

Learning Objectives	Identify	Identify risk factors and/or diagnostic indicators that may lead to hypertension.	
	Classify BP as outlined by ACC/AHA		
	Explain	Explain the proper way to take a BP.	
	Define and explain	Define and explain the criteria used to diagnose hypertension.	
	Identify	Identify first line treatment options for treatment of BP in patients and those with compelling indications according to JNC VIII, ACC/AHA	
	Explain	Explain the benefits, adverse drug reactions, interactions, contraindications, and monitoring for alternative treatment options for hypertension	
	Summarize	Summarize counseling points for antihypertensive drug classes.	
	Explain	Explain the rationale for and determine the appropriateness of combination therapy according to JNC VIII.	
	Design	Design a treatment and monitoring plan for patients with hypertension	

# Overview of HTN 75 million American adults have HTN Only about 54% of adults with HTN have the BP under control Persistently elevated blood pressure • Can damage the heart over time Hypertension Trends • Major risk factor for heart attack, stroke and kidney failure • Lifetime risk >90% by age of 55 • Unclear threshold of safety as evidenced by multiple changing recommendations No cure • Managed to minimize complications

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# Fluctuations in blood pressure BP normally follows a circadian rhythm • Lowest values occur during sleep • Starts to rise a few hours prior to awakening • Highest values occur midmorning Blood pressure can increase acutely • Physical activity • Emotional stress

# **Definition of HTN**

# Hypertension (HTN) or high blood pressure (HBP)

- · Patient language:
- Force of your blood moving against the walls of your arteries

# Systolic blood pressure (SBP)

- Peak blood pressure achieved during cardiac contraction (systole)
- Patient language:
- Top Number the pressure in the arteries when the heart beats

# Diastolic blood pressure (DBP)

Minimum pressure achieved in between contractions (diastole)

# Patient Language:

• Bottom Number – the pressure measured between heartbeats

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# Etiology

# **Primary HTN**

Formally referred to as essential HTN

Unknown cause

# Secondary HTN

Known cause

Examples: sleep apnea, CKD, primary aldosteronism

# Primary (essential) hypertension

> 90% of hypertensive patients

Usually results from unknown pathophysiologic etiology

• Several postulated mechanisms

Can't be cured

Genetic factors

• Monogenic and polygenic

Needs to be treated

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# Secondary hypertension

< 10% of hypertensive patients

HTN caused by something else

# **COMMON**

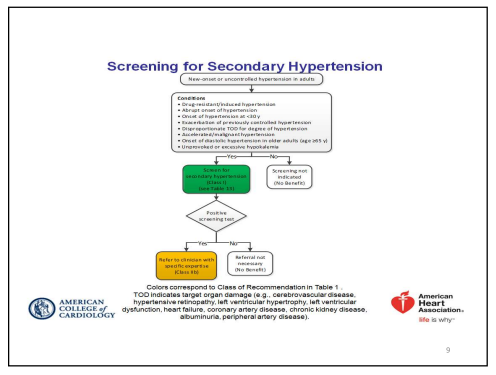
- Renovascular disease. Renal parenchymal disease, Primary aldosteronism
- Obstructive sleep apnea. Drug- or alcohol-induced

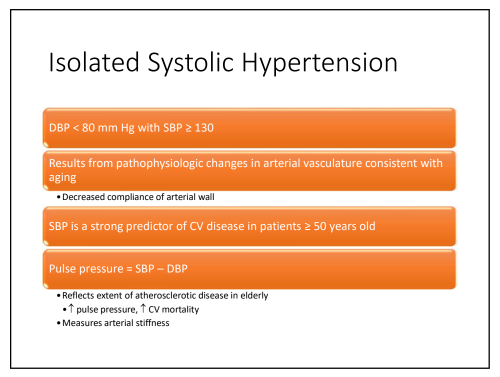
# **UNCOMMON**

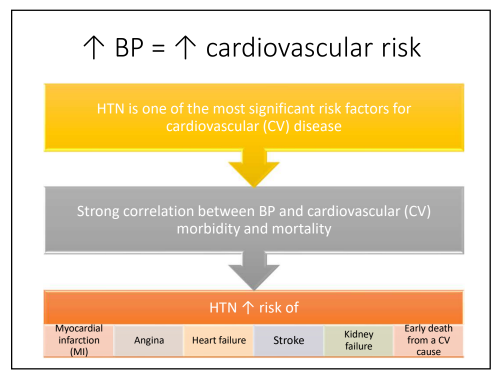
- Pheochromocytoma/ paraganglioma. Cushing's syndrome, Thyroid disease
- Hypo- or Hyperthyroidism
- Coarctation of the aorta (undiagnosed or unrepaired)

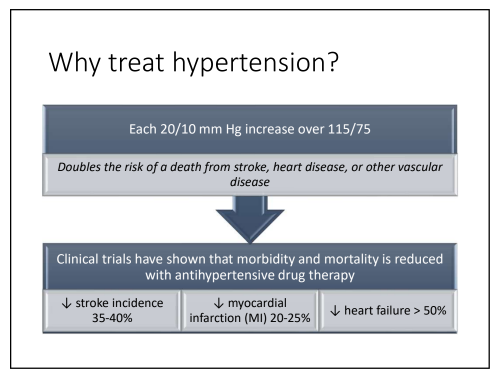
# Management:

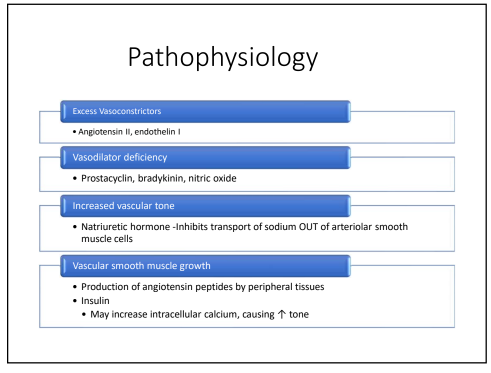
• Treat / correct the underlying comorbid condition!











# Pathophysiology Naintain BP through volume-pressure adaptive mechanism If BP ↓, kidneys increase sodium and water retention, leading to plasma volume expansion and ↑ BP If BP ↑, kidneys excrete more sodium and water to reduce plasma volume and cardiac output, therefore ↓ BP RAAS System

# Pathophysiology

## Renin

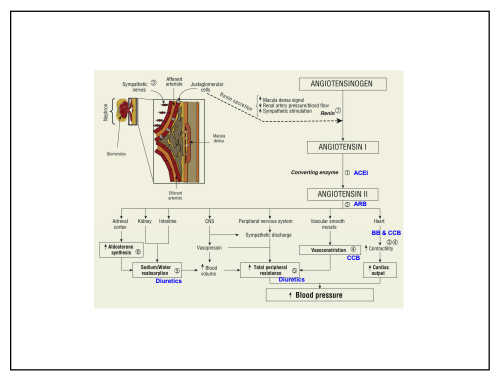
- Stored in juxtaglomerular cell
- Present in afferent arterioles of kidney
- Function as baroreceptor-sensing device
- Released in response to:
  - Intrarenal factors
  - Decreased renal artery pressure/renal blood flow
  - Catecholamine stimulation
- Extrarenal factors
- ullet in sodium and chloride delivered to the distal tubule
- ↓ serum potassium and/or intracellular calcium
- Catalyzes conversion of angiotensinogen to angiotensin I in the blood

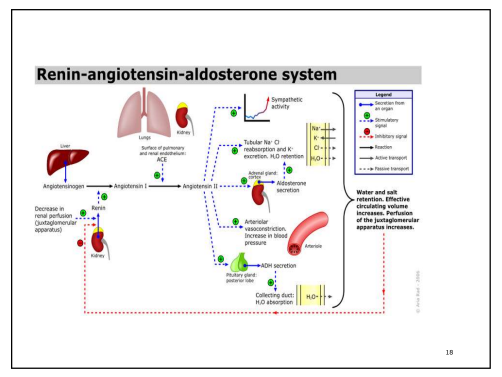
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# Pathophysiology

# Angiotensin

- Vasoconstriction
- Stimulation of catecholamine release
- Centrally mediated increases in sympathetic nervous system activity
- Stimulation of aldosterone synthesis from the adrenal cortex
- Sodium and water reabsorption
- Increases plasma volume, total peripheral resistance, and BP
- Myocardial fibrosis, vascular dysfunction



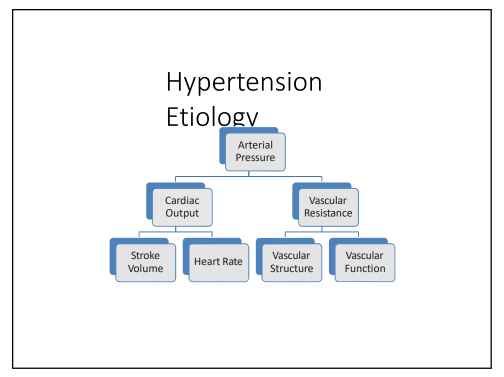


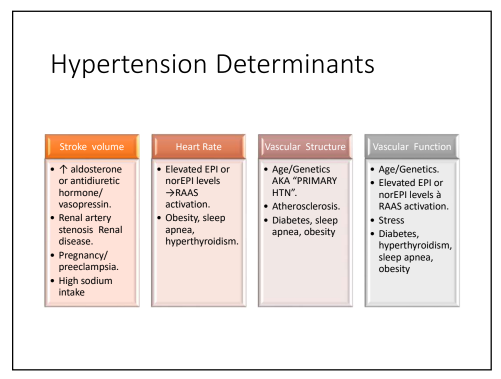
# Pathophysiology of HTN

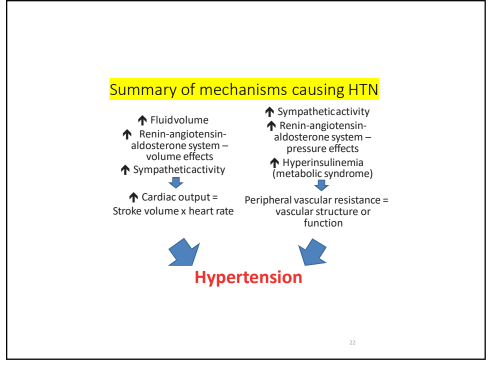
- BP = CO X TPR
  - Cardiac Output = HR X Stroke volume
    - Major determinant of SBP
  - Total peripheral resistance (TPR)
    - Major determinant of DBP
  - Drugs decrease BP by decreasing CO, TPR or both

