

Doctor 2019 - نبض - Medicine - MU

Hypertension in Pregnancy

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This sheet contains:

- **lecture slides**
- **Doctors notes**
- **additional notes and pictures from OBS & GYN books**

Introduction

-Hypertensive disorders of pregnancy are one of the leading causes of maternal mortality

-Worldwide: 50,000 women die each year

Hemodynamic Changes in Normal Pregnancy

The pregnancy decreases the baseline of blood pressure.. why?

-Systemic vascular resistance decreases

-Plasma volume and cardiac output increase

There is a physiological drop in BP detectable before the end of the first trimester due to vasodilation

Hypertension

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AHA SCIENTIFIC STATEMENTS

Hypertension in Pregnancy: Diagnosis, Blood Pressure Goals, and Pharmacotherapy: A Scientific Statement From the American Heart Association

-Hypertension is one the most common medical complications in pregnancy

-The major cause of maternal/fetal and neonatal morbidity and mortality

-**Hypertension** is :

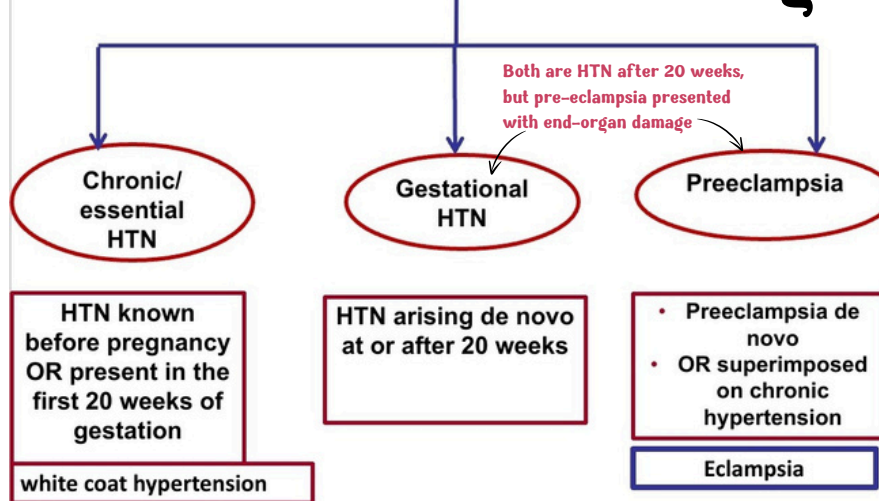
systolic blood pressure of ≥ 140 mm Hg and/or diastolic blood pressure of ≥ 90 mmHg on ≥ 2 occasions 4 hours apart

-**Proteinuria** is the presence of ≥ 300 mg of protein in a 24-hour collection of urine

-OR urinary protein to creatinine ratio of ≥ 30 mg/mmol 0.3 mg /dl

-OR two readings of at least ++ on dipstick analysis of a midstream or catheter urine specimen

Classification of HTN in Pregnancy



Chronic hypertension

-**Diagnosed** before pregnancy or before 20 weeks gestation (and persisting 12 weeks after pregnancy)

It has doubled in prevalence over the last decade and now complicates at least 100,000-pregnancies (2.36%) in the United States each year

The diagnostic criteria for hypertension in pregnancy were similar to those for non-pregnant individuals and included a systolic blood pressure (SBP) of ≥ 140 mm Hg or a diastolic blood pressure (DBP) of ≥ 90 mm Hg on at least 2 separate occasions more than 4 hours apart for mild hypertension

SBP ≥ 160 mm Hg or a DBP ≥ 110 mm Hg for **severe hypertension** (should be treated as we deal with pre-eclampsia)

-**Classified into:**

-Primary (idiopathic) or essential

-Secondary due to

- Renal: glomerulonephritis, renal artery stenosis , Polycystic kidneys .. etc
- Endocrine: crushing's syndrome,Conn's syndrome,Phaeochromocytoma ,

Thyrotoxicosis..etc

- Vascular: coarctation of the aorta

Gestational hypertension

-Hypertension after 20 weeks of gestation with no proteinuria

-High BP $\geq 140/90$ in 2 reading 4 hours apart

What are the effects of HTN on Pregnancy?

Maternal :

- Preeclampsia up to 50 % of those with severe chronic HTN
- Placental abruption up to 10%
- Cesarean delivery
- Cerebrovascular accidents
- Acute renal failure
- Congestive heart disease
- Liver failure
- DIC
- Death

Fetal :

- Fetal growth restriction
- Preterm birth
- Perinatal mortality

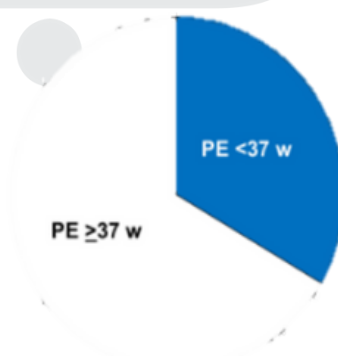
click here to watch
the video

Preeclampsia

- Multisystem syndrome developing during the second half of pregnancy
- characterized by hypertension and proteinuria OR in the absence of proteinuria the finding of maternal organ dysfunction

