#### Heart Failure

**Etiology And Diagnosis** 

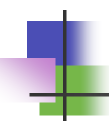
Dr Hanan ALBackr

### Definition:

Heart failure (HF) is a complex clinical syndrome that can result from any structural or functional cardiac disorder that impairs the ability of the ventricle to fill with or eject blood.

#### Prevalence

- Prevalence 0.4-2% overall, 3-5 % in over 65s, 10% of over 80s
- Commonest medical reason for admission
- Annual mortality of 60% over 80s
- > 10% also have AF
- Progressive condition median survival 5 years after diagnosis



#### REMEMBER LEFT VENTRICULAR FAILURE IS A TRUE LIFE THREATENING EMERGENCY



#### Etiology

- It is a common end point for many diseases of cardiovascular system
- It can be caused by :
  - -Inappropriate work load (volume or pressure overload)
  - -Restricted filling
  - -Myocyte loss

# Causes of left ventricular failure

• Volume over load: Regurgitate valve

High output status

• Pressure overload: Systemic hypertension

**Outflow obstruction** 

• Loss of muscles: Post MI, Chronic ischemia

Connective tissue diseases

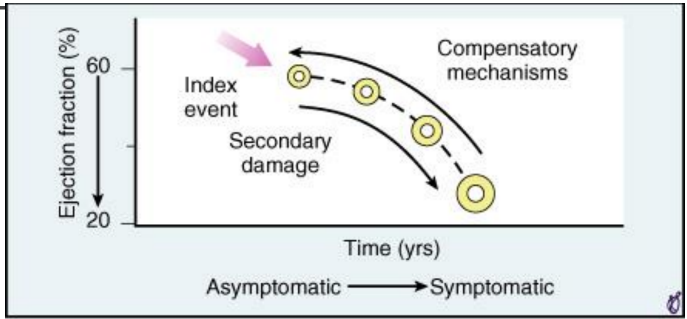
Infection, Poisons

(alcohol,cobalt,Doxorubicin)

• Restricted Filling: Pericardial diseases, Restrictive

cardiomyopathy, tachyarrhythmia

3ackground



- Heart failure pathophysiology
  - Index event
  - Compensatory mechanisms
  - Maladaptive mechanisms

