Cheatography

Hypertension Drugs Cheat Sheet

by Carm (Carmilaa) via cheatography.com/49544/cs/16839/

Introduction:

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

Compli Renal failure,
cat- Coronary disease,
ions: Heart failure, Stroke,
Diamentia

Classification:

Classific- ation:	Systolic BP	Diastolic BP:
Normal:	<120	<80
Pre-hy- perten- sion:	120- 139	80-89
Stage 1 HTN:	140- 159	90-99
Stage 2	>= 160	>=100

Types of Hypertension:

Essential	Primary hypert-	
Hypert-	ension, no identi-	
ension	fiable cause	
(85-90%):	(genetic), can't be	
	cured, can be	
	controlled	
Secondary	Specific identified	
Hypert-	cause (comorbid	
ension	disease or drug),	
(10-15%):	can be cured when	
	cause is eliminated	

Secondary Causes of HTN:

- > Genetic factors
- > Psychological stress

> Enviro- high salt diet, nmental decreased calcium and and phosphate Dietary intake, sedentary Factors: lifestyle

> renal, endocrine,
Diseases: vascular, renal
diseases

> Drugs: sympathomimetic

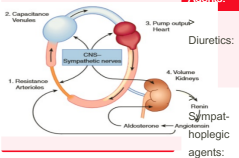
sympathomimetic amines, amphetamines, oral decongestants (eg. pseudoehedrine), corticosteroids, osteogens (C0Cs), NSAIDs, COX-2 inhibitors

Clinical presentation:

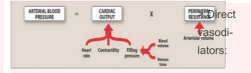
- Often incidental finding
- Adults should get BP checks once a year

- Headaches, visual
Severe disturbances, target
cases: organ damage
(stroke, ischemic
heart disease, renal
failure)

Normal regulation of BP:



Factors Influencing BP:



Potential Mechanisms of Parthogenesis:

BP= CO x PVR

Increased Increased fluid

Cardiac volume from
output: 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-a-

Classes of Antihypertensive

Reduce blood volume=Depletes the body of sodium, Venodilation

Reduce peripheral vascular resistance, Inhibit cardiac function, Increase venous pooling capacitance vessels

Reduce peripheral vascular resistance, Increase venous pooling capacitance vessels

> Angiot- Reduce peripheral
ensin vascular resistance,
antago- Reduce blood
nists: volume

Diuretics:

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

Antihypertensive Drugs:

ctivity





Published 28th August, 2018. Last updated 28th August, 2018. Page 1 of 3. Sponsored by **ApolloPad.com**Everyone has a novel in them. Finish
Yours!

https://apollopad.com



Hypertension Drugs Cheat Sheet

by Carm (Carmilaa) via cheatography.com/49544/cs/16839/

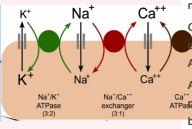
Diuretics: (cont)

- Altered sodium-sodium

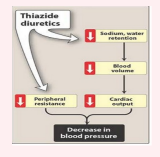
Venodi calcium exchange

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- Methyldopa, acting clonidine

Drugs:

Adreno- Beta-blockers, ceptor alpha-1 blockers antago-

nists

Ganglionic-blocking Agents:

Adrenergic

Ca**
ATPaseuron
blocking
agents:

No longer used clinically; hexamethonium

Block the release of noradrenaline, Reserpine, guanethidine,

debrisoquin,

clinically

Not/rarely used

Centrally-Acting Drugs:

- Methyl- rarely used dopa, except clondine clonidine
- Reduces sympathetic outflow
- Compen-salt retention satory response mechanism:
- Clonidine, Stimulate central guanabenz, alpha-2 adrenoguanfacine ceptors
- Results in the

 Methyldopa synthesis of a

 anologue false neurotranof L-dopa smitter

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: Betablockers

Non-selective: Propranolol

Beta-1 Betaxolol, bisoprselective: olol, esoprolol,

"BBEAM" atenolol, metoprolol,

Cardioselective

Vasodi- Also block alpha-1 lator receptors,Labetolol, beta-b- carvidelol, nebivolol lockers

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

Compensalt and water satory retention mechanism:

More effective when used with other drugs

Calcium Channel Blockers:

> amlodipine, isradipine,
Dihydr nicardipine, nimodiopy- pine, felodipine, nisoldridines ipine, lacidipine

> Verapamil, diltiazem,

Non- hydralazine dihyd-ropyri-

dines

Benzothiazepine (dilti-

Mechanism of action:

azem)

- 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:

Dihydr Primary vasodilators
opyridine decrease cardiac
CCBs contractility except
amlodipine and
felodipine



Published 28th August, 2018. Last updated 28th August, 2018. Page 2 of 3. Sponsored by **ApolloPad.com**Everyone has a novel in them. Finish
Yours!
https://apollopad.com

Cheatography

Hypertension Drugs Cheat Sheet

by Carm (Carmilaa) via cheatography.com/49544/cs/16839/

CCB: Mechanisms of Action: (cont)

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

Adverse effects:

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**: dihydropy-ridines
- 5. GIT: constipation (verapamil), nausea, and vomiting

Inhibitors of Angiotensin:

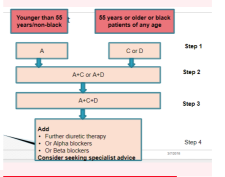
Angiotensin Captopril,
converting enalapril,
enzyme Ramipril,
inhibitors fosinopril,
(ACEIs) trandopril

Inhibitors of Angiotensin: (cont)

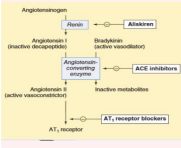
Angiot- Losartan,
ensin valsartan, telmisreceptor artan, irbesartan,
blockers candesartan
(ARBs)

Renin-inh- Aliskiren ibitors

Management Approach:



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

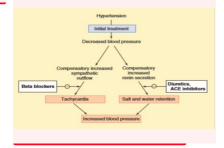
First-line agents: "ACD" drugs:

- 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)

Combination Treatment: Vasodilators:





By Carm (Carmilaa) cheatography.com/carmilaa/

Published 28th August, 2018. Last updated 28th August, 2018. Page 3 of 3.

Sponsored by **ApolloPad.com**Everyone has a novel in them. Finish
Yours!
https://apollopad.com