Preeclampsia superimposed on chronic hypertension

- PET superimposed on chronic hypertension :
- History of hypertension before conception
- the presence of hypertension before 20 weeks Gestation and proteinuria OR maternal organ dysfunction should develop after 20- weeks gestation



Prevalence

In 2-5% of pregnancies

The rate depends on the demographic characteristics of the population in Black women the rate is 2-3 times higher than in White women

In 1/3 of the cases the condition leads to delivery at <37 weeks' gestation (preterm PE) and in 2/3 delivery occurs at ≥37 weeks (term PE)

Maternal organ dysfunction

what are the investigations you should do ?

- Renal insufficiency - serum creatinine $\geq 90 \mu\text{mol/L}$ or 1.04 mg/dl
- Hepatic dysfunction - high serum hepatic transaminase levels (≥ 2 times the upper limit of normal) and/or severe persistent upper abdominal pain unresponsive to medication
- Neurological complications - eclampsia, stroke, confusion, hyperreflexia accompanied by clonus, severe headache accompanied by hyperreflexia, blindness or persistent visual scotomata
- Hematological complications - platelet count $<100,000/\text{dL}$
- Disseminated intravascular coagulation (DIC) or hemolysis
- Pulmonary edema

Complications important for MCQs

Maternal :

- Eclampsia (convulsions or coma in a woman with PET)
- Brain hemorrhage or stroke // -DIC
- HELLP syndrome (Hemolysis, Elevated Liver enzymes and Low Platelets)
- Other severe complications include:
Brain edema Blindness Renal & Hepatic failure Pulmonary edema Death

Long term complications

Doubling in lifetime risk of cardiovascular disease (CVD)
Including Hypertension / Ischemic heart disease / Stroke / Death from CVD

Fetal :

- Reduced blood supply to the placenta
- Impairment in fetal growth oxygenation and increased risk of stillbirth
- Premature delivery for maternal and / or fetal indications
- Babies are subjected to the additional risks arising from prematurity:
 - neonatal death -brain hemorrhage -seizures
- respiratory and feeding difficulties, jaundice, retinopathy, and prolonged hospitalization

-PET and eclampsia are implicated in about 25% of stillbirths and neonatal deaths and 15% of growth restricted neonates

- pre-eclampsia before the 20th week:
- molar pregnancy
 - multiple gestation

treatment of HELLP syndrome is delivery regardless of the gestational age

HELLP Syndrome

Haemolysis

Low platelets

Elevated liver enzymes

Childhood :

- Children exposed to PE before birth compared to those born after normal pregnancy have a doubling in risk of cerebral palsy (this risk is mediated through premature birth growth restriction or both)
- Higher blood pressure /Body mass index/ Increased risk for CVD /Diabetes in adult life

Pathogenesis

- In pregnancy the blastocyst implants into the maternal endometrium
- The outer layer of the blastocyst develops into the trophoblast
- This differentiates into villous trophoblasts and extravillous
- trophoblasts
- Villous trophoblasts give rise to chorionic villi that transport nutrients and oxygen between the fetus and the mother
- Extravillous trophoblasts which invade and transform the spiral arteries
- The trophoblasts replace the endothelial lining and destroy the musculoelastic tissue in the walls of the spiral arteries
- So that they are converted from tortuous narrow muscular vessels into large non-muscular channels thereby increasing maternal blood flow to the placenta

- This physiological process occurs in **two** stages:
- The **first** wave of trophoblastic invasion involves the spiral arteries in the decidua and starts at 8 weeks of gestation (endometrium of pregnancy)
- The **second** involves the spiral arteries in the inner third of the myometrium and occurs at 14-18 weeks

- In PEt**: the physiologic process of placentation is impaired > There is trophoblastic invasion in 50-70% of spiral arteries > does not extend into the myometrial segments > and is confined to the decidual part of the vessels

- The spiral arteries are less dilated than the normally transformed arteries and the blood supply the placenta is reduced
- The reason why in some pregnancies there is failed placentation is unknown but genetic- and immunological reasons may be implicated

Reduced perfusion of the placenta causes **oxidative stress** > release of trophoblast-derived factors > enter the maternal circulation and cause endothelial cell damage in: The kidney, liver, brain and placenta
exaggerated inflammatory response which underlines many of the changes observed in PE