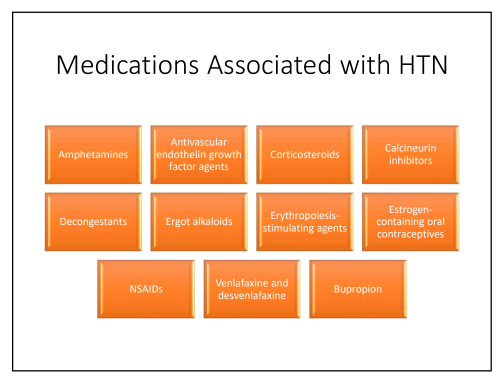


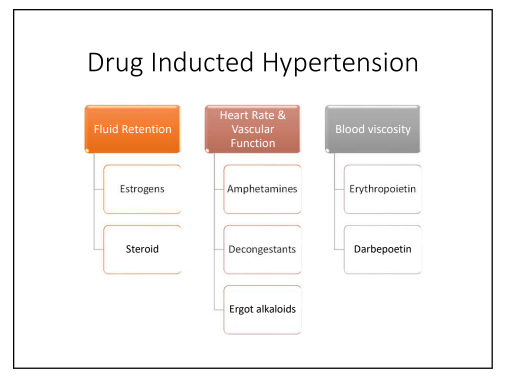
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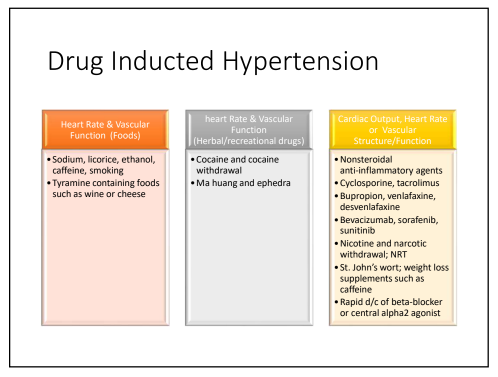


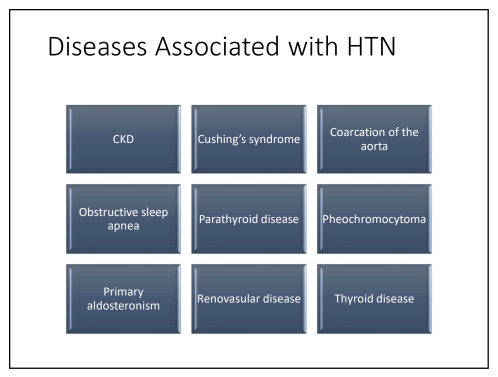


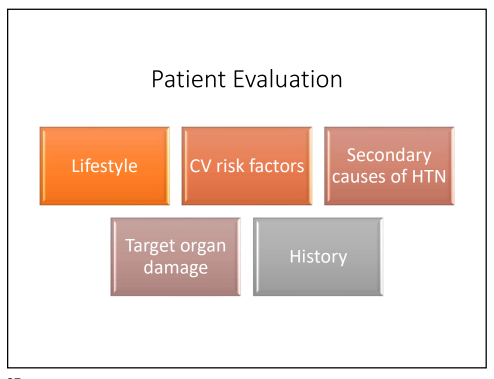












# Metabolic Syndrome

Associated with HTN & metabolic abnormalities such as↑ serum insulin levels

Metabolic syndrome is diagnosed if 3 of the 5 following are present:

- abdominal obesity (>40" men; >35" women)
- HTN (≥130/≥85 or taking antihypertensives)
- elevated fasting glucose (≥100 mg/dL or on diabetes meds)
- elevated TG (≥150 mg/dL or on lipid meds)
- low HDL (<40 mg/dL men; <50 mg/dL women)

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# CVD Risk Factors Common in Patients With Hypertension

### Modifiable Risk Factors

- Current cigarette smoking, secondhand smoking
- Diabetes mellitus
- Dyslipidemia/hypercholesterolemia
- Overweight/obesity
- Physical inactivity/low fitness
- Unhealthy diet

### Relatively Fixed Risk Factors

- CKD
- Family history
- Increased age
- Low socioeconomic/educational status
- Male sex
- Obstructive sleep apnea
- Psychosocial stress

https://www.acc.org/~/media/Non-Clinical/Files-PDFs-Excel-MS-Word-etc/Guidelines/2017/2017-Blood Pressure-Guideline.ppt

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# Basic and Optional Laboratory Tests for Primary Hypertension

Basic testing	Fasting blood glucose*			
	Complete blood count			
	Lipid profile			
	Serum creatinine with eGFR*			
	Serum sodium, potassium, calcium*			
	Thyroid-stimulating hormone			
	Urinalysis			
	Electrocardiogram			
Optional testing	al testing Echocardiogram			
	Uric acid			
	Urinary albumin to creatinine ratio			

\*May be included in a comprehensive metabolic panel. eGFR indicates estimated glomerular filtration rate.



# How is blood pressure measured?

Sphygmomanometer and stethoscope

Measured in millimeters of mercury (mm Hg)

# Systolic blood pressure (SBP)

- Top number; peak value
- Measured during cardiac contraction

# Diastolic blood pressure (DBP)

- Bottom number; nadir value
- Measured after contraction when the cardiac chambers are filling

# Different Readings

Appropriate measurement!

In office readings

Home readings

Ambulatory monitoring

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# Out of Office Monitoring

# **Ambulatory**

- Document BP at frequent time intervals over 8 -24h
- Useful to determine nighttime high BP readings

# Home

- Measurements collected by patients average home BP over 1 week
- Check AM and HS
- · Arm cuffs more accurate than wrist or finger
- FABRICATED readings!
- Accurate if within 5mmHg of in-office reading wait 1 minute between readings

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# **Blood Pressure Measurement**

- Steps for Proper BP Measurement
  - Step 1: Prepare the patient
  - Step 2: User proper technique for BP measurement
  - Step 3: Take the proper measurements needed for diagnosis and treatment of elevated BP/HTN
  - Step 4: Properly document accurate BP readings
  - Step 5: Average the readings
  - Step 6: Provide BP readings to the patient

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# Step 1: Properly prepare the patient

- Have the pt relax, sitting in a chair (feet on floor, back supported) for > 5 min.
- Avoid caffeine, exercise, and smoking for at least 30 minutes before measurement
- Ensure the pt has emptied his/her bladder
- Neither the patient nor the observer should talk during the rest period or during the measurement
- Remove all clothing covering the location of cuff placement
- Note: Measurements made while pt is sitting/lying on examining table do not fulfill these criteria



# Step 2: Use proper technique

- Use a BP measurement device that has been validated, and ensure the device is calibrated periodically
- Support the patient's arm (ex: rest on a desk)
- Position the middle of the cuff on the pt's upper arm at the level of the right atrium (midpoint of the sternum)
- Use the correct cuff size (bladder encircles 80% of the arm).
- Note if larger or smaller than normal cuff size is used
- Either the stethoscope diaphragm or bell may be used for auscultatory readings

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# | Step 3: Take proper measurements

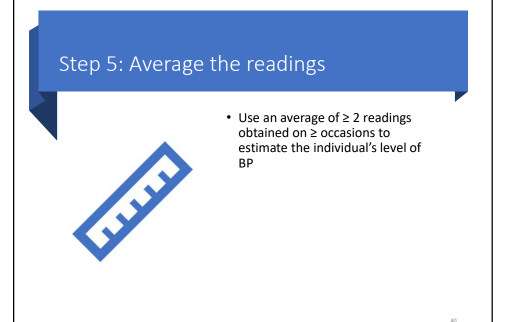
- At first visit, record BP in both arms, Use the arm that gives higher reading for subsequent readings
- Separate repeated measurements by 1-2 min
- For auscultatory determinations, use a palpated estimate of radial pulse obliteration pressure to estimate SBP. Inflate the cuff 20-30 mmHg above this level for an auscultatory determination of the BP level (more info in notes)
- For auscultatory readings, deflate the cuff pressure by 2 mmHg per second, and listen for Korotkoff sounds



Step 4: Properly document accurate BP readings

- Record SBP and DBP
- If using auscultatory technique, record SBP and DBP as the onset of the first Korotkoff sound and disappearance of all Korotkoff sounds, respectively, using the nearest even number
- Note the time and most recent BP medication taken before measurements

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# Step 6: Provide BP readings to patient



 Provide the patient the SBP/DBP readings both verbally and in writing

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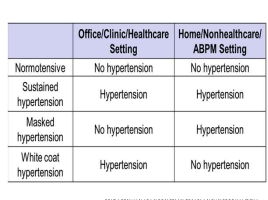
Counseling the Patient: Monitoring Blood Pressure

# Accurate monitoring

- Proper cuff technique
- Proper preparation
  - Relaxed in chair for 5 minutes
  - No exercise, smoking, or caffeine before

