ANMC GUIDELINES FOR MANAGEMENT OF HYPERTENSIVE DISORDERS IN PREGNANCY

INTRODUCTION
Hypertensive disorders of pregnancy include chronic hypertension, gestational hypertension, preeclampsia without severe features, preeclampsia with severe features, eclampsia, and HELLP.

It is important to understand that hypertensive disorders of pregnancy are progressive, and any diagnosis requires close follow up.

Patients with chronic hypertension are 4-5x more likely to develop preeclampsia (superimposed preeclampsia) than patients without hypertension. Thus, of patients with chronic hypertension 15-40% will develop superimposed preeclampsia. Patients who develop superimposed preeclampsia have higher rates of adverse maternal and fetal outcomes including a 3x higher risk of placental abruption, increased risk of fetal growth restriction, and increased risk of perinatal death.

Patients with chronic hypertension also have an increased risk of gestational diabetes, cesarean delivery, and postpartum hemorrhage.

When managed expectantly, 10-20% of patients with gestational hypertension will develop severe hypertension – this risk is greatest with gestational hypertension is diagnosed before 30wga. Of patients with gestational hypertension, 0.5% will ultimately progress to eclampsia, and 1-2% will develop HELLP syndrome.

In the rare circumstance that intravenous bolus labetalol, hydralazine, or immediate release oral nifedipine fails to relieve acute-onset, severe hypertension and is given in successive appropriate doses, consultation with an anesthesiologist, maternal–fetal medicine subspecialist, or critical care subspecialist to discuss second-line intervention may be recommended. Second-line alternatives to consider may include nicardipine or esmolol by infusion pump.

Eclampsia is a rare outcome that can occur in any patient with preeclampsia even without severe features or while on magnesium therapy. There is not high-quality data to inform the length of magnesium therapy postpartum, and seizures due to eclampsia are less common postpartum. There is some data showing that discontinuation earlier than 24 hours may not increase the risk of seizure. The benefits of earlier discontinuation in appropriate candidates may be improve patient satisfaction and safety as well as decrease time to ambulation, breastfeeding, and hospital discharge.

I. DEFINITIONS

Elevated blood pressure
- Systolic BP (SBP) ≥ 140mmHg or diastolic BP (DBP) ≥ 90mmHg
Severe range blood pressure
- SBP ≥ 160 mmHg and/or DBP ≥ 110 mmHg

Proteinuria: Abnormal amount of protein in urine
- The diagnosis of proteinuria is made with a Total Protein / Creat > 0.3, or a 24 hour urine specimen > 300 mg, or 2+ dipstick
  - If Total P/CR < 0.15 negative for significant proteinuria
  - If > 0.3 positive for significant proteinuria
  - If 0.16-0.29 get 24-hour urine for total protein
- Proteinuria can be misrepresented on contaminated urine specimens. Consider a reflex CCUA on all P:C > 0.15. If necessary, repeat the test on a clean specimen, e.g., 0-5 squamous cells, or on a catheterized specimen.

Thrombocytopenia
- Platelet count < 100,000/microliter

Renal insufficiency
- Creatinine (Cr) > 1.1mg/dL or a doubling of serum Cr concentration from baseline in the absence of other kidney disease

Impaired liver function
- Elevated liver transaminases (AST and ALT) to twice normal concentration
- Persistent severe right upper quadrant (RUQ) pain and/or severe epigastric pain not accounted by alternative diagnoses

Pulmonary edema
- Signs and symptoms of pulmonary edema include shortness of breath, crackles on lung exam, chest x-ray with diffuse infiltrates, and hypoxemia

Cerebral or visual symptoms
- Headache not improved with Tylenol
- Scotoma – seeing spots in vision

II. DIAGNOSIS

Preeclampsia
Newly elevated blood pressures after 20 weeks gestational age or within the first 6wks of the post-partum period
- Two readings of SBP ≥ 140 mmHg and/or DBP ≥ 90 at least 4 hours apart
- SBP ≥ 160 and/or DBP ≥ 110 twice – can be within a short interval (15 minutes)

AND
- Proteinuria

OR
In the absence of proteinuria, new onset of hypertension with the new onset any the following:
- Thrombocytopenia
- Renal insufficiency
- Impaired liver function
- Pulmonary edema
- New onset headache unresponsive to acetaminophen and not accounted for by alternative diagnoses
- Visual disturbances

**Preeclampsia with severe features (any of these)**

Diagnosis of preeclampsia (as above) with ANY of the following:
- Systolic blood pressure of 160 mm Hg or higher, or diastolic blood pressure of 110 mm Hg or higher on two occasions at least 4 hours apart while the patient is on bed rest (unless antihypertensive therapy is initiated before this time)
- Thrombocytopenia
- Impaired liver function
- Renal insufficiency
- Pulmonary edema
- New-onset headache unresponsive to acetaminophen and not accounted for by alternative diagnoses
- Visual disturbances

