Haematological disorders during pregnancy

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Haematological physiological changes during pregnancy
Haematological physiologic changes during pregnancy

- Anemia < hemodilution (↑ plasma volume up to 40%)
  < iron deficiency
  < folic acid deficiency

Normal hemoglobin level: > 11 g/dl during pregnancy

- Mild elevation of neutrophils

- Platelet count decreases by 10%

→ Left shift of the whole distribution of platelet counts at term

→ Platelets 120-150 x10⁹/l are frequent at the 3rd trimester
Thrombocytopenia during pregnancy
Thrombocytopenia during pregnancy

• The most common haematological abnormality during pregnancy: 7-15%
Thrombocytopenia during pregnancy

• 75% of thrombocytopenia during pregnancy are gestational thrombocytopenia, a benign phenomenon with no significant bleeding-risk for the mother or the fœtus.

• BUT
  – Difficult to distinguish from ITP
  – Thrombopenia can be an indicator of severe complications like eclampsia, DIC, TTP…

• Assessment is required to exclude severe complications and evaluate the bleeding risk for the mother and the fœtus, especially when platelets < 100 x10⁹/l
Differential diagnosis of thrombocytopenia in pregnancy

- **Isolated thrombocytopenia**
  - Gestational
  - ITP
  - Drug-induced
  - Type IIb von Willebrand disease

- **Thrombocytopenia associated with systemic disorders**
  - **Pregnancy specific**
    - Preeclampsia
    - HELLP
    - Acute fatty liver
  - **Not pregnancy specific**
    - Thrombotic microangiopathies: TTP, HUS
    - Systemic lupus
    - Antiphospholipid antibodies
    - Disseminated intra-vascular coagulation (DIC)
    - Viral infection (HIV, CMV, EBV)
    - Bone marrow dysfunction
    - Nutritional deficiency (folate deficiency)
Gestational thrombocytopenia (GT)

- 5-8% of healthy pregnant women
- Mild to moderate thrombopenia (>70-80 x10⁹/l)
- Cause: accelerated platelet consumption + increased plasma volume?
- Generally occurs in the 3rd trimester
- No maternal bleeding risk
- No fetal or neonatal thrombopenia or bleeding risk
- Platelets are normal before pregnancy and return to normal within 2-12 weeks postpartum
- Anti-platelet antibodies can be present (screening not recommended)
Immune thrombocytic thrombocytopenia (ITP) in pregnancy

- 3% of thrombocytopenia in pregnancy
- Moderate thrombocytopenia: $50-100 \times 10^9$ /l but lower
- Auto-immune disorder: antibodies (IgG) against platelet glycoproteins (GPIIb/IIIa and GPIb/IX) → destruction in reticuloendothelial system (spleen)
- Detection of antibodies is possible but not recommended
- Thrombocytopenia can be present before conception or early in pregnancy
- Normal bone marrow biopsy and spleen size
- Exclusion diagnosis
- Risk of maternal bleeding and fetal thrombocytopenia (IgG cross the placenta)
Gestational thrombocytopenia versus ITP

Important to distinguish GT from ITP to optimize management

<table>
<thead>
<tr>
<th></th>
<th>GT</th>
<th>ITP</th>
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<tbody>
<tr>
<td>Before pregnancy</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Mild</td>
<td>++</td>
<td>-/+</td>
</tr>
<tr>
<td>Moderate (&lt; 50 x10^9/l)</td>
<td>-</td>
<td>++</td>
</tr>
<tr>
<td>Antiplatelets antibodies</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>Fetal / neonatal thrombopenia</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Normal platelets in post-partum</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Early in pregnancy</td>
<td>-/+</td>
<td>++</td>
</tr>
<tr>
<td>Late in pregnancy</td>
<td>+</td>
<td>+</td>
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</tbody>
</table>
Recommendation: (All Grade C)
• Asymptomatic women with platelet counts $> 20 \times 10^9/l$ do not need treatment until delivery is imminent
• Platelet counts $> 50 \times 10^9/l$ are safe for normal vaginal delivery in patients with otherwise normal coagulation
• Platelet counts $> 80 \times 10^9/l$ are safe for caesarean section, spinal or epidural anaesthesia in patients with otherwise normal coagulation