# How often?

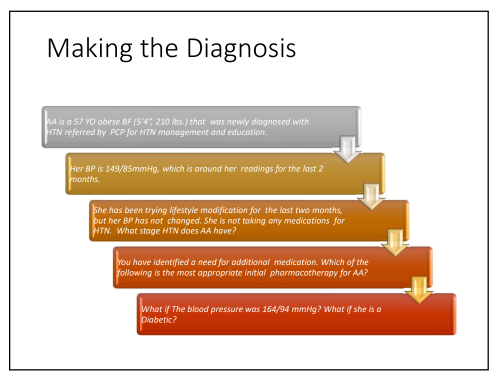
- Daily
  - Average of 2 readings 1 minute apart
  - Before medications in the morning
  - Before supper in the evening

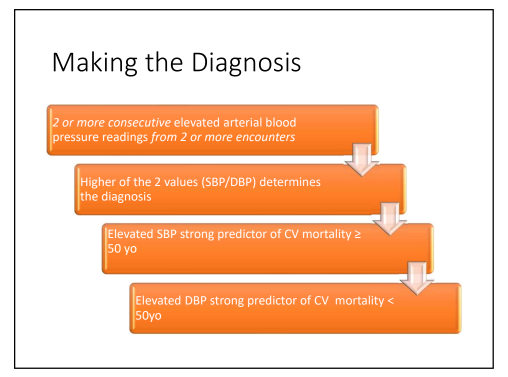


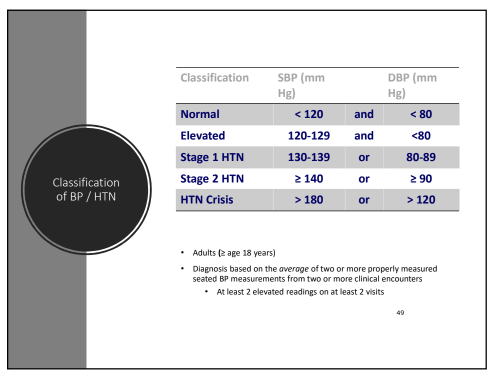
2017 ACC/AHA/AAPA/ABC/ACPM/AGS/ APhA/ASH/ASPC/NMA/PCNA
Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults

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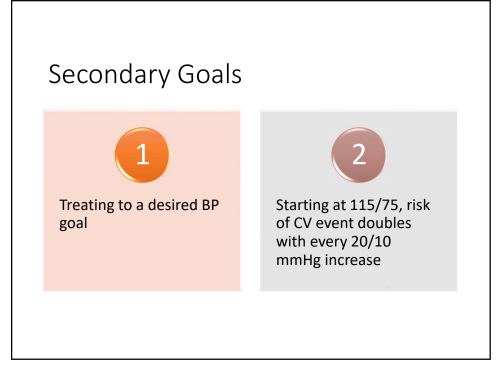












# Risks vs benefits

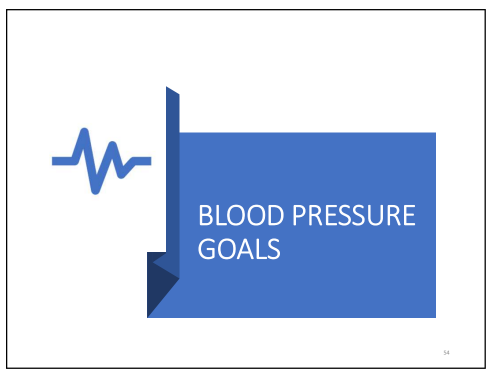
# Target organ damage

- Heart disease angina, MI, HF
- Cerebrovascular disease stroke, TIA
- Kidney disease
- Retinopathy
- Peripheral arterial disease

# Benefits of controlling BP

- Reduction in target organ damage
- Reduction is stroke risk
- · Reduction in MI risk
- Reduction in HF risk

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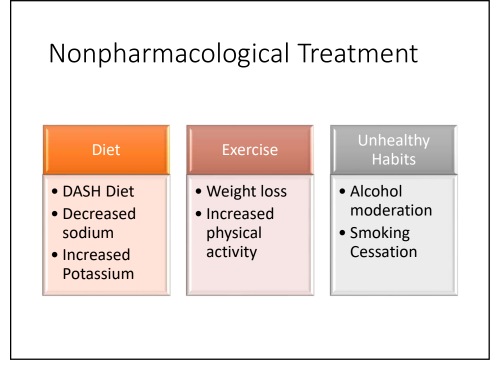


## **Comparison of BP Target Recommendations BP Target BP Categories** < 150/90 mm Hg for patients ≥ JNC 8, 2014 < 140/90 mm Hg for patients < < 120 80-89 Prehypertension 120-139 60, diabetes, Stage 1 hypertension 140-159 90-99 and chronic kidney disease Stage 2 hypertension ≥ 160 ≥ 100 ≤ 130/80 mm Hg SBP DBP ACC/AHA Normal < 120 < 80 2017 Elevated 120-129 < 80 Stage 1 hypertension 130-139 80-89 Stage 2 hypertension ≥ 140 ≥ 90

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# Sprint Trial Population included ≥50 yrs, baseline SBP ≥ 130, elevated CV risk but not diabetes or stroke • Elevated risk = CKD, 10-year Framingham risk score 15%, ≥75 yrs • Target BP < 140 vs. < 120 Mean SBP 121 mmHg vs. 136 after 1 year Primary composite outcomes (myocardial infarction, acute coronary syndrome not resulting in myocardial infarction, stroke, acute decompensated heart failure, or death from cardiovascular causes) better with lower BPs All-cause mortality was also significantly lower in the intensive-treatment group Rates of serious adverse events of hypotension, syncope, electrolyte abnormalities, and acute kidney injury or failure, but not of injurious falls, were higher in the intensive-treatment group than in the standard- treatment group





# Nonpharmacological Treatment

	Nonpharmacologic Intervention		Approximate Impact on SBP	
		Dose	Hypertension	Normotension
Weight loss	Weight/body fat	Ideal body weight is best goal but at least 1 kg reduction in body weight for most adults who are overweight. Expect about 1 mm Hg for every 1 kg reduction in body weight.	-5 mm Hg	-2/3 mm Hg
Healthy diet	DASH dietary pattern	Diet rich in fruits, vegetables, whole grains, and low-fat dairy products with reduced content of saturated and trans I fat	-11 mm Hg	-3 mm Hg
Reduced Intake of dietary sodium	Dietary sodium	<1,500 mg/d is optimal goal but at least 1,000 mg/d reduction in most adults	-5/6 mm Hg	-2/3 mm Hg
Enhanced intake of dietary potassium	Dietary potassium	3,500-5,000 mg/d, preferably by consumption of a diet rich in potassium	-4/5 mm Hg	-2 mm Hg
Physical activity	Aerobic	120-150 min/wk     65%-75% heart rate reserve	-5/8 mm Hg	-2/4 mm Hg
	Dynamic Resistance	90-150 min/wk     50%-80% 1 rep maximum     6 exercises, 3 sets/exercise, 10 repetitions/set	-4 mm Hg	-2 mm Hg
	Isometric Resistance	4 x 2 min (hand grip), 1 min rest between exercises, 30%-40% maximum voluntary contraction, 3 sessions/wk     8-10 wk	-5 mm Hg	-4 mm Hg
Moderation in alcohol intake	Alcohol consumption	In individuals who drink alcohol, reduce alcohol† to: • Men: ≤2 drinks daily • Women: ≤1 drink daily	-4 mm Hg	-3 mm Hg

2017 ACC/AHA/AAPA/ABC/ACPM/AGS/ APhA/ASH/ASPC/NMA/PCNA
Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults

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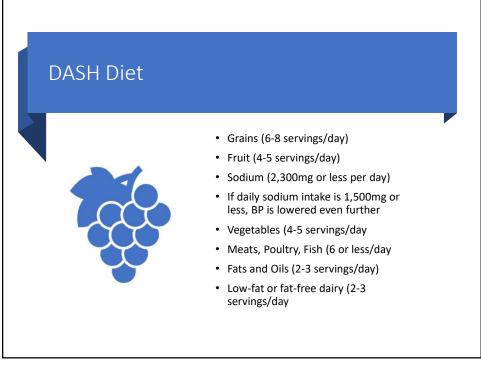
# DASH Diet



- Dietary Approaches to Stop Hypertension
  - Vegetables, fruits, whole grains
  - Fat-free or low-fat dairy products
  - Poultry, beans, nuts, vegetable oils
  - Limit food high in saturated fat fatty meats, full-fat dairy products,

tropical oils

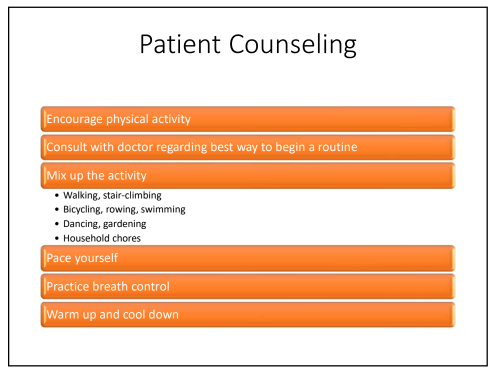
• Limit sugar-sweetened beverages and sweets