**HELLP syndrome**

[Hemolysis (abnormal peripheral smear, elevated bilirubin, elevated LDH), Elevated Liver enzymes, Low Platelets]
- LDH > 600 IU/L
- AST > 2x normal
- ALT > 2x normal
- Platelets < 100,000 x 10⁹/L
(Not all lab abnormalities are required for diagnosis)

- Common symptoms: RUQ pain, malaise, N/V

**Eclampsia**

New onset generalized seizures in pregnancy. Eclampsia should be suspected in all cases of generalized tonic-clonic seizures in a pregnant patient without prior history of seizure disorder and no other clear cause of seizure.

**Chronic Hypertension (CHTN)**

Hypertension that pre-dates pregnancy OR onset of hypertension before 20th week of gestation OR use of antihypertensive medications before pregnancy OR failure to normalize blood pressure after 12wks postpartum

**CHTN with Superimposed Preeclampsia**

Preeclampsia in a woman with a history of hypertension before pregnancy or before 20 wks of gestation

**CHTN with Superimposed Preeclampsia w/severe features.**
Preeclampsia in a woman with a history of hypertension before pregnancy or before 20 wks of gestation with the presence of severe symptoms

**Gestational hypertension**
Blood pressure elevation after 20 weeks GA without proteinuria
- Two readings of SBP ≥ 140 mmHg and/or DBP ≥ 90 at least 4 hours apart
- SBP ≥ 160 and/or DBP ≥ 110 twice – can be within a short interval (15 minutes)

If blood pressure elevation persists ≥12 weeks postpartum the diagnosis is changed to chronic hypertension.

**SCREENING AND INITIAL EVALUATION**
ALL women should have Blood pressure (BP) evaluation at initial and all prenatal visits

There is no role for universal urine dipstick testing to screen for preeclampsia in routine prenatal care in the asymptomatic non-hypertensive patient. On the other hand, do obtain a clean catch urine sample if the patient has:
- BP > 140/90
- Symptoms of preeclampsia
- Signs / symptoms of a urinary tract infection
- Multiple gestation
- Chronic hypertension or currently on anti-hypertension medication.

**Pregnancy and health history**
- Determine if patient has a history of chronic hypertension, gestational hypertension, preeclampsia, or eclampsia
  - This may require a review of past medical records with attention to vital signs at clinic and hospital visits.
  - Note pre-term or early term deliveries and perinatal outcomes.
- Women with chronic hypertension (CHTN), pre-gestational diabetes, chronic kidney disease, a multiple gestation pregnancy; a history of gestational hypertension, preeclampsia, eclampsia, or HELLP should also have:
  - Comprehensive metabolic panel (CMP) – to evaluate creatinine (Cr), potassium (K), and liver function (AST, ALT)
  - Baseline evaluation for proteinuria

When to refer for additional evaluation
- Abnormal kidney function – consider renal ultrasound, referral to OB/GYN
  - Cr > 1.1
  - Proteinuria
  - K < 3.0
- History of chronic hypertension >4 years
  - Most women have essential hypertension, but 10% of women with CHTN will have underlying renal or endocrine disorders.

**EARLY INTERVENTION TO PREVENT PREECLAMPSIA**

Low dose aspirin
Evidence supports daily low dose aspirin (81mg) when started between 12-28wk of pregnancy (ideally before 16wk) reduces the occurrence of preeclampsia, preterm birth, and IUGR in women at increased risk.

The harms of low-dose aspirin in pregnancy are small

Aspirin is RECOMMENDED for the following women:
- History of pre-eclampsia
- Multifetal gestation
- Chronic hypertension
- Preexisting Type 1 or Type 2 DM
- Renal disease
- Autoimmune disease (such as systemic lupus erythematosus or antiphospholipid antibody syndrome)

Daily aspirin should be continued until delivery

Calcium supplementation (1.5-2g) in women with low baseline calcium intake (<600mg/day)
- Most multivitamins and prenatal vitamins only have 200-300mg of calcium
- Recommend 1.5-2g supplemental calcium in divided doses for Alaskan Native patients with low dietary calcium intake especially if patient is lactose intolerance or have a known low dietary calcium intake.

