3: moderate limitation in physical activity. Comfortable at rest. Less than ordinary activity causes symptoms

Grade 4: sever limitation: symptoms at rest.

Stable angina-Diagnosis

- History : angina pectoris is clinical diagnosis
- Physical exam
- Electrocardiogram: 12 ECG, 24 ECG
- Stress ECG : diagnostic and prognostic information
- Radioactive studies: thalium scan,...
- Echocardiography
- CT Coronary angiography
- Serum lipid(LDL, HDL, TG), FBG, CBC
- Coronary angiography

Types of stress test

Exercise tolerance test: ST segment depression

 Exercise or dobutamine Echocardiogram: Wall motion abnormalities

 Exercise or dipyridamole Thallium: Decrease uptake of the nuclear isotope during exercise

Bruce Protocol for Treadmill Testing

STAGE	TIME	SPEED (mph)	GRADE (%)	METS
REST	00.00	0.0	0.0	1.0
1	03.00	1.7	10.0	4.6
2	03.00	2.5	12.0	7.0
3	03.00	3.4	14.0	10.1
4	03.00	4.2	16.0	12.9
5	03.00	5.0	18.0	15.1
6	03.00	5.5	20.0	16.9
7	03.00	6.8	22.0	19.2

Resting Electrocardiogram







Types of stress test

 Exercise or dipyridamole Thallium: Decrease uptake of the nuclear isotope during exercise

Normal Myocardial Perfusion



Myocardial Ischemia



Myocardial Infarction



Types of stress test

Exercise or dobutamine Echocardiogram: Wall motion abnormalities



Imaging Techniques Used to Assess Atherosclerosis

Invasive techniques

- Coronary angiography
- Intravascular ultrasound (IVUS)

Non-invasive techniques

- Magnetic resonance imaging (MRI)
- Computed tomography (CT)
- Ultrasound (B-mode)

Intravascular Ultrasound (IVUS) Showing Atheromatous Plaque

Angiogram



Reproduced from *Circulation* 2001;**103**:604–616, with permission from Lippincott William & Wilkins.

Cardiac CATH



CARDIAC CT ANGIOGRAPHY

Management goals of stable angina

To improve prognosis (mortality reduction)

- Modification of risk factors
- Aspirin
- Lipid-lowering therapy
- ACE-Inhibitor
- Revascularization procedures (PTCA, CABG)

To decrease anginal symptoms

Medical treatment

Treatment of stable angina

1- General measures

2- Medical therapy: Increase O2 supply Decrease O2 demand

3-Revasularization: PCI (percutaneous coronary intervension) CABG (coronary artery bypass grafting)

TREATMENT OF STABLE ANGINA General Measures

- Correction of established risk factors(reversible)
- weight reduction (ideal body weight)
 - Areobic exercise: improve functional capacity, well-being sensation

Treatment of: anemia, thyrotoxicosis, arrhythmias,...

MEDICAL THERAPY OF STABLE ANGINA Prognostic: Aspirin, Statines, ACEI

Symptomatic: Nitrate, B-, CA-blocker, (nicorandil, ranolazine, ivabradine)

INCREASE O2 Supply

1-Increase diastolic time: B-blocker
2-Decrease coronary tone: nitrate, ca-blocker
3-Decrease LV diastolic pressure: nitrate
4-Correct coronary stenosis: PCI, CABG
5-Increase O2 capacity of blood:

transfusion if anemia

DECREASE O2 Demand

 1-Decrease heart rate: B-blocker, ca-blocker
 2-Decrease contractility: B-blocker, ca-blocker
 3- Decrease wall tension (LV pressure and cavity radius): nitrate
 4- metabolic: trimetazidine

Treatment in practice

1-General measures
2-Aspirin
3-Nitrate: S/L, Oral, dermal
3-B-blocker
4-Statins: LDL>100 mg/dl(70mg/dl)
5-Ca-blocker
6-Angio: PTCA,CABG

New medical and invasive therapies for refractory angina

Inhibition of fatty acid metabolism: trimetazidine Potassium channel activators: Nicorandil. Ranolazine: interact with sodium channel Ivabradine: SA inhibitor Endothelin Receptor Blockers: bosentan Testosteron: improve endoth dysfunction. Enhanced external balloon counterpulsation Spinal cord stimulation. Laser revascularization, angiogenesis. **Prognostic Indicators of Coronary Artery Disease:**

1- Left Ventricular Function:
 Normal: 50-75%
 < 50% associated with increased mortality

2- Vessel(s) involved: severity and extent of ischemia

mortality/year 2% single vessel-----12% left main stem

Differential diagnosis of angina

1- Neuromuscular disorder
 2- Respiratory disorders
 3-Upper GI disorder
 4- Psychological
 5- Syndrome X

Cardiac Syndrome X

Typical, exertional angina with positive exercise stress test

Anatomically normal coronary arteries Reduced capacity of vasodilation in microvasculature

F>M

Young > Elderly Excellent prognosis Antianginal therapy is rarely effective

Long term prognosis very good

Case History

A 45 Year old male, presented with recurrent attcaks of chest pain last few mintes, during attacks of chest pain the ECG abnormalities as attached ECG. After pain subside the ECG back to normal.

What is the diagnosis?

During chest pain



After sublingual isosorbid dinitrate tablet



Case History

A 45 Year old male, presented with recurrent attcaks of chest pain last few mintes, during attacks of chest pain the ECG abnormalities as attached ECG. After pain subside the ECG back to normal.

What is the diagnosis?

VARIANT ANGINA-PRINZMETAL ANGINA

Chest pain with ST-Segment elevation Usually at rest, at night Troponin: negative Female > male

Spasm of large epicardial coronary vessel during the attack transmural ischemia 70% on top of atherosclerosis Vasospastic symptoms in other organs: Migraine, rhynauds

Can cause arrhythmias and death

Treatment: CA-blocker, Nitrate

B-blocker is contraindicated

Prognosis: 5 year mortality < 5%

Thank you