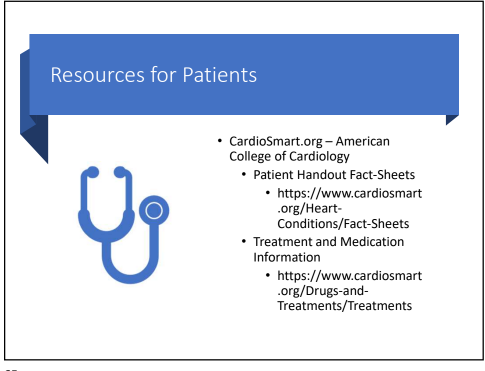
# Exercise

- American Heart Association Recommendations:
  - Aim for 90-150 minutes of aerobic and/or dynamic resistance exercises per week
  - Get the equivalent of 150 minutes per week of moderate- intensity physical activity (such as brisk walking)
  - Perform physical activity in at least 10minute intervals and
  - spread throughout the week
  - Include flexibility and stretching
  - Include muscle-strengthening activity at least twice per week





# Smoking Cessation Counsel on benefits of quitting as often as possible Medications + support = improved quit rates Offer encouragement as it often takes more than one try to quit Benefits of Quitting Smoking: Of smoke-free living: the carbon monoxide levels in your blood pressure and heart rate recover from the cigarette-induced spike. It to three months of smoke-free living: the carbon monoxide levels in your blood elevels in you





# Treatment Recommendations

Initiation of antihypertensive drug therapy, first line agents include thiazide diuretics, CCBs, and ACE inhibitors or ARBs

Stage 1 HTN and goal BP <130/80 – initiation of antihypertensive drug therapy with a single antihypertensive drug is reasonable with dosage titration and sequential addition of other agents to achieve the BP target

Stage 2 HTN and an average BP more than 20/10 mmHG above BP target – initiation of antihypertensive drug therapy with 2 first-line agents of different classes is recommended

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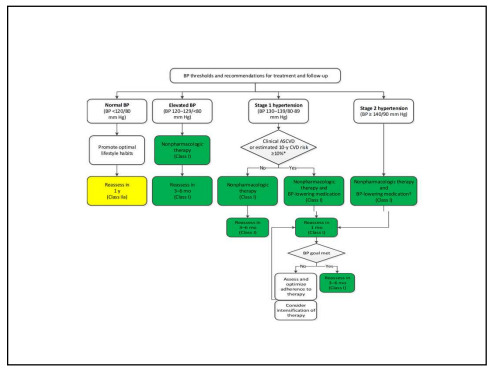
# Follow-up Recommendations

# After initial BP elevation

- Elevated BP or Stage 1 HTN with 10-year ASCVD risk less than 10% nonpharmacological therapy and repeat BP evaluation in 3-6 months
- Stage 1 HTN with 10-year ASCVD risk of 10% or greater nonpharmacological therapy AND antihypertensive treatment and follow-up in 1 month
- Stage 2 HTN evaluated by or referred to PCP within 1 month of initial diagnosis
- nonpharmacological therapy AND 2 antihypertensive drug therapies and follow-up in 1 month

# After initiating antihypertensive drug therapy

 Initiating a new or adjusted drug regiment for HTN should have follow-up evaluation of adherence and response at monthly intervals until goal is reached



# Pharmacotherapy Options

# First Line

- Thiazides
- ACEIs
- ARBs
- Calcium Channel Blockers
- Beta-1 Blockers \*\*

# Second Line

- Potassium Sparing Diuretics (possibly loop diuretics in CKD and HF)
- Aldosterone Antagonists
- Direct Renin Inhibitors
- Direct Vasodilators
- Centrally Acting Alpha-2 Antagonists
- Peripheral Adrenergic Inhibitors
- Alpha-1 Agonists

# Thiazide

### MOA

- ↑ excretion of Na, Cl, H2O
- Inhibit Na ion transport across renal tubular epithelium
- Inhibit active Cl reabsorption at distal ascending limb or distal tubule
- Decrease SV and CO
- Reduce TPR

# Contraindications

- Cross-sensitivity with other thiazides or sulfonamides, anuria, renal decompensation, hemodialysis
- Drug interactions
- Lithium, dofetilide, NSAIDs

# Drug interactions

 Lithium, dofetilide, NSAIDs

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# Thiazide

# Cautions

- Lose effectiveness when CrCl < 30 ml/min</li>
- Metolazone can still be used
- Use caution in patients with sulfonamide allergy
- May precipitate gout (especially if not on uric acidlowering therapy), systemic lupus erythematosus, and change in glucose control

# Monitoring parameters

- -SCr/BUN, Electrolytes, uric acid, Glucose, lipids, Blood pressure, dizziness
- Assess weight, Intake & Output (I&O) reports daily to determine fluid loss

# Thiazide Diuretics/Adverse reactions

### Нуро

- Hypokalemia
- Hyponatremia
- Hypomagnesemia
- Hypochloremia

### HYPER-

- Hypercalcemia
- Hyperuricemia
- Hyperglycemia
- Hyperlipidemia

### Photosensitivity

Higher risk of new onset diabetes (vs ACEI, ARBs, CCB, BB)

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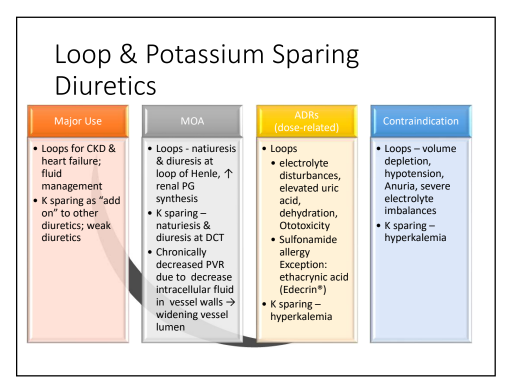
# Thiazide Diuretics

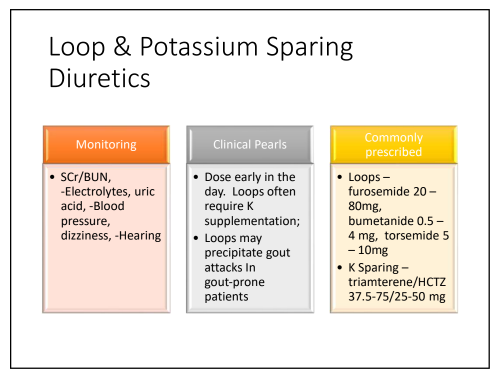
### Clinical Pearls

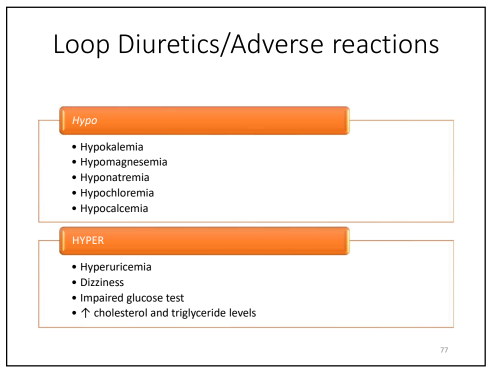
- Chlorthalidone preferred based on prolonged half-life and proven trial reduction of CVD
- Dose in morning and early afternoon if 2nd dose is needed
- Use in caution with patients with history of acute gout unless on uric acid lowering therapy
- Check electrolytes at baseline and as clinically necessary
- Cautions
- Lose effectiveness when CrCl < 30 ml/min
- Metolazone can still be used
- Use caution in patients with sulfonamide allergy
- May precipitate gout (especially if not on uric acid-lowering therapy), systemic lupus erythematosus, and change in glucose control

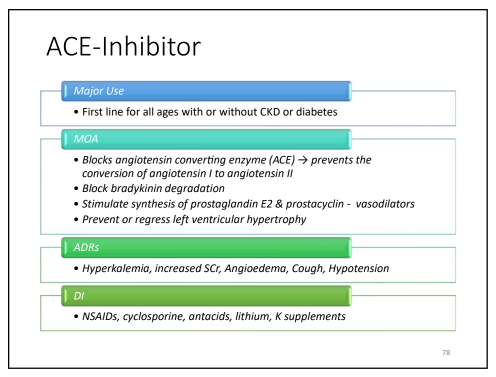
### Commonly prescribed

- Hydrochlorothiazide 12.5 25 mg
- -Chlorthalidone 12.5 50mg
- -Indapamide 1.25 5mg









# **ACE-Inhibitors**

### Contraindications

- Angioedema related to previous treatment with ACE-inhibitor
- Idiopathic or hereditary angioedema
- Pregnancy
- Do NOT use with ARBs or direct renin inhibitor

### Cautions

- Aortic stenosis
- Renal artery stenosis (unstented unilateral OR bilateral) or renal impairment → could cause acute renal failure

### Clinical Pearls

- Shown to work better in Caucasians than AA
- Acute kidney failure adjust dose or d/c if > 35% increase in SCr from baseline
- Dose increase slowly; can decrease or stop quickly
- Do not use in combination with ARB or DRI
- Usually once daily dosing, Twice daily dosing may be needed to maintain 24-hour BP control

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# **ACE-Inhibitors**

Generic	Brand	Usual Daily Dose (mg)
Benazepril	Lotensin®	10 - 40 1 or 2 doses per day
Captopril	Capoten®	25 - 150 2 or 3 doses per day
Enalapril	Vasotec®	5 - 40 1 or 2 doses per day
Fosinopril	Monopril®	10 - 40 Daily
Lisinopril	Prinivil®, Zestril®	10 - 40 Daily

# **ACE-Inhibitors**

Generic	Brand	Usual Daily Dose (mg)
Moexipril	Univasc®	7.5 - 30 1 or 2 doses per day
Perindopril	Aceon®	4 - 16 1 or 2 doses per day
Quinapril	Accupril®	10 - 80 Daily
Ramipril	Altace	2.5 - 20 Daily
Trandolapril	Mavik	<b>1 - 4 Daily</b> 81

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# Angiotensin Receptor Blockers

### Major Use

- First line in all ages with or without CKD or diabetes
- More data to support renoprotective effects

### MO

- Binds to the AT1 angiotensin II receptor, which prevents angiotensin II from binding to the receptor
- Blocks the vasoconstriction and aldosterone secreting effects of angiotensin

### Contraindications

- Angioedema related to previous treatment with ARB
- $\bullet\,$  If angioedema with ACEI, can receive ARB 6 weeks after ACEI is discontinued
- Pregnancy
- Do NOT use with ACEI or direct renin inhibitor

### Cautions

 Aortic/mitral stenosis, unstented unilateral or bilateral renal artery stenosis, renal impairment

# Angiotensin Receptor Blockers

### Adverse reactions

- Angioedema, Dyspepsia, Dyspnea
- Hyperglycemia, Hyperkalemia, Hypertriglyceridemia, Hyperuricemia
- ullet in serum creatinine

### Drug interactions Lithium, NSAIDs

Monitoring parameters Potassium, renal function, blood pressure, Scr.