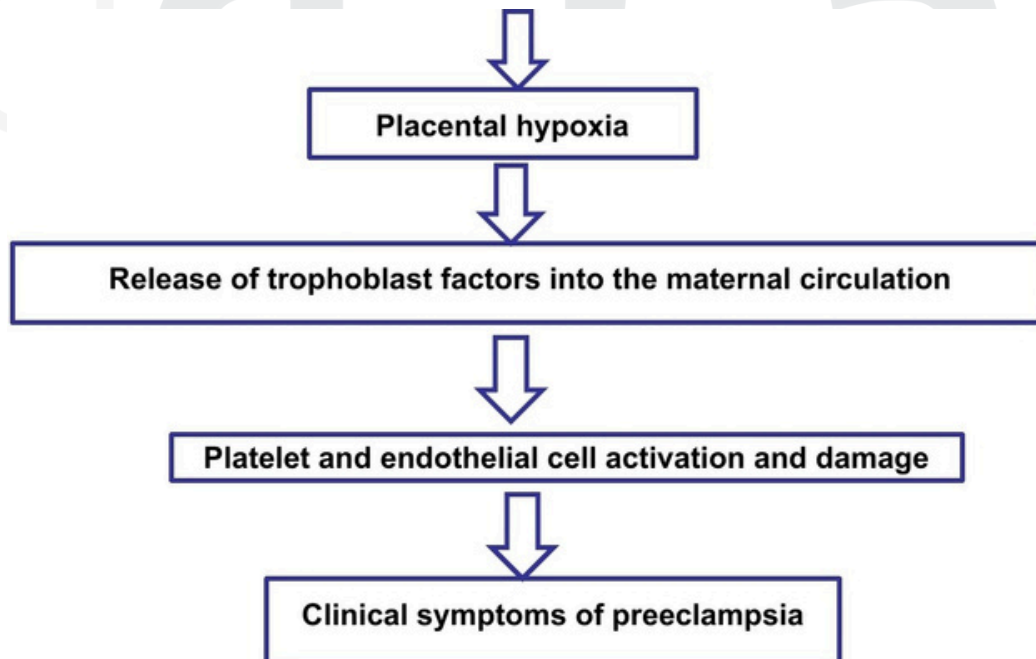
-Placental-derived factors released in response to stress include :
the anti-angiogenic protein **sFLT1** (which is **increased** in PE) whereas the circulating concentration of the angiogenic placental growth factor (**PlGF**) is **reduced** in PE
This angiogenic imbalance results in increased maternal vascular inflammation and generalized endothelial dysfunction

-In **preterm** PE which is characterized by impaired placentation

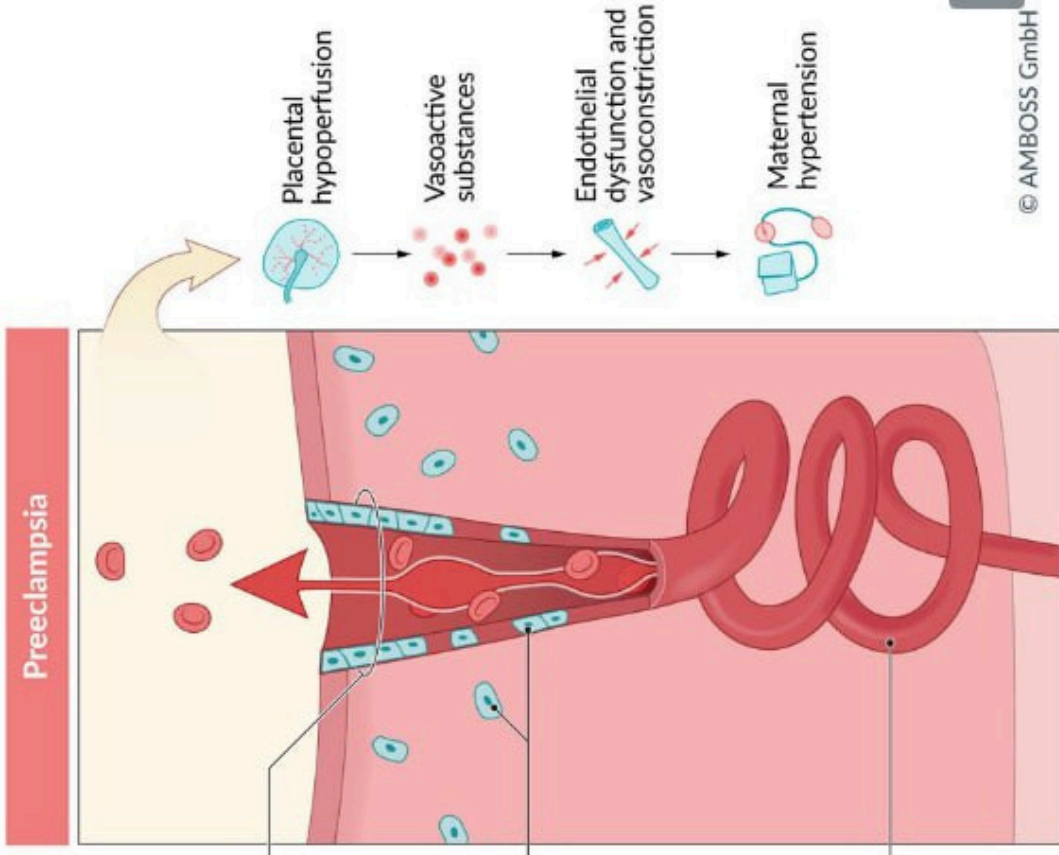
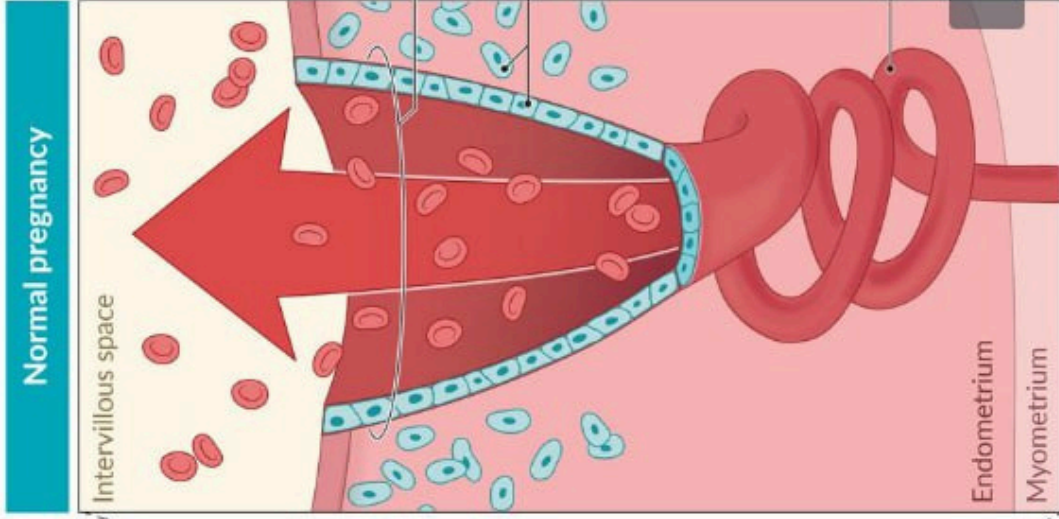
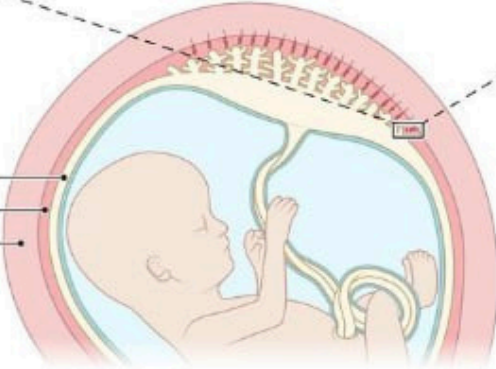
-In **term** PE placentation is usually normal

