

### Introduction:

- > most common CVS disease
- > elevated arterial BP damages vessels in renal, heart and brain

**Complications:** Renal failure, Coronary disease, Heart failure, Stroke, Dementia

### Classification:

Classification:	Systolic BP	Diastolic BP:
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Normal:	<120	<80
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Pre-hypertension:	120-139	80-89
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Stage 1 HTN:	140-159	90-99
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Stage 2 HTN:	>= 160	>=100
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### Types of Hypertension:

**Essential Hypertension (85-90%):** Primary hypertension, no identifiable cause (genetic), can't be cured, can be controlled

**Secondary Hypertension (10-15%):** Specific identified cause (comorbid disease or drug), can be cured when cause is eliminated

### Secondary Causes of HTN:

- > Genetic factors
- > Psychological stress
- > Environmental and Dietary Factors: high salt diet, decreased calcium and phosphate intake, sedentary lifestyle

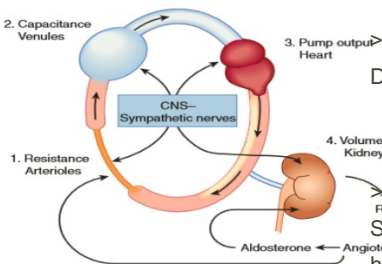
> renal, endocrine, Diseases: vascular, renal diseases

- > Drugs: sympathomimetic amines, amphetamines, oral decongestants (eg. pseudoephedrine), corticosteroids, osteogens (COCs), NSAIDs, COX-2 inhibitors

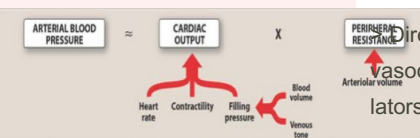
### Clinical presentation:

- Often incidental finding
- Adults should get BP checks once a year
- Severe cases: **Headaches**, visual disturbances, target organ damage (stroke, ischemic heart disease, renal failure)

### Normal regulation of BP:



### Factors Influencing BP:



### Potential Mechanisms of Pathogenesis:

$$BP = CO \times PVR$$

**Increased Cardiac output:** Increased fluid volume from excessive sodium intake or renal sodium retention. Venous constriction: due to excess stimulation of RAAS

**Increased Peripheral resistance:** Excess stimulation of RAAS Sympathetic nervous system over-activity

### Classes of Antihypertensive Agents:

**Diuretics:** Reduce blood volume=Depletes the body of sodium, Venodilation. Reduce peripheral vascular resistance, Inhibit cardiac function, Increase venous pooling capacitance vessels

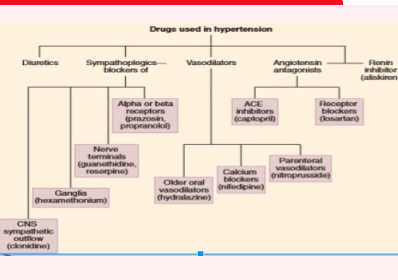
**Direct Vasodilators:** Reduce peripheral vascular resistance, Increase venous pooling capacitance vessels

> **Angiotensin antagonists:** Reduce peripheral vascular resistance, Reduce blood volume

### Diuretics:

- Reduce blood volume and cardiac output
- Cardiac output returns to normal
- But peripheral vascular resistance declines
- Sodium contributes to vascular resistance = Increase vessel stiffness

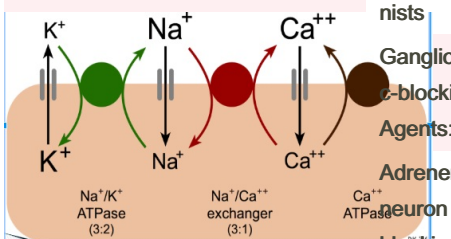
### Antihypertensive Drugs:



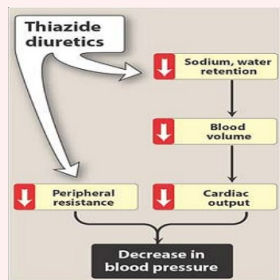
### Diuretics: (cont)

- Altered sodium-sodium  
Venodi calcium exchange  
lation

### Vasodilation mechanism by Diuretics:



### Thiazide Diuretics:



### Use of Diuretics:

> Lower BP by 10-15 mmHg in most patients

> Thiazide diuretics –mild to moderate HTN

> Loop diuretics – severe HTN and hypertensive emergencies

### Adverse Effects (Thiazide Diuretics):

Potassium depletion – hypokalemia

Magnesium depletion

Hyperuricemia- gouty attacks

Glucose intolerance

Increase serum lipid concentrations

### Sympathoplegic Drugs:

**Centrally-acting Drugs:** Methyldopa, clonidine

**Adrenoceptor antagonists:** Beta-blockers, alpha-1 blockers

**Ganglionic blocking Agents:** No longer used clinically ; hexamethonium

**Adrenergic blocking agents:** Block the release of noradrenaline, Reserpine, guanethidine, debrisoquin, Not/rarely used clinically

### Centrally-Acting Drugs:

- Methyl- rarely used  
dopa, except clonidine

- Reduces sympathetic outflow

- Compensatory response mechanism: salt retention

- **Clonidine**, Stimulate central guanabenz, alpha-2 adrenoceptor

- Methyldopa Results in the synthesis of a false neurotransmitter  
– analogue of L-dopa