### Clinical Pearls

- ACE/ARB combination therapy only with severe nephrotic syndrome
- Combination ACE/ARB therapy not recommended for HTN
- Alternative for ACEI-induced cough
- Lower risk of angioedema; not recommended
- If angioedema with ACEi, patient can start on ARB 6 weeks after discontinuation of ACEi
- Dose increase slowly; can decrease or stop quickly

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# Angiotensin Receptor Blockers

Generic	Brand	Usual Daily Dose (mg)
Azilsartan	Edarbi™	40 - 80 Daily
Candesartan	<b>Atacand</b> ®	8 - 32 Daily
Eprosartan	Teveten®	400 - 800 1 or 2 doses per day
Irbesartan	<b>Avapro</b> ®	150 - 300 Daily
Losartan	Cozaar®	25 - 100 1 or 2 doses per day
Olmesartan	Benicar®	20 - 40 Daily
Telmisartan	Micardis®	20 - 80 Daily
Valsartan	Diovan®	80 - 320 1 or 2 doses per day

## Calcium Channel Blockers

Inhibits calcium ion from entering the "slow channels" (select voltage-sensitive areas of vascular smooth muscle and myocardium during depolarization)

Produces a relaxation of coronary vascular smooth muscle and coronary vasodilation

Increases myocardial oxygen delivery in patients with vasospastic angina

Non-dihydropyridines ONLY slow automaticity and conduction of AV node

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# Calcium Channel Blockers

### Contraindications

- Non-dihydropyridines
  - Severe LV dysfunction, cardiogenic shock, sick sinus syndrome, 2nd or 3rd degree AV block
- Dihydropyridines
- Hypersensitivity, advanced aortic stenosis

### Cautions

- Avoid in heart failure with reduced ejection fraction (amlodipine or felodipine may be used if needed)
- Hepatic impairment, hypertrophic cardiomyopathy, renal impairment
- Avoid routine use of non-dihydropyridines with BB due to risk of bradycardia and heart block

### Adverse reactions

- · Non-dihydropyridines
- Edema, HA, 1st degree AV block, hypotension, flushing, rash, gout, constipation (moreso with verapamil), diarrhea, myalgias, dyspnea, gingival hyperplasia (verapamil)
- Dihydropyridines
  - Peripheral edema, HA, somnolence, male sexual dysfunction, abdominal pain, dyspepsia, gingival hyperplasia, muscle cramps

# Calcium Channel Blockers

### Major use:

- first line for all ages with or without diabetes
- NDHP rate control in atrial fibrillation, CHF (diastolic, EF preserved)

### Non-dihydropyridines D/I

- CYP 3A4 inducers and inhibitors
- Amiodarone, azole antifungals, benzodiazepines, carbamazepine, dabigatran, digoxin, dronedarone, seizure medications, macrolide antibiotics, protease inhibitors, ranolazine, risperidone, conivaptan, tolvaptan

### Dihydropyridines D/I

- Azole antifungals, barbiturates, clopidogrel, conivaptan, fosphenytoin, macrolide antibiotics, seizure medications, neuromuscular blockers, protease inhibitors, CYP3A4 and 1A2 inducers and inhibitors
- ullet grapefruit ullet serum concentration of DHP (but you have to drink LOTS of it)

### Monitoring parameters

• HR, BP, peripheral edema & dyspnea (worsening CHF)

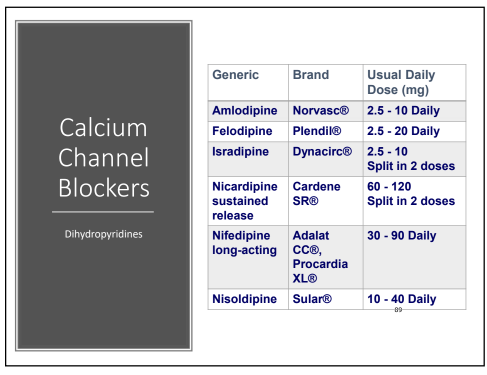
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### Calcium Channel Blockers

• Non-dihydropyridines

Generic	Brand	Usual Daily Dose (mg)
Diltiazem extended release (capsule)	Cardizem CD®, Dilacor XR®, Tiazac	180 - 420 Daily
Diltiazem extended release (tablet)	Cardizem LA	120 - 540 Daily
Verapamil immediate release	Calan®, Isoptin®	80 - 320 Split in 2 doses
Verapamil extended release (tablet)	Calan SR®, Isoptin SR®	120 - 480 1 or 2 doses per day
Verapamil extended release (capsule)	Covera-HS®, Verelan PM®	120 - 480 Daily (at bedtime) 100 - 400 Daily (at bedtime)

Both diltizem and verapamil available as IV



# $\beta$ – blockers Mechanism of Action

Competitively block beta adrenergic receptors

### Effect is dependent on type of receptor

- Beta<sub>1</sub> blockade
  - ↓HR, contractility, cardiac output
- Beta 2 blockade
  - Vasoconstriction
  - Bronchoconstriction

# **β** - blockers

### Contraindications

 Sinus bradycardia, second- or third-degree heart block, cardiogenic shock, overt heart failure, sick sinus syndrome, uncompensated heart failure, pulmonary edema

### Cautions

- Should NOT be withdrawn abruptly
  - Taper over 1-2 weeks
- Bronchospastic disease (non-selective BB should be avoided), DM, heart failure

### Adverse reactions

- Hypotension. Bradycardia, Dizziness, Fatigue, Insomnia, nightmares
- Decreased libido or impotence, Bronchospasm, Depression

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# **β** - blockers

## **Drug interactions**

• Digoxin, theophylline, sulfonylureas, dronedarone

## Monitoring parameters

• HR, BP

### Potentially favorable effects

 Useful for atrial tachyarrhythmias/fibrillation, migraine, thyrotoxicosis (short term), essential tremor, perioperative hypertension

# $\beta$ – blockers/Lipid Solubility

### High

- Largely metabolized by the liver
- Penetrate CNS
- Provide better effects for non-CV conditions
  - Migraine headache prevention, essential tremor, thyrotoxicosis

### Low

• Excreted unchanged by kidneys

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# $\theta$ – blockers (BB)

Non-selective beta blockers (1st generation)

• Bind to beta<sub>1</sub> and beta<sub>2</sub> receptors

ardioselective beta blockers ♥ (2<sup>nd</sup> Generation)

- Bind to beta<sub>1</sub> receptors
- Can bind to beta<sub>2</sub> at higher doses

BB with vasodilatory properties

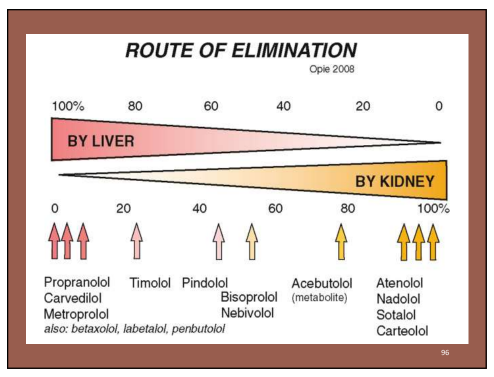
- α-adrenergic blockade
- Direct vasodilation

BB with intrinsic sympathomimetic activity (ISA)

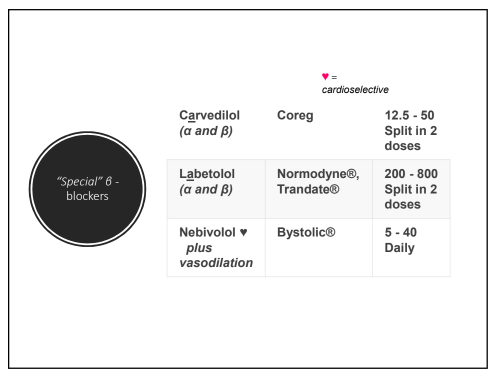
Act as both agonist and antagonist at beta receptors

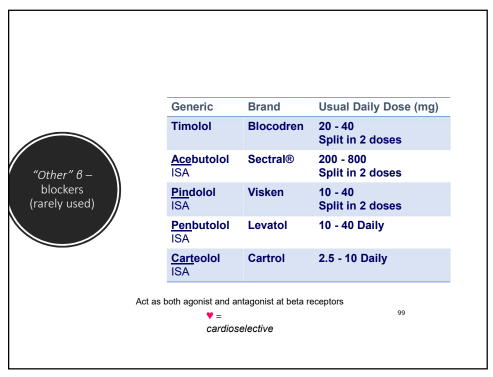
# Bolockers Combined alpha and beta blockers Carvedilol (Coreg®, Coreg CR®) Labetalol (Normodyne®, Trandate®) Vasodilators Nebivolol (Bystolic®) Intrinsic Sympathomimetic (AVOID) Acebutelol (Sectral®) ◆ Penbutolol (Levatol®) Pindolol Carteolol