*British Journal of Haematology, 2003, 120, 574–596*
**Treatment of ITP during pregnancy**

- **Corticosteroids – IVIG**
- Similar response for both treatments
- No effect on fetal platelet count
- Splenectomy has been performed during 1st or 2d trimester
- Immunosuppressive agents avoided if possible (azathioprine is safe, ciclosporine)
- Rituximab (used in case reports) contra-indicated in pregnancy
- The risk for thrombopenia in the neonates remains in splenectomized mothers (persistance of IgG).
Treatment of ITP during pregnancy

Corticosteroids

- 1-2 mg/kg prednisone
- Response within 7 days
- Tapering after 2-3 weeks
- Side effects: hypertension, hyperglycemia, weight gain, psychosis, osteoporosis
- Indicated if platelets <50 x10^9/l

IVIG

- 0.4 g/kg/d for 5 days
- Response within 1-3 days
- Response duration 2-3 weeks
- Side effects: headache, anaphylactic choc, oedema
- Indicated if:
  - platelets < 50 x10^9/l
  - pre-operatively
  - bleeding with platelets < 30 x10^9/l
  - Cortico-resistance
- Costs!
Delivery in women with ITP

- The mode of delivery in women with ITP should be decided by primarily obstetric indications. There is no evidence to support the routine use of caesarean section (Grade B)

- Women undergoing operative delivery should be considered for thromboprophylaxis according to their individual clinical risk factors. Standard prophylactic doses of UFH or LMW heparin should be used if the maternal platelet count is $>100 \times 10^9/\text{l}$ (Grade C)
- Non-steroidal anti-inflammatory drugs should be avoided for post-partum or post-operative analgesia in women with platelet counts $<100 \times 10^9/\text{l}$ (Grade C)

*British Journal of Haematology, 2003, 120, 574–596*
Epidural analgesia in patients with ITP

- Platelet counts > $80 \times 10^9/l$ are safe for spinal/epidural anaesthesia or caesarean section if coagulation is otherwise normal

*British Journal of Haematology, 2003, 120, 574–596*

**Safe Epidural Analgesia in Thirty Parturients with Platelet Counts Between 69,000 and 98,000 mm$^{-3}$**

*ANESTH ANALG 1997;85:385–8*

Anesthetic management of 52 deliveries in parturients with idiopathic thrombocytopenic purpura

*Journal de Gynécologie Obstétrique et Biologie de la Reproduction 36 (2007) 384-388*
Management of neonate of a mother with ITP

- Antiplatelets antibodies can cross the placenta → fetal thrombopenia (5% of neonates have platelets <20,000/mm³) and 10-15% <50,000)
- Bleeding (intra-cranial haemorrhage) is uncommon
- Evaluation of the fetal platelet count by cordocentesis or fetal scalp sampling are not recommended (risks >> benefits)
- Avoid scalp electrodes, forceps, ventouse
- Measure cord platelets count and monitor platelets until nadir (2-5 days after delivery)
- If neonate’s platelet < 30-50 x10⁹/l : IVIG 1g/kg
- No treatment if neonate’s platelets >50 x10⁹/l
- If life-threatening haemorrhage : IVIG + platelet transfusion
Neonatal alloimmune thrombocytopenia

- Maternal antibodies against alloantigens carried on fetal platelets.
- Cause of severe thrombocytopenia in the fetus and neonate, can produce serious bleeding, intracranial haemorrhage and death.
- Platelet GPs are polymorphic, classified in a system of Human Platelet Antigen (HPA).
- Maternal-fetal incompatibility for HPA-1a is the most common cause of NAIT.
- A first affected neonate with NAIT is suspected when signs of bleeding are evident at or shortly after birth.
- Treatment: PS transfusion +/- IVIG.
Management of subsequent pregnancies

- Test the HPA phenotype of the father and the mother

- Estimate the degree of fetal thrombocytopenia to gauge the risk of antenatal intracranial haemorrhage.
  - Platelet count on a fetal blood sample. (invasive)
  - Mother’s seric titre of the anti-HPA antibody
  - Consider the severity of disease in affected siblings.