### Centrally-Acting Drugs: (cont)

Alpha-methylnoradrenaline = Stimulates central alpha-2 adrenoceptors

**Clonidine:** Reduces cardiac output, PVR, relaxation of capacitance vessels

Rarely causes postural hypotension

Adverse effects:

- Dry mouth

- Sedation

Contraindication: Patients with depression

**Caution: Abrupt discontinuation can lead to hypertensive crisis**

### Adrenoceptor antagonists: Beta-blockers

Non-selective: Propranolol

Beta-1 selective: Betaxolol, bisoprolol, esoprolol, "BBEAM" **atenolol, metoprolol**, Cardioselective

Vasodilator beta-blockers: Also block alpha-1 receptors, Labetolol, carvedilol, nebivolol

### Decrease cardiac output

Decrease peripheral vascular resistance

Inhibit stimulation of renin production by catecholamines

Adverse effects= Heart block, bronchoconstriction, diabetes, vivid dreams

### Alpha-1 blockers:

**Prazosin**, terazosin, doxazosin

Block alpha-1 receptors in arterioles and veins

Vasodilation Reduce peripheral resistance

Compensatory mechanism: salt and water retention

More effective when used with other drugs

### Calcium Channel Blockers:

> amlodipine, isradipine, Dihydropyridines: nicardipine, nimodipine, felodipine, nisoldipine, lacidipine

> Verapamil, diltiazem, Non-dihydropyridines: hydralazine

Benzothiazepine (diltiazem)

### Mechanism of action:

- Inhibit calcium influx through voltage-dependent L-type calcium channels  
- Relax arteriolar smooth muscle, reduce peripheral vascular resistance  
- Cause coronary and peripheral vasodilation

### CCB: Mechanisms of Action:

Dihydropyridine CCBs: Primary vasodilators (reduce PVR), All decrease cardiac contractility except amlodipine and felodipine



### CCB: Mechanisms of Action: (cont)

Non-dihydropyridines (diltiazem, verapamil) directly block the AV node, decrease heart rate, decrease cardiac contraction

**Adverse effects:** 1. Flushing, peripheral oedema, tachycardia, bradycardia, heartblock

2. Headache, flushing, dizziness, palpitations, hypotension occur within a few hours of dosing, Associated w high initial doses or rapid increase in dose, Common with short-acting preparations

3. Ankle oedema: Due to a rise in intracapillary pressure as a result of selective dilatation of precapillary arterioles, NOT due to sodium retention, Relieved by bed rest

4. Gum Hypertrophy: dihydropyridines

5. GIT: constipation (verapamil), nausea, and vomiting

### Inhibitors of Angiotensin:

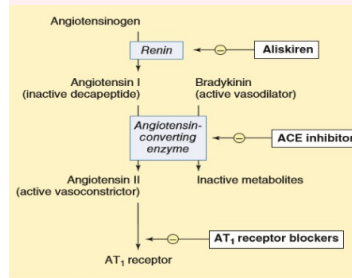
Angiotensin converting enzyme inhibitors (ACEIs)  
 Captopril, enalapril, Ramipril, fosinopril, trandopril

### Inhibitors of Angiotensin: (cont)

Angiotensin receptor blockers (ARBs)  
 Losartan, valsartan, telmisartan, irbesartan, candesartan

Renin-inhibitors  
 Aliskiren

### Inhibitors of Angiotensin:



### Adverse Effects

ACEIs =  
 Dry cough  
 Can cause hyperkalemia – potassium monitoring essential  
 Angioedema (substance P?)

ARBs =

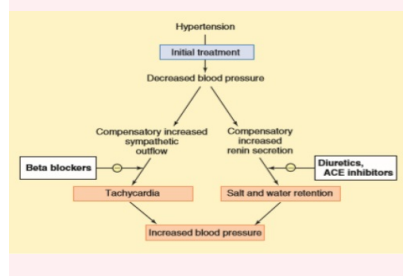
No dry cough

### Hyperkalemia

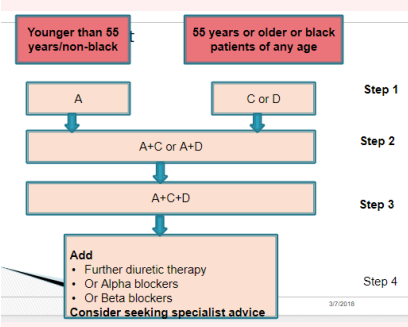
Angiodema is less common than ACEIs

Contraindicated in pregnancy

### Combination Treatment: Vasodilators:



### Management Approach:



### First-line agents: "ACD" drugs:

- A: ACEIs and ARBs
- C: Calcium channel antagonists
- D: Diuretics (Thiazides)

### Second-line agents:

- Beta-adrenoceptor blockers
- Aldosterone antagonists (spironolactone, eplerenone)
- Alpha-blockers (doxazosin, prazosin, terazosin)
- Loop diuretics (frusemide, torsemide)
- Direct vasodilators (hydralazine, minoxidil) [last-line of therapy]
- Central α-2 agonists (clonidine)
- Adrenergic antagonists (reserpine)