Vitamin D supplementation – Recommended for all pregnant Alaskan women to be supplemented with 1,000 IU/day in addition to a daily prenatal vitamin containing 400 IU/day (not to exceed 4,000 IU/day)

The following interventions are NOT recommended for prevention of pre-eclampsia
- Supplementation with vitamin C or vitamin E
- Restriction of dietary salt intake
- Bedrest or restriction of physical activity

EVALUATION

Screening for hypertensive disorders in pregnancy is recommended for all pregnant women at the first prenatal visit and throughout the remainder of pregnancy.

Initiate an evaluation for hypertensive disorders of pregnancy when:
- Elevated BP with systolic BP (SBP) 140 to < 160 mmHg or diastolic BP (DBP) 90 to < 110 on TWO occasions at least 4hr apart
  - If one abnormal reading, consider having patient present to triage/hospital or stay in clinic for repeat BP
  - Elevated BP should be confirmed before initiating laboratory work up.
- New onset severe range blood pressure SBP ≥160 or DBP ≥ 110 should have work up initiated immediately and patient sent to triage for evaluation
- New onset symptoms
  - Headache not improved with acetaminophen
  - Visual changes (seeing spots in vision)
  - Persistent right upper quadrant or epigastric pain
- New onset proteinuria or renal impairment
- Fetal growth restriction

Evaluation should include:
- Clean catch urinalysis (CCUA) and urine total protein: creatinine ratio
  - If >5 squamous cells or evidence of UTI, then invalid for protein assessment
- Ultrasound for fetal growth
- NST
- CMP (assessment of Cr, AST, ALT)
- CBC

**MANAGEMENT**

**CHRONIC HYPERTENSION**
- Initial prenatal evaluation as above
- If severe hypertension and/or cardiac, renal, or connective tissue disorders – early OB consultation or referral
- For pregnant women w/CHTN and poorly controlled BP, the use of home BP monitoring is suggested
- For women with well controlled CHTN who are accustomed to exercising, it is recommended that moderate exercise be continued during pregnancy

**Antihypertensive Therapy**
- ACE inhibitors are contraindicated and should be stopped as soon as pregnancy is diagnosed
- All BP medications can be continued (except ACE inhibitors above)
- If initial blood pressure is well controlled < 140/90, no changes are necessary
  - first line meds in pregnancy: labetalol, nifedipine, and hydralazine
  - If persistent chronic hypertension when systolic pressure is 140 mm Hg or more, diastolic pressure is 90 mm Hg or more, or both then adjust medication to stay within threshold. (See Appendix III)
  - In the setting of comorbidities or underlying impaired renal function, treating at lower blood pressure thresholds may be appropriate.

**Monitoring**
- Prenatal care visits Q 4wks until 32wks then Q2wks if BP remains well controlled
- Initial ultrasonography (US) at 8-10 wks
- Anatomy US at 20-22wk
- Growth US repeated at 28-32wga and then Q3-4 weeks
- Antenatal testing
  - Well controlled: No meds
    - On meds: NST + AFI Q week starting at 36wk
  - Not well controlled
    - On meds: NST 2x /wk + AFI Q week starting at 32wk
  - If FGR develops:
    - At Dx: NST 2x /wk + AFI Q week + Doppler 1x /wk
    - If oligohydramnios
      - At Dx: NST / wk plus repeat AFI / MVP 24 hrs initially

If oligohydramnios
Labs: Baseline and if BP is difficult to control or exam changes
- CBC, CMP (Cr, AST/ALT), and urine P:C ratio or urine dip for proteinuria
- Obtain baseline EKG:
  - If poorly controlled hypertension for more than 4 years or those suspected of having long-standing hypertension based on age (older than 30 years) are more likely to have cardiac hypertrophic changes, cardiomegaly, and ischemic heart disease

Delivery
- Well controlled:
  - No meds 39-39 6/7 wks
  - On meds 39-39 6/7 wks
- Not well controlled
  - On meds > 37wks

GESTATIONAL HYPERTENSION
- Diagnosis ≥ 37wga -> DELIVER
- Diagnosis < 37wga
  - Daily kick counts
  - Prenatal visits: Every 4 weeks until 32 weeks, then every 2 weeks until 36 weeks, then weekly
  - BP check twice weekly
    - One check can be at home if patient has BP cuff and is reliable
  - Antenatal testing w/ NST + AFI weekly at diagnosis after 32 wks
  - If FGR develops:
    - At Dx NST 2x/wk + AFI Q week + Doppler 1x/wk
    - If oligohydramnios
      - At Dx NST/wk plus repeat AFI/MVP 24 hrs initially
  - Labs w/each prenatal visit
    - CBC, CMP (Cr, AST/ALT), and urine P:C ratio or urine dip for proteinuria
    - Growth US every 3-4 weeks
  *If develops laboratory values or signs/symptoms consistent with pre-eclampsia, adjust management to new diagnosis.