- Antenatal therapy to the mother to improve fetal thrombocytopenia and reduce the risk of pre and postnatal bleeding.
  - High dose IVIG with or without corticosteroids
diagnosis FNAIT

obstetric history

sibling without ICH

IVIG weekly 28 wks-birth 0.5 gr/kg

optional: predelivery FBS+IUPT*

induction of labour 37 wks #

Immediate platelet count from cord blood. Matched platelets available, transfuse if <30 x10^9/l. Neonatal head ultrasound.

sibling with ICH

IVIG weekly 16 wks-birth 1.0 gr/kg

Individually based IVIG weekly 1.0 gr/kg

induction of labour #
or elective CS 36 wks

during pregnancy

fetus with ICH

Fetal MRI

Multidisciplinary meeting^*

Individual basis IVIG weekly 1.0 gr/kg

TOP

elective CS 34 wks

M. Kamphuis Prénat Diagn 2011
Hypertensive disorders

- **Preeclampsia**: hypertension + proteinuria + oedema
- Occurs in 6% of all pregnancies
- Thrombopenia in 20-50%
- Platelet consumption (adhesion to damaged endothelium)
- Haemorrhage is uncommon unless DIC develops

- **HELLP** (hemolysis, elevated liver enzymes, low platelets)
- Moderate thrombopenia
- Treatment: MgSO4, delivery
- Platelets transfusion if bleeding or platelets < 50 x10⁹/l
Possible mechanism of thrombopenia in eclampsia and HELLP

Abnormal insertion of the placenta
\[ \Rightarrow \text{utero-placental perfusion} \]
\[ \Rightarrow \text{hypoxia} \]

Endothelial dysfunction

Vasoconstriction

NO+prostacyclins ↔ TXA2 + angiotensin II

Platelets activation
Hemostasis disturbances
Fibrin formation and platelet consumption
Thrombotic microangiopathies (TTP-HUS)

• Classical pentade:
  – Thrombocytopenia
  – Microangiopathic hemolytic anemia
  – Neurologic dysfunction
  – Renal failure
  – Fever

• Biological findings
  – Thrombocytopenia < 20 \times 10^9/l
  – Anemia < 10 g/dl, hemolysis (elevated reticulocytosis, LDH, indirect bilirubin, haptoglobin)
  – Schistocytes

• Treatment: plasmapheresis, fresh frozen plasma, corticoids, immunosuppresive agents, rituximab…
Pathophysiology of TTP

Blood Flow

Adhesion, Rolling, Activation, Recruitment

Protease

No Protease

Thrombus

Normal Multimers

“Ultra-Large” Multimers

(TTP?)
Disseminated intravascular coagulation (DIC)

Systemic activation of coagulation

- Intravascular deposition of fibrin
  - Thrombosis of small and midsize vessels and organ failure
- Depletion of platelets and coagulation factors
  - Bleeding
• Causes of DIC during pregnancy
  – Placenta abruptio
  – Eclampsia and preeclampsia
  – Abortion
  – Acute fatty liver (3rd trimester, abdominal pain, mental disturbances, cholestasis, DIC, hypoglycemia)
  – Intra uterine fetal death (IUFD)
  – Sepsis
  – Amniotic embolism

• Biological findings
  – ↑ prothrombin time and APTT
  – ↓ fibrinogen
  – Severe thrombopenia
  – Microangiopathic hemolytic anaemia (schistocytes)
  – ↑ fibrin and degradation fibrin product (D-dimers), TAT
DIC : treatment