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	β - blocke	rs
		♥ = cardioselective
Generic	Brand	Usual Daily Dose (mg)
Atenolol •	Tenormin®	25 - 100 Daily
Betaxolol ♥	Kerlone®	5 - 20 Daily
Bisoprolol •	Zebeta	2.5 - 10 Daily
Esmolol •	Brevibloc®	IV only – bolus then continuous infusion
Metoprolol tartrate ♥	Lopressor®	50 - 400 2 or 3 doses per day
Metoprolol succinate ♥	Toprol XL®	50 - 200 Daily
Nadolol	Corgard®	40 - 120 Daily
Propranolol	Inderal	80 - 640 Split in 2 doses
Propranolol (long-acting)	Inderal LA®	60 - 180 Daily



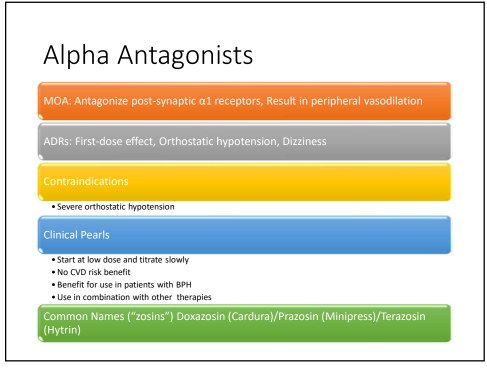


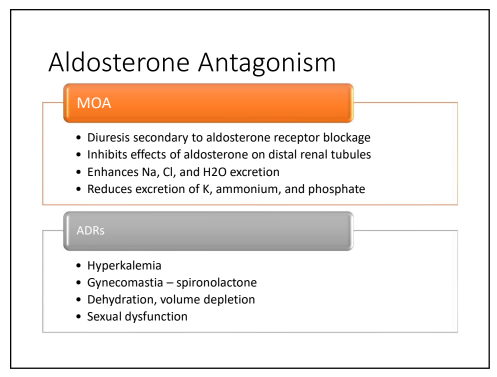


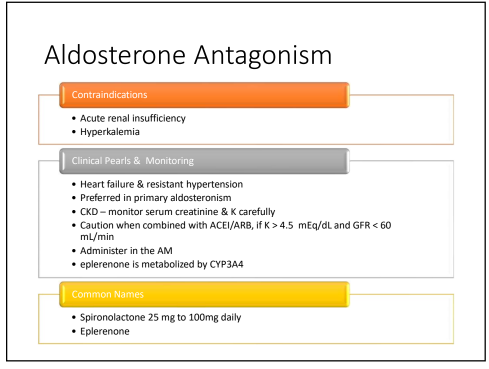
# Alternative Agents Direct arterial vasodilator Hydralazine – is commonly used Alpha blockers Doxazosin (Cardura®), Prazosin (Minipress®), Terazosin (Hytrin®) Direct renin inhibitor Aliskiren (Tekturna®) Centrally acting antihypertensives Clonidine Methyldopa (drug of choice in pregnancy!)

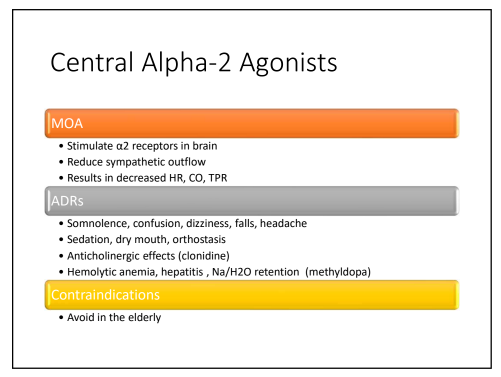
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# Renin Inhibitor MOI: Directly inhibits renin, Inhibits conversion of angiotensinogen to Angiotensin I Results in decreased production of Angiotensin II ADRs: Angioedema rarely, Hyperkalemia Contraindications: Pregnancy Category D Clinical Pearls Alternative or combination therapy Once daily dosing only Taken with high fat meals will reduce absorption No CV risk benefits Avoid combination of ACEI/ARB + K Common Names: Aliskiren, Tekturna









# Central Alpha-2 Agonists

### Clinical Pearls & Monitoring

- Ambulation, alertness
- Concurrent diuretic
- Hepatic function, WBC (methyldopa)
- Avoid abrupt discontinuation
- Must be tapered
- Methyldopa can be used in pregnancy
- Generally last line therapy due to CNS effects

### Common Medications

- Methyldopa 750mg to 3000mg/day BID to TID
- Clonidine 0.1mg to 0.3mg TID
- Guanfacine (Tenex)

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# Peripheral Vasodilators MOA • Arterial smooth muscle vasodilation, NO formation (hydralazine) and K+ channel mediated (hydralazine and minoxidil) • Directly relax smooth muscle in arterioles • Results in peripheral vasodilation ADRS • Reflex tachycardia, Headache, worsening angina • Sodium and water retention, edema • Lupus (hydralazine) immune disorder • Hirsutism (minoxidil) Contraindications • SLE, CAD

# Peripheral Vasodilators

### Clinical Pearls & Monitoring

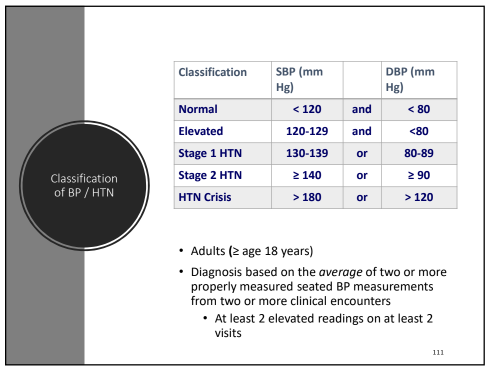
- Muscle weakness (hydralazine)
- Admin w/diuretic and  $\beta$  receptor antagonist, rarely used alone.
- Minoxidil requires a loop diuretic and can cause pericardial
- effusion
- Third-line or later

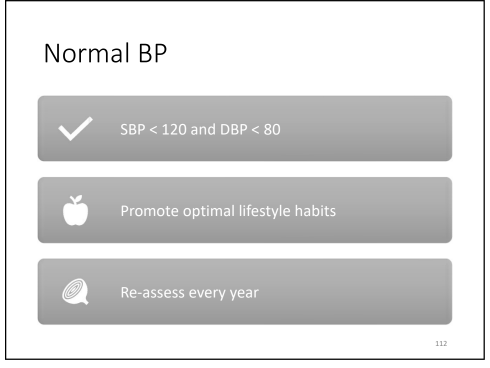
### Common Medications

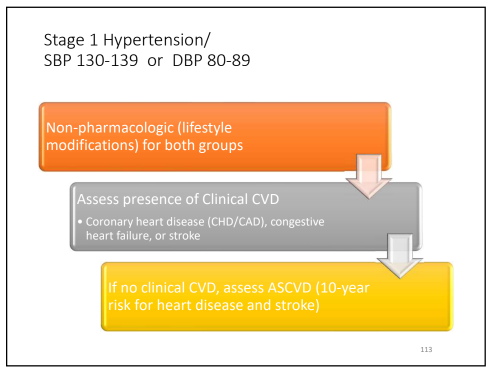
- Minoxidil 5mg to 40mg/day in divided doses
- Hydralazine 40mg to 300mg/day in divided doses

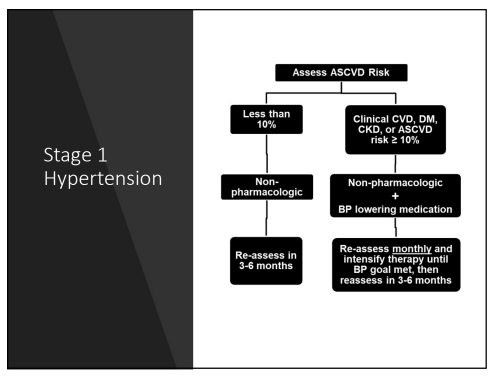
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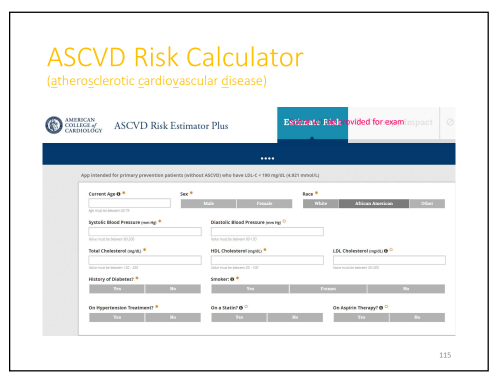