-In women with **medical disorders**, such as chronic hypertension, there is endothelial dysfunction even before pregnancy in such cases PE can develop in the absence or lower degree of impaired placentation

Impaired trophoblastic invasion of the maternal spiral arteries



Myometrium
Endometrium
Placenta



Prevention of preeclampsia

- In 1543 BC the Egyptians used extracts from the willow tree for pain relief
- In 400 BC Hippocrates used powder made from the bark and leaves of the willow tree for headaches, pains and fevers
- In 1828 Johann Buchner at the University of Munich, extracted the active ingredient of the willow plant and called it salicin (Latin term for willow)
- In 1915 Bayer developed aspirin tablets
- In 1979 Crandon and Isherwood reported that women who had taken aspirin regularly during pregnancy were less likely to have PE than women who had not

The rate of PE is not reduced by:

bed rest

restriction of physical activity or dietary manipulations, including limitation of salt intake or supplementation with magnesium, zinc, folate vitamins C, D, and E, or fish oil

Dietary calcium supplementation in women with low-calcium diets may halve the rate of P
Preliminary data suggest that prophylactic use of pravastatin may also benefit women at high-risk of PET

● **Low dose aspirin** → is given at the 12th or 13th week (before the 16th week) for high-risk patients.

- The prophylactic use of low-dose aspirin in the prevention of PE has been studied extensively
- A meta-analysis of trials showed that the administration of low-dose aspirin in high-risk pregnancies resulted in a 10% lower incidence of PE However in most studies aspirin was started after 16 weeks gestation and at a dose of <100 mg/day
- Other meta-analyses showed that aspirin started before 16 weeks resulted in halving the rate of PE, whereas aspirin started after 16 weeks did not have a significant benefit
- In addition, the beneficial effect of aspirin was dose-dependent, with a greater reduction in the incidence of PE being associated with a dose of >100 mg/day