Delivery
- 37-38 wks

PREECLAMPSIA WITHOUT SEVERE FEATURES
- Transfer if higher level of care as needed for further evaluation and management
- DELIVER if:
  - Diagnosis ≥ 37wga
  - Diagnosis ≥ 34wga < 37 wga and if any of the following
    - Reversed end-diastolic flow on umbilical artery Doppler studies
- Oligohydramnios (AFI <5cm, MVP < 2cm)
- Persistent BPP ≤ 6/10
- Fetal death
- Lethal anomaly or extreme prematurity

Consider course of late-preterm corticosteroids, if not previously given during this pregnancy, with initiation of induction of labor – not to delay delivery process.

- Expectant Management if diagnosis < 37wga without any findings noted above
  - Transfer to ANMC at diagnosis or 37wks if undelivered
  - Inpatient vs outpatient management
    - Outpatient management of preeclampsia without severe features can be considered if medical adherence with home care and follow-up guidelines is assured and followed.

- Inpatient management:
  - Regular diet
  - Bed rest NOT recommended
  - Vital signs Q6hrs while awake (BP, HR, urine output)
  - Daily weight, best in AM
  - Daily evaluation for CNS and GI symptoms, fetal movement, vaginal bleeding, contractions
  - At least daily NST
  - Fluid restriction NOT recommended

- Outpatient management:
  - Daily patient self-assessment of fetal movement and maternal symptoms
  - Weekly provider visits
  - BP check twice weekly – these can be done at time of antenatal testing and/or provider visit
  - Antenatal testing w/ NST twice weekly + AFI Q week at diagnosis
  - If FGR develops:
    At Dx NST 2x /wk + AFI Q week + Doppler 1x /wk
    If oligohydramnios
    At Dx NST / wk plus repeat AFI / MVP 24 hrs initially
  - Labs w/each weekly prenatal visit
    - CBC, CMP (Cr, AST/ALT), and urine P:C ratio or urine dip for proteinuria
    - Growth US every 3-4 weeks

- Antihypertensive Medications
  - Persistent hypertension when systolic pressure is 160 mm Hg or more, diastolic pressure is 110 mm Hg or more, or both. (See Appendix III)

* If patient develops signs or symptoms of severe features or evidence of worsening disease, adjust management to management of preeclampsia with severe features.
**PREECLAMPSIA WITH SEVERE FEATURES**

**Initial Steps**
- Prompt OB consultation
- Consider initial infusion of 500 ml of lactated ringers (LR) IV as a bolus, as many patients will have vasoconstriction, within overall daily fluid restriction.
  - Do not bolus fluid if: SOB or SpO2 < 95%, or renal impairment
  - Consider continued IV fluids if evidence of hemoconcentration or oliguria
- Treat severe range BP as soon as possible (30-60 minutes) with antihypertensive medications – see Appendix IV
- Start Magnesium sulfate for seizure prophylaxis – see Appendix V
- Transfer to tertiary care center once stable (stretcher, medical escort)

**Maternal stabilization and DELIVER immediately if:**
- ≥ 34wga (can initiate course of late pre-term corticosteroids with induction)
- < 34wga and any of the following:
  - Maternal
    - Uncontrolled severe hypertension
    - Persistent headache
    - Persistent epigastric pain
    - Persistent CNS Sx
    - Stroke, MI
    - HELLP
    - Worsening renal function
    - Eclampsia
    - Pulmonary edema
    - Placental abruption
  - Fetal
    - Non-reassuring fetal heart status
    - Intrauterine fetal demise
    - Lethal anomaly or extreme prematurity
    - Persistent reversed end diastolic umbilical artery flow

Consider course of corticosteroids, or rescue corticosteroids as per guideline, with initiation of induction of labor – not to delay delivery process.

- During this time, patient should be treated with:
  - MagSO4 for seizure prophylaxis
  - Antihypertensive medication
  - Labor induction

- **EXPECTANT MANAGEMENT** if <34wk and stable (without any of the signs or symptoms listed above), manage as an inpatient until 34 0/7wga with the following:
  - Administer corticosteroids or rescue corticosteroids as per guideline
  - MagSO4 for seizure prophylaxis for initial 48hours
  - Antihypertensive medication
- Vital signs, intake and output, symptoms of severe pre-eclampsia Q6 hrs while awake
- Daily weight, best in AM
- CBC, CMP daily; can do every other day if values are stable
- NST daily
- Growth US Q 2-3wks
- Umbilical artery Doppler studies Qwk if intrauterine growth restriction present
- DELIVER if worsening disease or non-reassuring fetal status
- Fluid restriction NOT recommended if stable