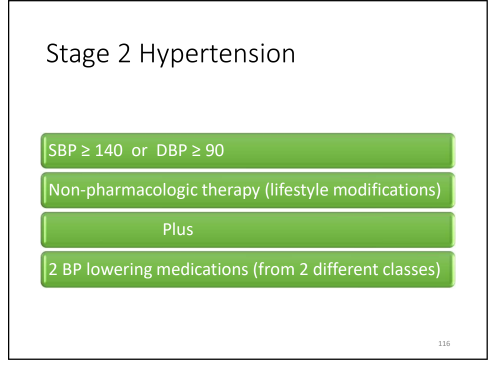


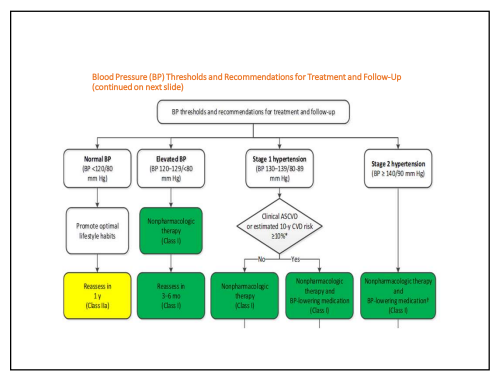


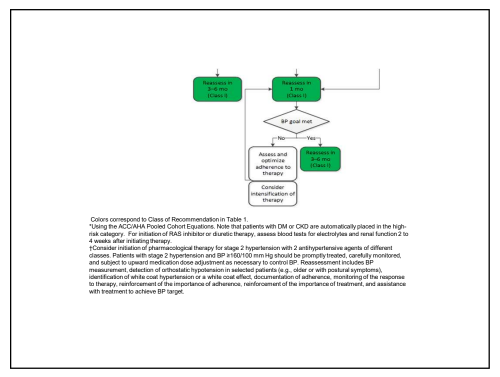


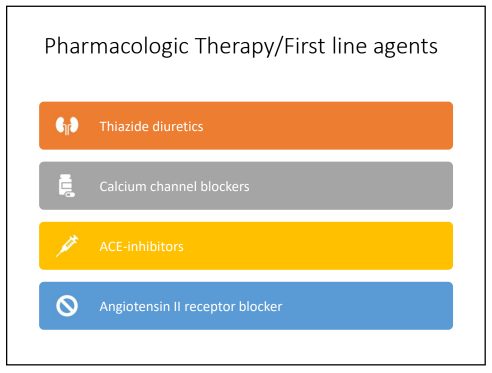


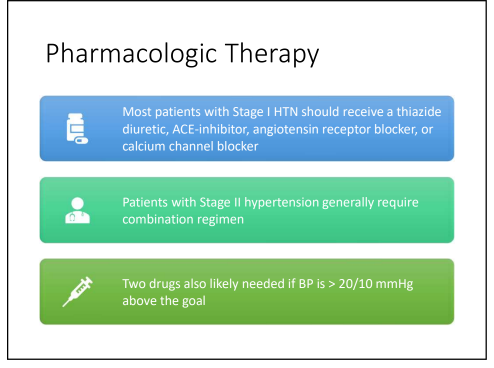


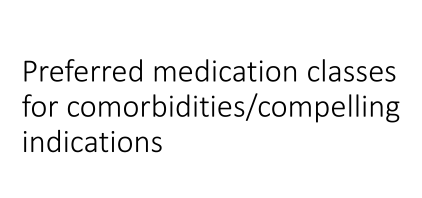


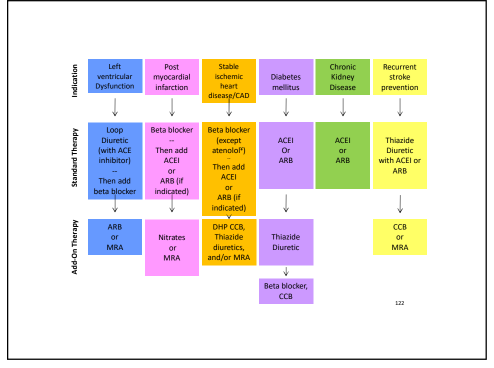












# Heart Failure

### **Diuretics**

- Thiazides better for BP lowering
- Loops better for volume control for LVD and may be necessary if volume overload is a problem

### **ACEI/ARBs**

### **B-Blockers**

- Improved outcomes with 3 specific agents:
- Carvedilol, metoprolol succinate, bisoprolol

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# Heart Failure

Mineralocortocoid receptor antagonists (MRA)

- Spironolactone or eplerenone
- Class II-IV HF with LVD (Class II, EF<35%; Class III-IV EF<40%)

Drugs to avoid in HF pts with HTN:

- Non-dihydropyridines
- Verapamil, diltiazem
- Clonidine
- Minoxidi

Only use alpha-blockers if other drugs are inadequate to achieve BP control

# Post Myocardial Infarction

### β-Blockers

- Start with a short acting B1 selective without intrinsic sympathomimetic activity
- Given with nitrates in acute MI

### Using non-dihydropyridine CCB

- If BB is CI and there is no LVD
- If pt has supraventricular tachycardia
- Do NOT use if bradyarrhythmia or impaired LV function

### CCB – dihydropyridine

Long acting

Note CCB can ↑ mortality if LVD &/or pulmonary edema

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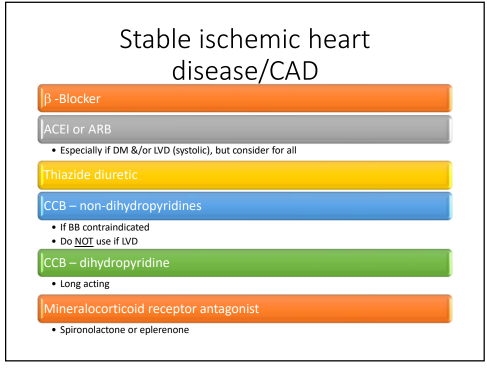
# Post Myocardial Infarction

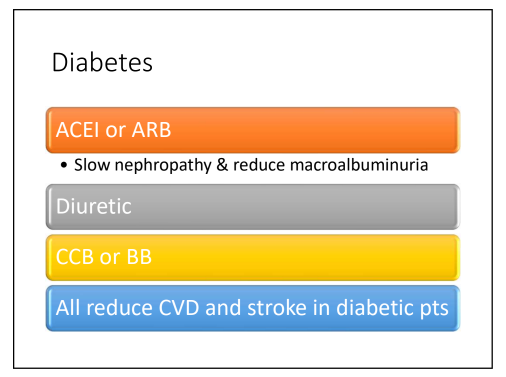
### **ACEI**

- Use in pts with anterior MI (when stable) and persistent HTN, LVD, HF, or DM
- Benefit if infarct is large (STEMI) &/or history of previous infarction or HF
- ARB can also be used, but lower level of evidence

### Mineralocortocoid receptor antagonists (MRA)

- Spironolactone or eplerenone
- Use in STEMI with LVD & HF





# Chronic Kidney Disease (CKD)

### **ACEI or ARB**

- CKD 3 or higher
- Preferred if albuminuria present in stage 1 & 2 CKD
- ≥ 300 mg/day or ≥300 mg/g creatinine
- Delay progression of renal disease
- Rise in serum creatinine (SCr) up to 35% above baseline is acceptable
- Do not hold therapy unless hyperkalemia develops

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# Chronic Kidney Disease (CKD)

In absence of albuminuria, CCB or thiazide diuretics can be used

Loop diuretics are usually needed with advanced renal disease to control volume status (in combination with other medications)

After kidney transplant, it's reasonable to use CCB

# **Recurrent Stroke Prevention**

Thiazide diuretic, ACE or ARB

### Thiazide Diuretic + ACEI (or ARB)

• Combination of diuretic and ACEI reduces rates of recurrent stroke

After first line, BP reduction appears to be more important than agent choice

• May add CCB or mineralocortocoid receptor antagonists (MRA)

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# For patients without compelling indications

## Non-black patients with HTN/Initial therapy

- Thiazide-type diuretics
  - Thiazides, chlorthalidone, indapamide
- Calcium channel blockers (CCB)
- Angiotensin converting enzyme inhibitors (ACEI)
- Angiotensin receptor blockers (ARB)

### Black patients with HTN

- Thiazide-type diuretics
- Thiazides, chlorthalidone, indapamide
- CCB

# Chronic Kidney Disease and HTN

Regardless of race or diabetic status, ACEI or ARB should be used to improve kidney outcomes

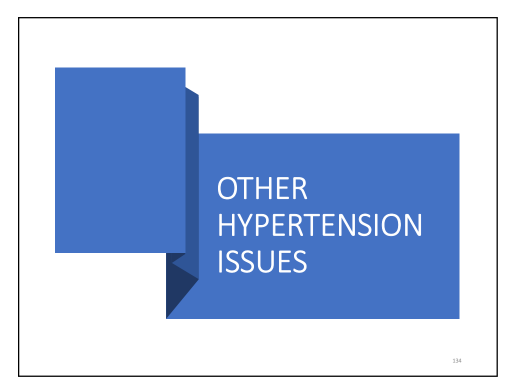
Initial therapy ACEI or ARB

May use ACEI/ARB as add on therapy Do not use ACEI and ARB together

Clinical pearls

- If CKD and proteinuria initial therapy should include ACEI or ARB
- Higher likelihood of progression to end stage renal disease (ESRD)
- If ACEI/ARB not used as initial therapy, it can be added as second-line drug if necessary, to achieve goal BP
- Most patients with CKD and HTN require more than one drug to reach goal BP
- ACEI/ARB with thiazide-type diuretic or CCB

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# Orthostatic Hypotension

- - Diabetes
  - Dehydration
  - ↓baroreceptor activity (age)
  - Autonomic insufficiency (CKD)
  - Venodilators (α-blockers, mixed α/β-blockers, nitrates, phosphodiesterase inhibitors)

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# Resistant Hypertension

### Definition

 Failure to achieve BP goal despite 3 or more BP medications on optimum doses

### Causes

- Drugs inadequate doses, inappropriate choices, BPelevating agents
- Fluid overload
- Nonadherence
- Obesity, alcohol, sleep apnea, excess dietary sodium
- Poor blood pressure measurement technique
- White coat/pseudohypertension

# **Resistant Hypertension**

### reatment

- Remove/treat secondary causes see earlier slides
- Maximize diuretic therapy
- Add a mineralocorticoid receptor antagonist
- Add other agents with different MOAs
- Use loop diuretics in patients with CKD and/or patients receiving potent vasodilators (minoxidil)
- Identify and correct barriers to adherence
- Weight loss, limit alcohol, sodium restriction
- Potassium supplementation
- Home/ambulatory monitoring, Osler's sign