ASPREE trial

International multicenter study

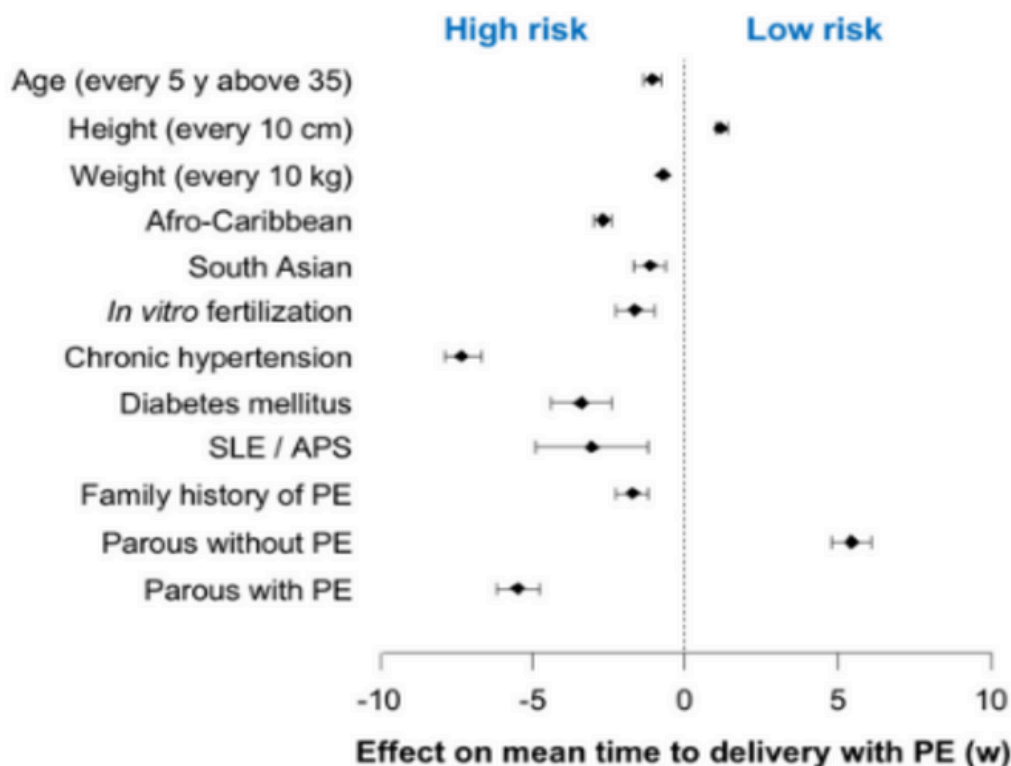
Routine screening for preterm PE was carried out at **11-13 weeks** gestation by the FMF algorithm that combines maternal factors and biomarkers in about 27,000 singleton pregnancies

The use of aspirin was associated with a 62% reduction in the incidence of preterm PE and 82% reduction in the incidence

Maternal risk factors

- Advancing maternal age
- Increasing weight
- Afro-Caribbean and South Asian racial origin
- Medical history of chronic hypertension
- Diabetes mellitus
- Systemic lupus erythematosus or antiphospholipid syndrome
- Conception by in vitro fertilization
- family history or personal history of PE
- The risk of PE in women in their first pregnancy is three times higher than in women with previous pregnancies that were not complicated by PE
- Women who had PE in a first pregnancy are up to 10 times more likely to develop PE in a second pregnancy
- The risk for PE is lower in tall than in short women
- Decreased in parous women with no previous PE
- The protective effect against PE of a previous pregnancy without PE, decreases with the time interval between the previous and the current pregnancy so that after 15 years the risk of PE is about the same as that in nulliparous women

- previous delivery with a normal baby & no complications is a good prognostic factor for this pregnancy
- Smoking is not a cause of pre-eclampsia, it prevents it



Screening

Screening at 11–13 weeks

–Combined screening by maternal risk factors, mean arterial pressure MAP, uterine arteries UTPI and PLGF predicts about 90% of early PE (<34 weeks), 75% of preterm PE (<37 weeks) and 45% of term PE (≥37 weeks)

Mean arterial pressure (MAP)

In the prediction of PE, the measurement of MAP is more useful than the measurement of systolic- and diastolic blood pressure

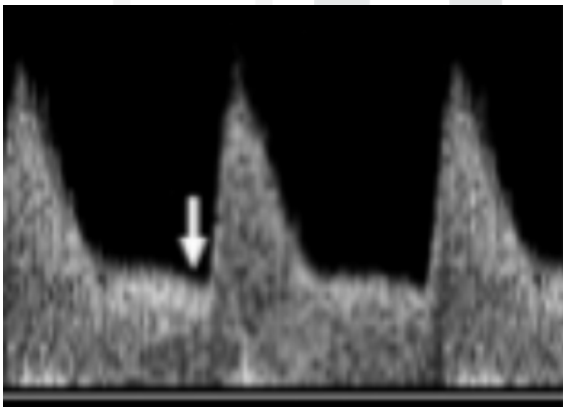
–The MAP is defined as the average arterial pressure during a single cardiac cycle and is calculated from the following formula