**ECLAMPSIA**

**Initial Steps – maternal stabilization**
- Place patient in ‘position of safety’ until more alert
- Give magnesium STAT IM/IV (See Appendix V)
  - If eclampsia occurs with no MagSO4 ongoing, then give 6 gm bolus and start 2 gm/hr
  - If eclampsia occurs while MagSO4 drip already ongoing, then re-bolus with 2-4 gm and increase drip to 3gm/hr IV
  - If additional seizure with MagSO4 drip already ongoing, then re-bolus with 2-4 gm and place drip at 3gm/hr IV
  - If eclampsia, obtain a magnesium level when feasible, but do not delay bolus(s) or maintenance level changes while awaiting magnesium results
- Treat severe hypertension (See Appendix IV)
- Oxygen by mask at 8 L/minute when seizures resolved
- Continuous fetal monitoring,
  - Anticipate non-reassuring fetal heart rate in the immediate postictal period.
  - Allow for in utero resuscitation (with standard measures)
  - If non-reassuring FHR persists after 30 minutes, stat OB consultation
- Transport to tertiary care center after stabilization (stretcher, medical escort, magnesium sulfate, oxygen, IV Fluids)

**Deliver AFTER maternal stabilization**
- **SEIZURE IS NOT AN INDICATION FOR CESAREAN DELIVERY**
  - Urgent cesarean delivery as a result of responding to the non-reassuring fetal tracing immediately after a seizure is cautioned, as the risk of maternal cerebrovascular hemorrhage is extremely high at this time.
  - Continue MagSO4 during the cesarean delivery
  - Please stabilize with adequate magnesium sulfate first if a cesarean delivery must be done despite the risk, e.g., status epilepticus
  - Remember ‘Primum non nocere’ a.k.a. ‘do no harm’, most eclamptic seizures resolve spontaneously

**SPECIAL CIRCUMSTANCES**

Cesarean delivery while on MagSO4
DO NOT DISCONTINUE MagSO4
- The ½ life of MagSO4 is 5 hours. Discontinuing at the time of cesarean delivery only minimally decreases the concentration and potentially increases the risk of a seizure.

Fluid Management
- Maternal death in hypertensive disease of pregnancy is much more the result of pulmonary edema than renal failure, so judicious fluid administration seems prudent.
- In uncomplicated preeclampsia, total fluids should be restricted to 100 -120 mL/h (3000 cc/24 hr total: IVF, boluses, PO, piggybacks), and the urine output monitored hourly.
- If one feels the patient remains vasoconstricted (urine output < 20 cc/ hr) and may benefit from additional fluid boluses, please consult an OB/GYN.

POSTPARTUM MANAGEMENT
- If on MagSO4, continue for 12-24 hours from delivery
- Consider 12 hr MagSO4 duration if: Diuresis for two consecutive hours, absence of symptoms [headache, visual changes, epigastric pain], and absence of severe hypertension.
- Depending on clinical condition, consider continued Pitocin IV drip
- Avoid methergine (could cause vasospasm and increase BP)
- Monitor intake and output every 4-8 hours until discharge
- Continue or initiate antihypertensive therapy when SBP ≥ 150mmHg or DBP ≥ 100mmHg on two occasions at least 4 hours apart
- If SBP ≥ 160mmHg or DBP ≥ 110mmHg on two checks 15 minutes apart, treat with IV antihypertensive medications within 30 minutes and initiate MagSO4
- If using NSAIDs in those patients without renal dysfunction, then one may need to adjust anti-HTN meds accordingly
- Women should be discharged with information about signs and symptoms of pre-eclampsia
- Treat w/ MagSO4 for at least 24hrs if patient develops severe hypertension or preeclampsia associated with:
  - Headache not improved with acetaminophen
  - Visual changes (spots in vision)
  - Altered mental status
  - Persistent right upper quadrant or epigastric pain
  - Shortness of breath
  - Thrombocytopenia
  - Abnormal liver enzymes

- Consider straight catheter urine collection for accurate assessment of proteinuria in postpartum period

Women with a history of a pregnancy affected by a diagnosis of least preeclampsia are at increased risk of cardiovascular disease later in life. This risk is twice as high as the baseline risk in all women with preeclampsia, and 4-8x higher in women who had recurrent preeclampsia or a delivery before 34 0/7wga due to preeclampsia.
Thus, additional recommendations include:
- Referral to primary care for evaluation and management of arteriosclerotic cardiovascular disease risk factors
- At least yearly assessment of blood pressure, fasting blood glucose, and BMI
- Pre-conception counseling and assessment prior to next pregnancy

POST-DISCHARGE
In women with hypertension in pregnancy, BP usually decreases within the first 48hrs after delivery, but will increase again around 3-6 days postpartum and should be monitored closely. The majority of PP patients can discontinue BP by 3-4 wks. If still hypertensive at 6 wks, then facilitate a smooth transition to the primary care provider.
- Monitor for 72hr postpartum, if discharged before this time, she should return for blood pressure check at 3 days postpartum
- Blood pressure check again at 7-10 days postpartum

BP >160/110 or symptoms
- Obtain CMP and CBC (and pre-eclampsia screen if no recent PEC)
- Consult with OB
- Prepare to send to nearest hospital.