- Treat basic disease
  - Delivery
  - Control hypertension
  - Antibiotics

- Maintain and restore blood volume
  - Transfusion: red blood cell, platelets, FFP
  - Fluids perfusion

- Heparin if thrombosis, antithrombin

- Tranexamic acid if bleeding (with caution)
Other causes of thrombocytopenia during pregnancy

- Infection:
  - Viral: EBV, CMV, HIV
  - Mycoplasma
  - Bacteria
  - Parasites (malaria)
  - Mycobacteria, rickettsia

- Systemic lupus and antiphospholipid syndrome (APS)

- Haematological malignancies

- Congenital:
  - May-Hegglin
  - von Willebrand type IIb
  - Familial
Drug-induced thrombocytopenia during pregnancy

- **Drug induced**: recovery of platelets 5-7 days after withdrawn

<table>
<thead>
<tr>
<th>Class</th>
<th>Medication</th>
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<tbody>
<tr>
<td>Antibiotics</td>
<td>Ampicillin Penicillin Rifampin</td>
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<tr>
<td>Diuretics</td>
<td>Thiazides Furosemide</td>
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<td>Aspirin Acetaminophen Indocin</td>
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<tr>
<td>Anticonvulsants</td>
<td>Phenytoin Valproic Acid Carbamazepine</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>Metyldopa Heparin Digitalis Ranitidine Cimetidine Procainamide Gold compounds Cis-platinum Cyclosporin</td>
</tr>
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</table>
## Thrombocytopenia induced by heparin

- **UFH > LMWH**

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<th>2</th>
<th>1</th>
<th>0</th>
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<tbody>
<tr>
<td>Platelets</td>
<td>20-100 or ↓ 50%</td>
<td>10-20 or ↓ 30-50%</td>
<td>&lt;10 or ↓ 30%</td>
</tr>
<tr>
<td>Timing</td>
<td>5-10 days</td>
<td>&gt; 10 days</td>
<td>&lt; 4 days</td>
</tr>
<tr>
<td>Thrombosis</td>
<td>Proven Cutaneous necrosis</td>
<td>suspected</td>
<td>no</td>
</tr>
<tr>
<td>Other possible cause</td>
<td>no</td>
<td>possible</td>
<td>yes</td>
</tr>
</tbody>
</table>

- Antibodies against PF4+ or platelets aggregation with heparin
- Stop heparin
- Hirudin or daparinoïd
Work-up of thrombocytopenia during pregnancy summary

- Medical and familial history
- Timing of thrombopenia: before pregnancy? Gestational age
- Risk for HIV
- Bleeding (bruising, gingivorragia, menometrorraghia, purpura)
- Drug use
- Physical examination (blood pressure, oedema, splenomegaly, adenopathy, signs of systemic disease…)
- Blood count and peripheral smear
- Liver and renal function, auto-immune, viral serology
- Antiphospholipid antibodies
- Hemostasis tests: APTT, PT, INR, fibrinogen, D-Dimers
- Proteinuria
Pseudothrombopenia

- In vitro artefact usually caused by antibodies antiplatelets against the cryptic site IIb-IIIa unmasked by EDTA
- Control platelets counts with citrate and heparin
- Control platelets count on blood smear (clumps)
Thrombopenia during pregnancy
Exclude pseudothrombocytopenia

No history of ITP, drug, medical disorders

Peripheral smear

History of ITP

Pl < 50
Prednisone
IVIG

Pl > 75
Only low Pl

APL
AAN
HIV, virus

Normal, Pl > 75
probable ITP

Drugs, medical disorder
Stop drugs, treat medical disorder

Hemolytic anemia

Hypertension
Proteinuria
Liver enzymes
Preeclampsia
HELLP

Fever
Neurological Purpura

Bleeding
Thrombosis
Hemostasis

TTP
DIC

Delivery
MgSO4
Plasmapheresis
Delivery transfusion

Follow platelets monthly
Repeat screening if < 75
Haemostasis in normal pregnancy
PREGNANCY = HYPERCOAGULABILITY