MAP = 2/3 diastolic blood pressure + 1/3 systolic blood pressure

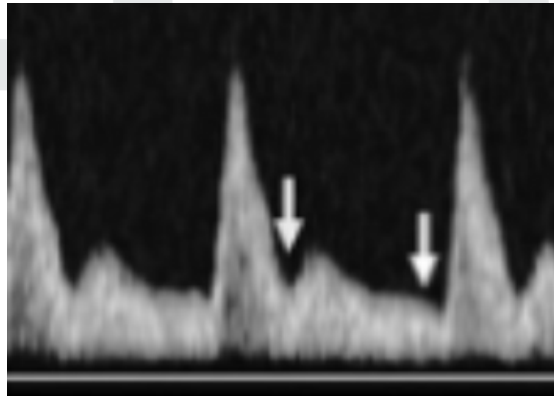
Measurement of uterine artery PI (UTPI)

The UTPI can be measured by either transabdominal or transvaginal sonography in the period between 13–15 weeks & should be measured bilaterally

Waveform has good end-diastolic flow



shows high resistance of flow with **early diastolic notch** and low end-diastolic flow



seeing this notch when screening after the 13th week of gestation, this patient is at risk of pre-eclampsia

Soluble FMS-like tyrosine kinase-1 (sFlt-1)

–An anti-angiogenic factor that is thought to play a central role in the pathogenesis of PE
–Exogenous sFLT-1 administered to pregnant rats induces hypertension, proteinuria, and glomerular endotheliosis **we do this test on the 11th–13th week**
Increase in patients with PE

Placental growth factor PIGF

we do this test on the 11th-13th week
decrease in patients with PEt

- Measure the amount of PIGF in blood plasma or serum
- PIGF is a protein involved in placental angiogenesis (the development of new blood vessels)
- In normal pregnancy PIGF levels rise and peak at 26-30 weeks
- PIGF levels do not rise during pregnancy may be placental dysfunction
- In Preeclampsia level of PIGF can be abnormally low

Pregnancy associated plasma protein-A (PAPP-A)

-PAPP-A is produced by the placenta and is thought to play an important role in placental growth and development

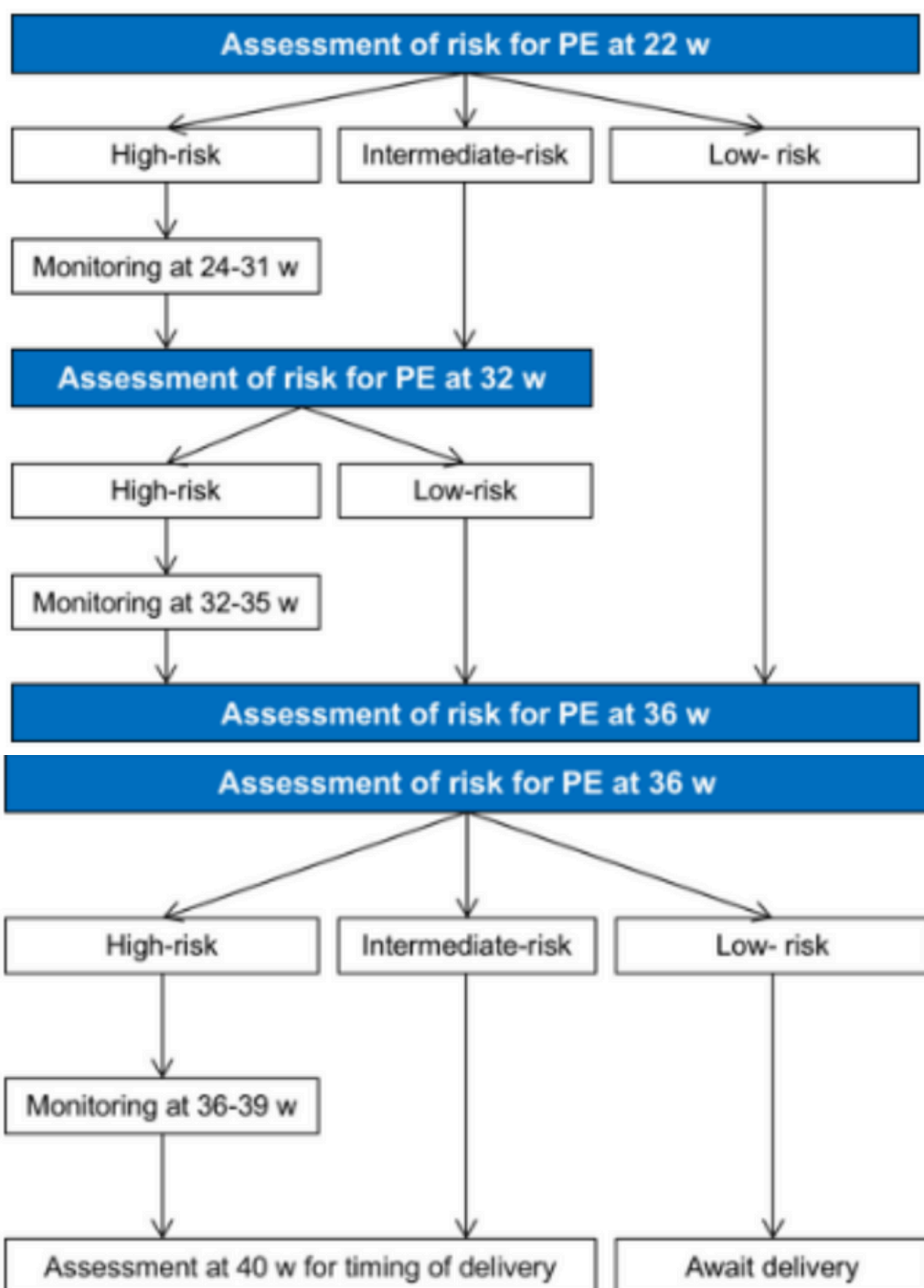
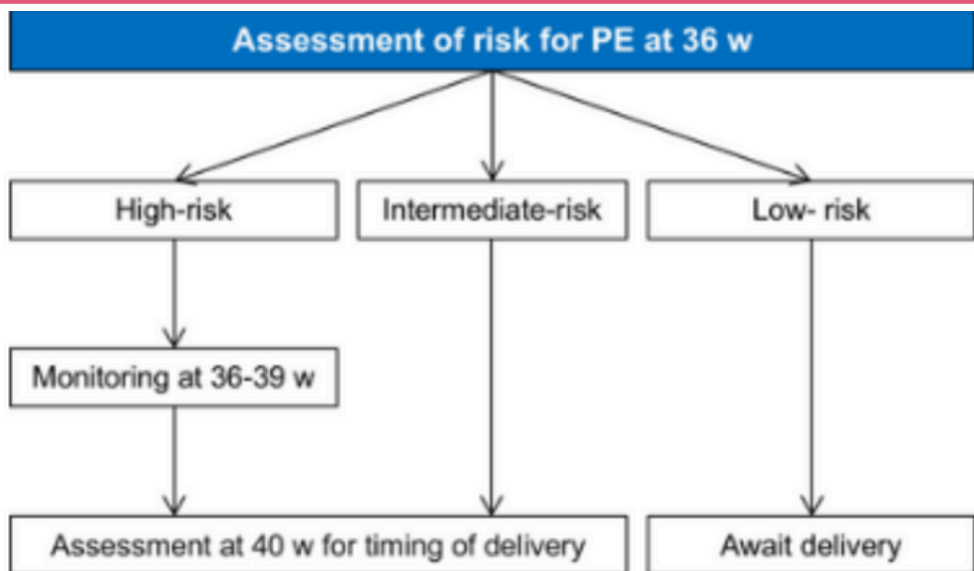
Maternal serum levels of PAPP-A in the first-trimester of pregnancy- are decreased in pregnancies with fetal trisomies 21, 18 and 13