BP >150/100
- If on nifedipine < 60 mg bid, then increase by 30 mg q day, follow up in 7 days
- If on labetalol, increase by 100 mg bid, follow up in 7 days

BP >140/90
- Continue current regimen, follow up in 7 days

BP < 140/90:
First step
- On labetalol:
  - Decrease daily dosing interval, follow up in 7 days
  - TID to BID, follow up in 7 days
  - BID to QD, follow up in 7 days

- On nifedipine:
  - Decrease daily dosing interval, follow up in 7 days
  - BID to QD, follow up in 7 days

Second step
If on lowest number of daily dose interval already, then
On labetalol: decrease dose size by increment
- 400 mg-> 300 mg, follow up in 7 days
- 300 mg-> 200 mg, follow up in 7 days
- 200 mg ->100 mg follow up in 7 days
- 100 mg-> discontinue

On nifedipine: decrease dose size by increment
- 60 mg -> 30 mg, follow up in 7 days
- 30 mg -> discontinue

BP <110/70 or symptoms of hypotension….
- Stop medication, follow up in 7 days

Bothersome side effects (i.e. HA from nifedipine) switch to labetalol
- If on nifedipine 30 mg BID -> labetalol 200 BID
- If on nifedipine 60 mg BID -> labetalol 300 BID

If PP BP is > 150 / 100 at 5 wks PP, then:
- Change lifestyle
- Consult OB/GYN
- If changes made, then re-appt one week
- Smooth transition to primary care provider at 6 wks PP

Pre-eclampsia and eclampsia can develop up to 6wks postpartum and should be considered when a patient presents with elevated blood pressures, headache, visual changes, and/or right upper quadrant or epigastric pain in the postpartum period.

- If new onset PEC or GHTN, then obtain CMP, CBC, and pre-eclampsia screen
- If prior PEC or GTHN, then it is not necessary to repeat pre-eclampsia screen

If both severe HTN and severe features develop:
- Admit, Consult OB/GYN
- Initiate MagS04 as above

### SUMMARY OF DELIVERY TIMES AND MONITORING

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<th>AFI</th>
<th>Delivery</th>
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<td>1x/wk</td>
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### APPENDICES

I. Measuring Blood Pressure
II. Diagnosing proteinuria
III. Treating antepartum hypertension
IV. Treating severe hypertension
V. Magnesium administration
I. MEASURING BLOOD PRESSURE
The systolic blood pressure is the pressure at which a heartbeat is first heard, the diastolic blood pressure is the pressure at which the sound disappears (Korotkoff phase V). To reduce inaccurate readings:
- An appropriate size cuff should be used (length 1.5 times upper arm circumference or a cuff with a bladder that encircles 80% or more of the arm).
- BP cuff should be placed in middle of upper arm in line with her sternum
- The blood pressure level should be taken with the patient sitting comfortably in an upright position with legs uncrossed after a 10-minute or longer rest period.
- For patients in the hospital, the blood pressure can be taken with either the patient sitting up or in the left lateral recumbent position with the patient's arm at the level of the heart.
- BP should be taken while she is not talking or moving
- The patient should not use tobacco or caffeine for 30 minutes preceding the measurement.
- Although validated electronic devices can be used, a mercury sphygmomanometer is preferred because it is the most accurate device

II. DIAGNOSING PROTEINURIA
Urine must be clean catch specimen and not collected in the setting of urinary tract infection – a dirty catch or UTI can falsely elevate the amount of protein in urine. If there are >5 squamous cells on urinalysis, suspect dirty catch.