### Refer to specialist

- Refer to specialist for known or suspected secondary cause(s) of HTN
- Refer to HTN specialist if BP remains uncontrolled after 6 mon of treatment

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# Pregnancy

### Preferred meds

- Methyldopa
- Nifedipine
- Labetalol

Hydralazine may also be used

### ACEIs and ARBS should NOT be used

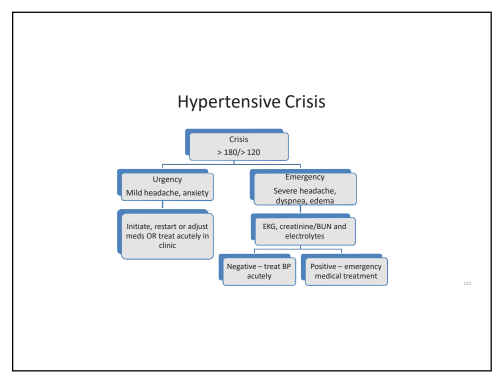
- Potential for fetal defects
- Should be avoided in women likely to become pregnant also

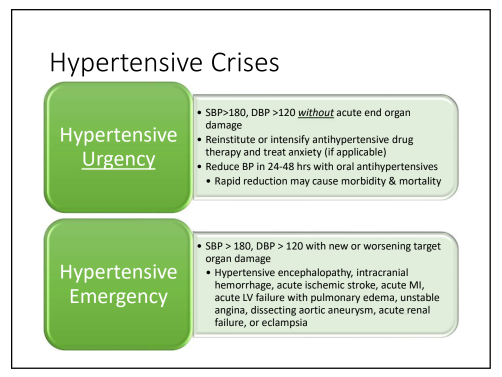
Drug	Disadvantages – how to monitor
Thiazide/Loop diuretics	Urinary Frequency – take earlier in the day Electrolyte abnormalities – K, Na; monitor more frequently Worsening of gout – monitor uric acid
ACEI/ARB	Acute renal failure – avoid if Scr rises > 35% Hyperkalemia – low potassium diet Profound BP lowering w/volume depletion – dose low, go slow
CCBs	Peripheral edema – elevate legs, avoid excess Na Reflex tachycardia – consider combined use with BBs Profound BP lowering – dose low, go slow Bradycardia (nonDHPs) – avoid use with BBs Constipation – laxatives, fiber, fluids Isolated systolic hypertension - preferred
BBs (beta1 preferred)	Bradycardia – avoid use with nonDHP CCBs
Clonidine	Anticholinergic effects - depression, urinary retention, sedation, falls, confusion, vivid dreams, third- or fourth-line agent
α - Antagonists	Orthostasis, dizziness – take at bedtime, dose slowly, use generally for benign prostatic hypertrophy symptoms; little CV benefit

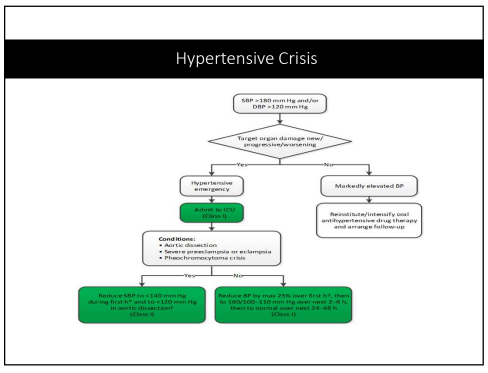


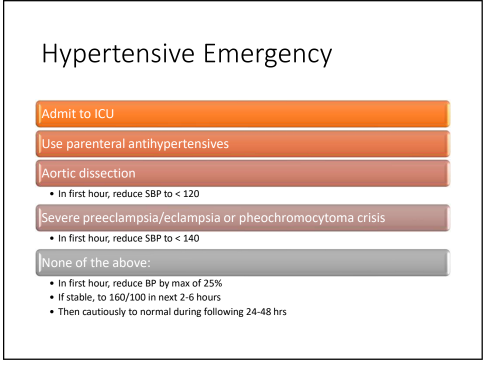
# Hypertensive Emergencies

- Causes
  - Vascular sclerosis
  - Renal parenchymal disease
  - Cocaine, amphetamine or stimulant abuse
  - Rapid clonidine withdrawal
  - Endocrine disease pheochromocytoma, hyperaldosteronism, Cushings
  - CNS trauma, Guillain-Barré syndrome
  - Coarctation of aorta
  - Pre-eclampsia
  - Postoperative









# Hypertensive Emergency

### Acute Intracerebral hemorrhage

- Continuous IV antihypertensive infusion with close BP monitoring
- Immediate reduction of SBP to < 140 can be harmful

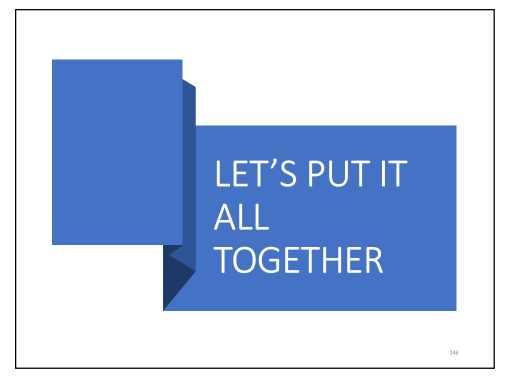
### Acute ischemic stroke

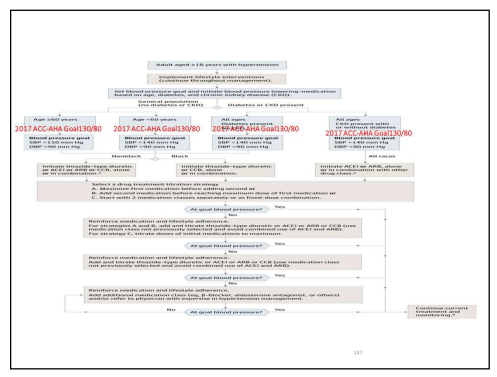
- IV thrombolysis candidates
- Lower SBP to < 185 and DBP to < 110 before initiating thrombolysis
- Maintain BP < 180/105 for first 24 hrs after thrombolysis
- Non-thrombolysis candidates
  - If BP > 220/110, lower BP 15% during first 24 hours
  - IF BP ≤ 220/110, no benefit of treating HTN in first 48-72 hrs

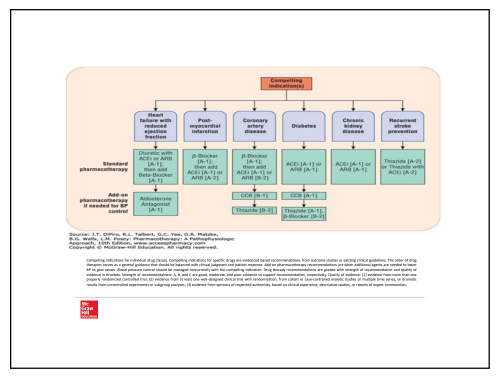
Start/resume antihypertensives  $\geq$  72 hrs from symptom onset with stable neurological status if SBP is  $\geq$  140 or DBP is  $\geq$  90

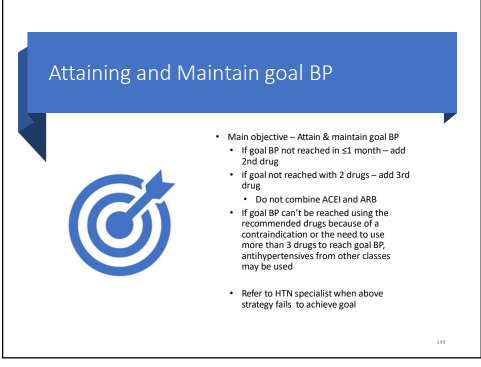
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# Monitoring Parameters

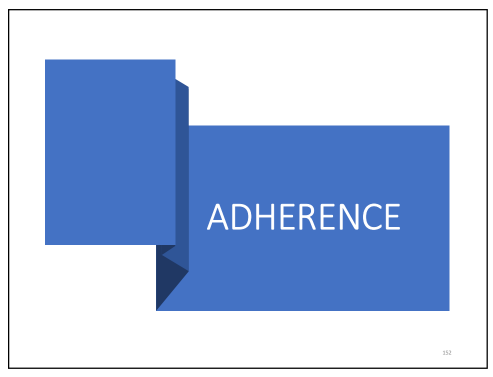
### BP monitoring

- 2 to 4 weeks after changing or initiating therapy
- 6 to 12 months when controlled or stable
- Home or more frequent monitoring if uncontrolled or suspect organ damage

### Organ disease progression

- Signs: EKG, SCr, proteinuria, retinal exam
- Symptoms: ischemic chest pain (or pressure), palpitations, dizziness, dyspnea, orthopnea, headache, sudden change in vision, one-sided weakness, slurred speech, and loss of balance

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# Adherence

### Identify

• Identify any drug therapy problems – indication, safety, efficacy, adherence!

### Simplify

- Simplify medication regimen
  - Drug combinations, sustained release formulations

### **Empower**

- Empower informed patients
- Explicit instruction, good counseling techniques, teach back, literacy issues
- Pill hoxes
- Phone or email reminders of refills, phone/office appointments

### Maintain

• Maintain follow-up with patient, home blood pressure monitoring

### Team

• Team-base, collaborative models of care

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# Summary

Hypertension increases risk for cardiovascular morbidity and mortality

Lifestyle modifications should be encouraged for all patients

Stage I and II HTN typically require pharmacologic therapy

• Stage II often requires more than one agent

Specific treatment recommendations are defined for compelling indications, ischemic heart disease, African Americans, and elderly pts