- In pregnancies that develop PE, compared to unaffected pregnancies,- maternal serum PAPP-A is decreased during the first-trimester, not significantly different in the second trimester and increased in the early third-trimester

Screening at 20-24 weeks measure the BP
(if all previous tests are normal)

Screening at 30-34 weeks measure the BP
(if all previous tests are normal)

Method of screening	Detection rate		
	PE <34 w	PE <37 w	PE ≥37
Maternal factors	52%	47%	36%
Maternal factors plus:			
MAP	72%	60%	44%
MAP, UTPI	96%	80%	44%
MAP, PLGF	94%	75%	44%
MAP, sFLT-1	77%	65%	44%
MAP, UTPI, PLGF	94%	85%	45%
MAP, UTPI, PLGF, sFLT-1	100%	85%	45%



Managements

Antenatal appointments

Weekly if HTN poorly controlled or admission

Every 2 to 4 weeks if well controlled

Chronic hypertension:

STOP :

-ACE inhibitors or ARBs (within 2 days of notification of pregnancy)

-Diuretics

ARB cases of fetal renal failure, lung dysplasia, cranial hypoplasia

ACE case fetal renal damage

-All these outcomes may be due to fetal hypotension and reduced renal blood flow perfusion

kidney ischemia - anuria - oligohydramnios

Treatment of chronic hypertension in pregnancy **methyldopa + aspirin**

Start antihypertensive SBP \geq 140 mmHg , DBP \geq 90 mmHg

Consider labetalol

Consider Nifedipine for women in whom labetalol is not suitable

Consider methyldopa if both labetalol and Nifedipine are not suitable

Offer pregnant women with chronic hypertension aspirin 75 mg-150 mg once at night from 12 week

Methyldopa

- crosses the placenta
- increase the incidence of postpartum blues
- his effect is observed within
- 48 hrs, so it CANT be used in emergency!!



Antihypertensives used in pregnancy can be remembered with the mnemonic "Hypertensive Moms Need Love": Hydralazine, Methyldopa, Nifedipine, or Labetalol

Antihypertensive therapy:

Treating hypertension is mainly to reduce maternal complications, It will not improve the fetal condition

Acute treatment of severe hypertension: (In emergency cases)

Hydralazine: 5mg IV repeated every 20–30 min.