Proteinuria is diagnosed when:
- Spot urine w/ protein: creatinine (P:C) ratio ≥ 0.3mg/dL
  - If urine 0.15 ≤ P:C < 0.3 consider 24hr urine protein
- 24hr urine collection w/ ≥ 300mg
- Urine dipstick with ≥ 2+ protein on two separate occasions (only when no other method of evaluation is available)

III. TREATING ANTEPARTUM HYPERTENSION
- Angiotensin-converting enzyme (ACE) inhibitors, angiotensin receptor blockers (ARB) should be discontinued in pregnancy and avoided during pregnancy as they are associated with fetal anomalies
- No difference in outcome or safety between labetalol, methyldopa, and nifedipine.
- Labetalol should be avoided in women with history of asthma, heart disease, and congestive heart failure
- Goal BP is:
  - 120 < SBP < 160
IV. TREATING SEVERE HYPERTENSION

- **GOAL IS TREATMENT WITHIN 30-60 MINUTES**
- No significant difference between the efficacy or safety between hydralazine, labetalol, or nifedipine.
- If no IV, start with immediate release Nifedipine 10mg PO, if not available give Labetalol 200mg PO, repeat in 30min if needed
- Switch to IV medications when available
- Choice of medication should depend on clinician comfort with the medication and patient factors.
- Max out one intravenous agent before moving to a different IV agent – see algorithms below
- Consider starting an oral medication of another mechanism simultaneously with intravenous agent, e.g., if giving labetalol IV consider starting nifedipine XL PO
- Avoid labetalol in setting of bradycardia, history of asthma, heart disease, congestive heart failure
- Goal BP 140-150 systolic, 90-100 diastolic
- Once BP threshold achieved, repeat BP measurement Q 10 minutes x 1 hour, then Q 15 minutes x 1 hour, then Q 30 minutes x 1 hour, and then Q1hr x 4 hours.
V. MAGNESIUM ADMINISTRATION
See Contraindications and Alternatives below.

Eclampsia is a rare outcome that can occur in any patient with preeclampsia even without severe features or while on magnesium therapy. There is not high-quality data to inform the length of magnesium therapy postpartum, and seizures due to eclampsia are less common postpartum. There is some data showing that discontinuation earlier than 24 hours may not increase the risk of seizure. The benefits of earlier discontinuation in appropriate candidates may be improve patient satisfaction and safety as well as decrease time to ambulation, breastfeeding, and hospital discharge.

a. Administering Magnesium with an IV
   - Loading dose or seizure:
     - Administer Magnesium sulfate [40g/liter] with a 4-6g IV bolus over 20min, use with infusion pump
   - Maintenance: 1-2g/hr
   - Monitor:
     - Intake and output, consider placement of foley catheter
     - Reflexes, if depressed -> concern for magnesium toxicity, no established therapeutic level, concern if respiratory or cardiovascular depression
     - BP Q15-30 min and continuous fetal monitoring

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage</th>
<th>Comments</th>
<th>Onset of Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Labetalol</td>
<td>10–20 mg IV, then 20–80 mg every 10–30 minutes to a maximum cumulative dosage of 300 mg; or constant infusion 1–2 mg/min IV</td>
<td>Tachycardia is less common and fewer adverse effects than other agents. Avoid in women with asthma, preexisting myocardial disease, decompensated cardiac function, and heart block and bradycardia.</td>
<td>1–2 minutes</td>
</tr>
<tr>
<td>Hydralazine</td>
<td>5 mg IV or IM, then 5–10 mg IV every 20–40 minutes to a maximum cumulative dosage of 20 mg; or constant infusion of 0.5–10 mg/hr</td>
<td>Higher or frequent dosage associated with maternal hypotension, headaches, and abnormal fetal heart rate tracings; may be more common than other agents.</td>
<td>10–20 minutes</td>
</tr>
<tr>
<td>Nifedipine (immediate release)</td>
<td>10–20 mg orally, repeat in 20 minutes if needed; then 10–20 mg every 2–6 hours; maximum daily dose is 180 mg</td>
<td>May observe reflex tachycardia and headaches.</td>
<td>5–10 minutes</td>
</tr>
</tbody>
</table>

Abbreviations: IM, intramuscularly; IV, intravenously.
- Concern for MagSO4 toxicity: treat with calcium gluconate 10%, give 10cc IV/IO over 5 minutes at a rate of 200 mg/min
- If seizure:
  - If eclampsia occurs with no MagSO4 ongoing, then give 6 gm bolus and start 2 gm/hr
  - If eclampsia occurs while MagSO4 drip already ongoing, then re-bolus with 2-4 gm and increase drip to 3gm/hr IV
  - If additional seizure with MagSO4 drip already ongoing, then re-bolus with 2-4 gm and place drip at 3gm/hr IV
- Obtain a magnesium level when feasible, but do not delay bolus(s) or maintenance level changes while awaiting magnesium results
- Duration:
  - 12-24 hours from delivery
  - Consider 12 hr duration if: Diuresis for two consecutive hours, absence of symptoms [headache, visual changes, epigastric pain], and absence of severe hypertension.