## Pathophysiology

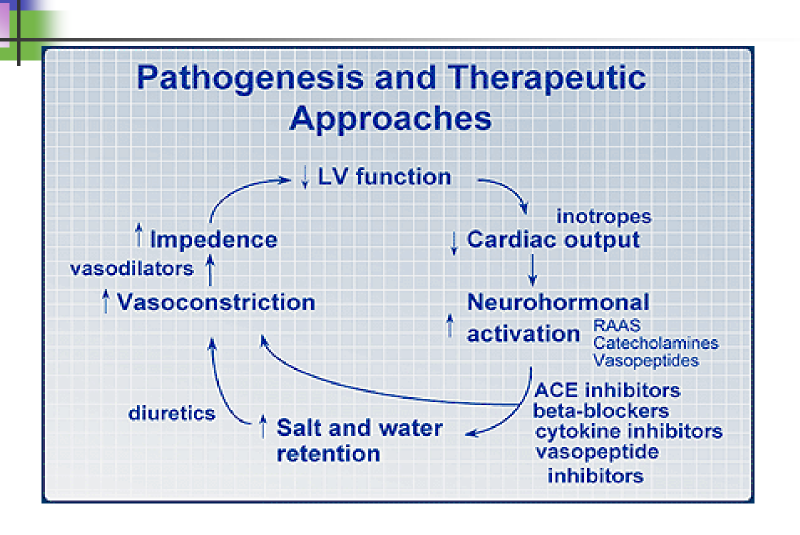
Hemodynamic changes

Neurohormonal changes

Cellular changes

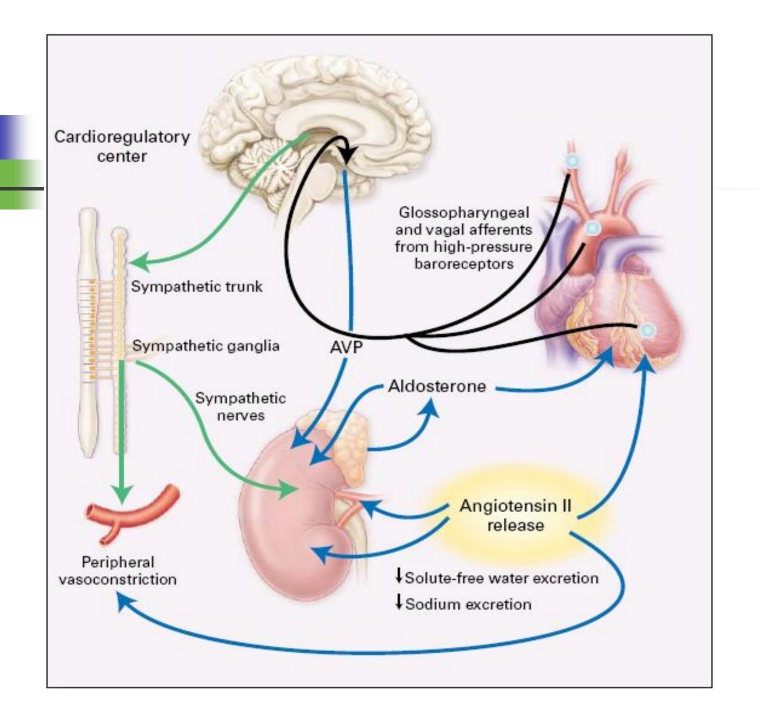
# Hemodynamic changes

 From hemodynamic stand point HF can be secondary to systolic dysfunction or diastolic dysfunction



#### Neurohormonal changes

N/H changes	Favorable effect	Unfavor. effect
↑ Sympathetic activity	<ul><li>↑ HR ,↑ contractility,</li><li>vasoconst. → ↑ V return,</li><li>↑ filling</li></ul>	Arteriolar constriction $\rightarrow$ After load $\rightarrow \uparrow$ workload $\rightarrow \uparrow$ O <sub>2</sub> consumption
↑ Renin-Angiotensin – Aldosterone	Salt & water retention→↑ VR	Vasoconstriction →  ↑ after load
↑ Vasopressin	Same effect	Same effect
↑ interleukins &TNFα	May have roles in myocyte hypertrophy	Apoptosis
↑Endothelin	Vasoconstriction→↑ VR	↑ After load



### **Symptoms**

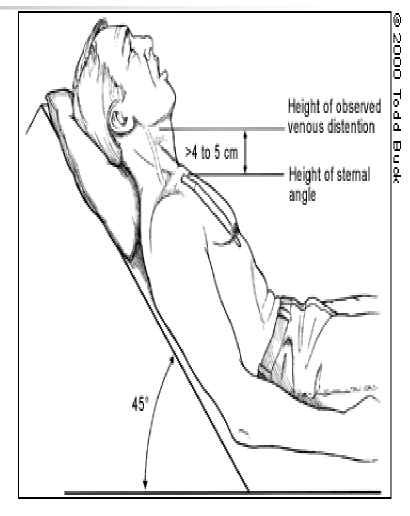
 SOB, Orthopnea, paroxysmal nocturnal dyspnea

Low cardiac output symptoms

 Abdominal symptoms: Anorexia, nausea, abdominal fullness, Rt hypochondrial pain

#### Physical Signs

- High diastolic BP & occasional decrease in systolic BP (decapitated BP)
- JVD
- Rales (Inspiratory)
- Displaced and sustained apical impulses
- Third heart sound low pitched sound that is heard during rapid filling of ventricle



### Physical signs (cont.)

 Mechanism of S<sub>3</sub> sudden deceleration of blood as elastic limits of the ventricles are reached

Vibration of the ventricular wall by blood filling

Common in children

## Physical signs (cont.)

- Fourth heart Sound (S<sub>4</sub>)
  - Usually at the end of diastole
  - Exact mechanism is not known
     Could be due to contraction of atrium against stiff ventricle

Pale, cold sweaty skin





- Systolic & Diastolic
- High Output Failure
  - Pregnancy, anemia, thyrotoxisis, A/V fistula, Beriberi, Pagets disease
- Low Output Failure
- Acute
  - large MI, aortic valve dysfunction---
- Chronic

# Forms of heart failure ( cont.)

### Right vs Left sided heart failure: Right sided heart failure:

Most common cause is left sided failure

Other causes included: Pulmonary embolisms

Other causes of pulmonary htn.

**RV** infarction

MS

Usually presents with: LL edema, ascites

hepatic congestion

cardiac cirrhosis (on the long run)

### Differential diagnosis

- Pericardial diseases
- Liver diseases
- Nephrotic syndrome
- Protein losing enteropathy



#### Laboratory Findings

- Anemia
- Hyperthyroid
- Chronic renal insuffiency, electrolytes abnormality
- Pre-renal azotemia
- Hemochromatosis

### Electrocardiogram

- Old MI or recent MI
- Arrhythmia
- Some forms of Cardiomyopathy are tachycardia related
- LBBB→*may help in management*

## Chest X-ray

- Size and shape of heart
- Evidence of pulmonary venous congestion (dilated or upper lobe veins → perivascular edema)
- Pleural effusion

### Echocardiogram

- Function of both ventricles
- Wall motion abnormality that may signify CAD
- Valvular abnormality
- Intra-cardiac shunts

## Cardiac Catheterization

When CAD or valvular is suspected

If heart transplant is indicated

In conclusion, congestive heart failure is often assumed to be a disease when in fact it is a syndrome caused by multiple disorders.