Protection from haemorrhage during delivery

Predisposes to thromboembolism
Effects of pregnancy on the clotting system

- Increase of clotting factors
  - ↑ FVIII, ↑ FVII, ↑ FX, ↑ Fibrinogen, ↑ von willebrand factor

- Factors II, V, IX, XI are stable

- Reduction of coagulation inhibitors
  - ↓ Proteins S
  - ↓ Antithrombin
  - Protein C remains stable

- Reduction of fibrinolysis inhibition (hypofibrinolysis)
  - ↑ PAI-1
Pregnancy in women with inherited bleeding disorder
Inherited bleeding disorders

- Carriers of Haemophilia
- Von Willebrand disease
- Other coagulation factor deficiency (FXI,.....)
Issues related to inherited bleeding disorders in pregnancy

- Miscarriages
- Follow-up of pregnancy
- Labour
- Epidural analgaesia
- Delivery
- Fetal complications
- Post-partum haemorrhage
Carriers of haemophilia

• Carriers of haemophilia may have a factor (VIII or IX) deficiency → check factor VIII level at 3rd trimester (FIX will not raise)

• Determine the gender of the baby in case of hemophilia (if male fetus, prenatal diagnosis and genetic counselling should be offered)

• Need for treatment with factor concentrate during pregnancy is rare

• For labour, delivery and postpartum, if factor level < 50 iu/dl
  - Factor concentrate (FVIII, vWF+FVIII, FIX)
  - Desmopressin for HA - VWD
  - Tranexamic acid
VWD during pregnancy

• **Variable and unpredictable increase of**:
  - FVIII
  - von Willebrand factor: > 12 weeks, not in vWD type 3
  → factors levels should be checked during the last trimester

• **Miscarriages**
  – Similar incidence in women with vWD (10-20 %) and without vWD (12-15%)
  – Miscarriages or abortion early in pregnancy carry an increased risk of maternal haemorrhage

• **Vaginal bleeding**
  – During first trimester: 33% in VWD 33% > mothers without vWD (16%)
  – Antepartum: frequency not higher than normal
Epidural analgesia

- Limited reported experience
- Many anaesthetists are reluctant to perform the procedure
- Can be safely performed if FVIII and von Willebrand factor (VWF) antigen and activity levels are above 50% at time of delivery
- Von Willebrand factor concentrates or DDAVP should be administered if FVIII and VWF levels below 50% 
- Any other clotting abnormality should be excluded (low platelets)
- Individualised assessment is required
Primary post-partum haemorrhage
- Blood loss > 500 ml during 24 hrs after birth of the infant
- 3 to 5 % in normal vaginal delivery
- 15 to 18.5 % in women with vWD

Secondary post-partum haemorrhage
- 1.3 % in normal individuals
- 20-25 % in patients with vWD (rapid fall of FVIII and VWF)

Always check VWF and FVIII before delivery and factors + haemoglobin on discharge
Management of peri- and post-partum haemorrhage

- Minimise genital and peritoneal trauma at delivery

- Factor levels should be kept > 50 % for 3-4 days after vaginal delivery and 4-5 days after cesarean section

- At time of delivery, a sample of cord-blood should be collected and carried promptly to the lab to exclude severe deficiency

- Good communication between anaesthesiaiologist, obstetrician and bleeding expert is mandatory
Management of the fetus born to a mother with a bleeding disorder

- Determine the gender of the baby in case of hemophilia
- Reduce the risk of bleeding complications at delivery by avoiding:
  - Instrumental deliveries (vacuum extractions, forceps)
  - Prolonged deliveries
  - Invasive monitoring techniques
- Desmopressin do not contra-indicate breast feeding
- After birth, avoid IM injection of Vit K, IM immunisations and postpone surgery when possible (circumcision)
Other rare bleeding disorders in pregnancy