Nifedipine: 10mg orally repeated at 30 min. IV infusion can be used in severe cases.

Labetalol: 10–20mg IV .The dose can be doubled every 10 minutes if a proper response is not achieved.

Magnesium Sulphate should be given in the management of all cases of severe preeclampsia to prevent eclampsia

- prevents eclampsia
- prevents seizures & ICH in baby

To prevent eclampsia by MgSO₄:

- give 4–6 g (loading dose) within 10 min
- then give 1–2 g/ hr for a 24 hrs

How to monitor the patient during this period?

يعني كيف أعرف انه الجرعات اللي عم أعطيها للمريضة كافية ومش زيادة؟

we should examine :

- the patient reflexes
- respiratory rate
- UOP

Time of delivery

In chronic hypertension no induce delivery before 37 weeks if

BP lower 160/110

After 37 weeks depends on senior obstetrician decision

If early birth is necessary offer :

- Antenatal corticosteroids
- Magnesium sulfate

Laboratory findings

–Urine analysis ---proteinuria

Microangiopathic hemolytic anemia---elevated serum lactate- dehydrogenase LDH or decreased serum Hepatoglobulin

–Elevated hematocrit ---due to third spacing fluid

–Elevated serum creatinine

–Elevated serum uric acid

–Elevated serum transaminases

–Thrombocytopenia

–Prolonged prothrombin and partial thromboplastin

–Decreased fibrinogen

Increased fibrin degradation products

Symptoms of end-organ damage

- headache (occipital)
- blurred vision
- epigastric pain
- right upper quadrant pain

you should mention them in the history of pregnant patient with hypertension

Eclampsia

- The occurrence of tonic-clonic convulsions (without any neurological disease) in a woman with pre-eclampsia
- Incidence: 5 in 10 000 deliveries and 1-2% of severe PE cases
- High maternal and fetal mortality
- It can occur antenatally, intra-partum and post-partum
- The pathophysiology is cerebral vasospasm leading to ischemia and cerebral edema

Management of eclampsia

During seizure: Maintain airway, Administer oxygen, and avoid supine hypotension

Anticonvulsant therapy:

-Magnesium sulfate 4-6 g IV followed by a maintenance infusion of 1-2 g / h

-Diazepam 20mg IV followed by a maintenance infusion as required

Phenynton

-Anticonvulsant should be continued for at least 24 h after the last convulsion

CS is indicated unless the mother is in active labour

Magnesium Sulfate (MgSO₄)

It can be given IV, IM or SC

The therapeutic level is 4-7mEq/L

The total dose of MgSO₄ should not exceed 24 gms in 24 hours

The dose of MgSO₄ is monitored by :

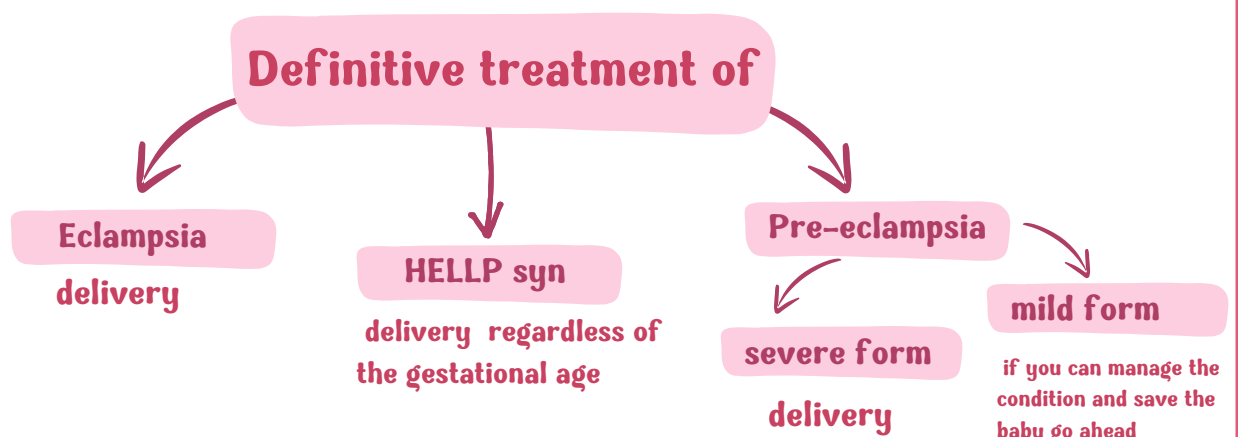
- Preserved patellar reflex. (7-10 mEq/L)
- Respiratory rate >16/min. (10-13 mEq/L)
- Urine output >100ml/4hours. (15-25 mEq/L)
- Serum Mg⁺⁺ level

Is stopped 24 hours after delivery

Antidote is ca gluconate

How to manage MgSO₄ overdose?

- stop it
- give ca gluconate 1g over 10 min



24-3 Hypertension in Pregnancy