### TREATMENT

- Correction of reversible causes
  - Ischemia
  - Valvular heart disease
  - Thyrotoxicosis and other high output status
  - Shunts
  - Arrhythmia
    - A fib, flutter, PJRT
  - Medications
    - Ca channel blockers, some antiarrhythmics

### Diet and Activity

- Salt restriction
- Fluid restriction
- Daily weight (tailor therapy)
- Gradual exertion programs

#### Diuretic Therapy

- The most effective symptomatic relief
- Mild symptoms
  - HCTZ
  - Chlorthalidone
  - Metolazone
  - Block Na reabsorbtion in loop of henle and distal convoluted tubules
  - Thiazides are ineffective with GFR < 30 --/min</p>

#### Diuretics (cont.)

#### Side Effects

- Pre-renal azotemia
- Skin rashes
- Neutropenia
- Thrombocytopenia
- Hyperglycemia
- ↑ Uric Acid
- Hepatic dysfunction

#### Diuretics (cont.)

- More severe heart failure → loop diuretics
  - **Lasix** (20 320 mg QD), Furosemide
  - Bumex (Bumetanide 1-8mg)
  - Torsemide (20-200mg)

**Mechanism of action**: Inhibit chloride reabsortion in ascending limb of loop of Henle results in natriuresis, kaliuresis and metabolic alkalosis

#### **Adverse reaction:**

pre-renal azotemia Hypokalemia Skin rash ototoxicity

#### K+ Sparing Agents

- Triamterene & amiloride acts on distal tubules to ↓ K secretion
- Spironolactone (Aldosterone inhibitor)

recent evidence suggests that it may improve survival in CHF patients due to the effect on reninangiotensin-aldosterone system with subsequent effect on myocardial remodeling and fibrosis



- Renin-angiotensin-aldosterone system is activation early in the course of heart failure and plays an important role in the progression of the syndrome
- Angiotensin converting enzyme inhibitors
- Angiotensin receptors blockers
- Spironolactone

#### Angiotensin Converting Enzyme Inhibitors

- They block the R-A-A system by inhibiting the conversion of angiotensin I to angiotensin II → vasodilation and ↓ Na retention
- ↓ Bradykinin degradation ↑ its level → ↑ PG secretion & nitric oxide
- Ace Inhibitors were found to improve survival in CHF patients
  - Delay onset & progression of HF in pts with asymptomatic LV dysfunction
  - ↓ cardiac remodeling

# Side effects of ACE inhibitors

- Angioedema
- Hypotension
- Renal insuffiency
- Rash
- cough

# Angiotensin II receptor blockers

Has comparable effect to ACE I

 Can be used in certain conditions when ACE I are contraindicated (angioneurotic edema, cough)

#### Digitalis Glycosides (Digoxin, Digitoxin)

- The role of digitalis has declined somewhat because of safety concern
- Recent studies have shown that digitals does not affect mortality in CHF patients but causes significant
  - Reduction in hospitalization
  - Reduction in symptoms of HF

#### **Digitalis** (cont.) Mechanism of Action

- +ve inotropic effect by ↑ intracellular Ca & enhancing actin-myosin cross bride formation (binds to the Na-K ATPase → inhibits Na pump → ↑ intracellular Na → ↑ Na-Ca exchange
- Vagotonic effect
- Arrhythmogenic effect

#### Digitalis Toxicity

- Narrow therapeutic to toxic ratio
- Non cardiac manifestations

Anorexia,

Nausea, vomiting,

Headache,

Xanthopsia sotoma,

Disorientation

#### Digitalis Toxicity

#### Cardiac manifestations

- Sinus bradycardia and arrest
- A/V block (usually 2<sup>nd</sup> degree)
- Atrial tachycardia with A/V Block
- Development of junctional rhythm in patients with a fib
- PVC's, VT/ V fib (bi-directional VT)

#### β Blockers

- Has been traditionally contraindicated in pts with CHF
- Now they are the main stay in treatment on CHF & may be the only medication that shows substantial improvement in LV function
- In addition to improved LV function multiple studies show improved survival
- The only contraindication is severe decompensated CHF

### Antiarrhythmics

- Most common cause of SCD in these patients is ventricular tachyarrhythmia
- Patients with h/o sustained VT or SCD → ICD implant

## New Methods

Implantable ventricular assist devices

Biventricular pacing (only in patient with LBBB & CHF)

Artificial Heart

#### Cardiac Transplant

It has become more widely used since the advances in immunosuppressive treatment

- Survival rate
  - 1 year 80% 90%
  - 5 years 70%

#### **Prognosis**

- Annual mortality rate depends on patients symptoms and LV function
- 5% in patients with mild symptoms and mild
   in LV function
- 30% to 50% in patient with advances LV dysfunction and severe symptoms
- 40% 50% of death is due to SCD