b. Administering Magnesium w/o an IV
- Loading dose or if seizing:
  1. Magnesium sulfate 50% give 5g IM (10mL) in each buttock (10g total) *
- Maintenance: Magnesium sulfate 50% give 5g IM (10mL) every 4-6 hours. *
- If no peripheral IV access, consider interosseous (IO) or central access
- * Add Lidocaine 1% 1mL to each injection to minimize discomfort
- Monitor:
  - Intake and output, consider placement of foley catheter
  - Reflexes, if depressed -> concern for magnesium toxicity
  - No established therapeutic level, concern if respiratory or cardiovascular depression
  - BP Q15-30 min and continuous fetal monitoring
- Concern for MagSO4 toxicity: Treat with calcium gluconate 10%, give 10cc IV/IO over 5 minutes at a rate of 200 mg/min (NB: Not IM)
- Repeat seizure: repeat loading dose
- Duration:
  - 12-24 hours from delivery
  - Consider 12 hr duration if: Diuresis for two consecutive hours, absence of symptoms [headache, visual changes, epigastric pain], and absence of severe hypertension.

VI. ALTERNATIVES TO MAGNESIUM

Contraindications to MagSO4:
Myasthenia gravis, hypocalcemia, moderate to severe renal failure, cardiac ischemia, heart block, or myocarditis.
Use with caution in pulmonary edema

Consult with OB/GYN.

In setting of contraindication to MagSO4 or no MagSO4 available:

- Lorazepam 2-4mg IV x 1, repeat once in 10-15 min if persistent seizure
- Diazepam 5-10mg IV Q 5-10min; max dose 30mg
- Phenytoin 15-20mg/kg IV x 1, repeat 10mg/kg x 1 in 20min if not hypotensive
- Keppra 500mg IV or PO Q12h

VII. ADMINISTERING CORTICOSTEROIDS

- Dosing options
  - Betamethasone 12mg IM Q 12 hours x 2 doses
  - Dexamethasone 6mg IM Q 6 hours x 4 doses

- When to administer in setting of preeclampsia
  - Consider administration of corticosteroids, and initiation of induction of labor with diagnosis of severe preeclampsia and a viable fetus at $\geq$ 34 weeks and < 37 weeks – if the patient has not previously had corticosteroids during this pregnancy.
  - It is suggested that corticosteroids, or rescue corticosteroids as per guideline, be administered and delivery deferred for 48 hours if:
    - Maternal and fetal conditions remain stable for women with severe preeclampsia and a viable fetus at 33 6/7 weeks or less of gestation and the patient has none of the following.
    - Maternal
      - Uncontrolled severe hypertension
      - Persistent headache
      - Persistent epigastric pain
      - Persistent CNS Sx
      - Stroke, MI
      - HELLP
      - Worsening renal function
      - Eclampsia
      - Pulmonary edema
      - Placental abruption

    - Fetal
      - Non-reassuring fetal heart status
      - Intrauterine fetal demise
      - Lethal anomaly or extreme prematurity
      - Persistent reversed end diastolic umbilical artery flow

It is recommended that corticosteroids, or rescue corticosteroids as per guideline, be given if the fetus is viable and at 33 6/7 weeks or less of gestation, but that delivery not be delayed after initial maternal stabilization regardless of gestational age for women with severe preeclampsia which is complicated by any of the above.
VIII. SUMMARY OF DELIVERY TIMES AND MONITORING

<table>
<thead>
<tr>
<th></th>
<th>Start</th>
<th>NST</th>
<th>AFI</th>
<th>Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chr HTN – no meds</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>39-39 6/7 wks</td>
</tr>
<tr>
<td>Chr HTN – controlled on meds</td>
<td>36</td>
<td>1x/wk</td>
<td>1x/wk</td>
<td>39-39 6/7 wks</td>
</tr>
<tr>
<td>Chr HTN – difficult to control</td>
<td>32</td>
<td>2x/wk</td>
<td>1x/wk</td>
<td>≥ 37</td>
</tr>
<tr>
<td>Gestational HTN</td>
<td>at Dx p 32</td>
<td>1x/wk</td>
<td>1x/wk</td>
<td>37-38 wks</td>
</tr>
<tr>
<td>Preeclampsia</td>
<td>at Dx</td>
<td>2x/wk</td>
<td>1x/wk</td>
<td>≥ 37</td>
</tr>
<tr>
<td>Preeclampsia, severe &lt; 34 wks</td>
<td>at Dx</td>
<td>Hosp</td>
<td>Hosp</td>
<td>Steroids</td>
</tr>
<tr>
<td>Preeclampsia, severe &gt; 34 wks</td>
<td>at Dx</td>
<td>Hosp</td>
<td>Hosp</td>
<td>at Dx</td>
</tr>
</tbody>
</table>

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