Global management like hemophilia and vWD

- **Afibrinogenaemia**
  - Infusion of fibrinogen or FFP → Fg level > 100 mg/dl

- **Factor XI deficiency**
  - No raise during pregnancy
  - Correlation between factor level and bleeding is difficult (< 15 iu/dl)
  - Fresh frozen plasma if required

- **Factor VII deficiency**
  - Tranexamic acid, Novoseven®
Management of pregnancies in women with inherited bleeding disorders

**Conception**
- Before conception
  - Measure factor level
  - Genetic counseling
  - ? Risk of transmission
- Determine fetus sex

**Delivery**
- Week 34
  - Control factor level
  - Prepare management plan with anaesthetist, gynecologist and paediatrician

Post-partum
- Control haemoglobin level
- Control neonate's factor level on blood cord
Gestational venous thrombo-embolism (diagnosis, prevention and treatment)
Venous thrombo-embolic disease (VTED)

• Risk of DVT/PE : 5-10 X during pregnancy

• Incidence : 0.86/1000 pregnancies (0.71: DVT - 0.15 : PE)

• 80 % DVT : left lower limb

• Proximal DVT (iliac and/or femoral veins) >> distal DVT during pregnancy
Pregnancy and risk factors of VTE

- **STASIS – Mechanical factors**
  - Slowdown of blood flow
  - Reduction of vascular tone
  - Compression of veins by uterus (left > right)

- **HYPERCOAGULABILITY**
  - Increase of FVIII and VWF
  - Reduction of inhibitors (protein S and antithrombin)
  - Hypofibrinolysis

- **VASCULAR INJURY / LESION**
  - Delivery
Individual risk factors for VTE

• BEFORE PREGNANCY
  – Past history of VTE disease (personal and family)
  – Thrombophilia
  – Age > 35 years
  – Obesity
  – Multiparity
  – Venous insufficiency
  – Nephrotic syndrome

• DURING PREGNANCY
  – Immobility
  – Hyperemesis
  – Dhydratation
  – Ovarian hyperstimulation
  – Pre-eclampsia
  – Air travel
  – Cesarean section
  – Infection or inflammatory process
Model of thrombotic risk
DVT = multifactorial disease

Adapted from Rosendaal FR: Lancet 353: 1167, 1999
Consequences and sequelae of VTE disease

- Maternal mortality
- Venous insufficiency
- Post-thrombotic syndrome (PTS)
- Recurrent DVT

Pulmonary Embolism $\xrightarrow{\text{Recurrent DVT}}$ Post-thrombotic Syndrome

$\text{Healthy Vein Valves} \& \text{Correct Blood Flow}$ $\xrightarrow{\text{Damaged Vein Valve} \& \text{Incorrect Blood Flow}}$
Antithrombotic agents

- **VKA (Sintrom, Marevan, Marcoumar)** contra-indicated (relative CI)
  - Crosses the placenta
  - Risk of bleeding
  - Teratogenic (6-12 weeks)

- **LMWH**: anticoagulant agent of choice >> UFH
  - Excellent bioavailability and predictable effect
  - Does not cross the placental barrier
  - Short half-life
  - Well-tolerated (rare thrombocytopenia, osteoporosis)

- **NOAC**: contra-indicated in pregnancy
Treatment of VTE during pregnancy

• **Doses and duration:**
  - 100 anti-Xa units /kg, twice a day (avoid once a day treatment)
  - If distal DVT : 100 U anti-Xa units /kg, once a day after 3 months of treatment
  - Throughout pregnancy and at least 6 weeks postpartum
  - 24 hours between last injection and epidural at therapeutic concentration

• **Monitoring (anti-Xa and platelets count):**
  - Anti-Xa activity should be measured after one week, then every 6 weeks
  - Expected value : 0.5-1.0 anti-Xa unit / ml (2-4 h post-injection)
  - Mandatory if < body weight <50 or >100 kg, renal insufficiency, increased bleeding risk
  - Platelet count : risk of Heparin Induced Thrombocytopenia (HIT) low with LMWH during pregnancy, full blood count after a week then 1x/month
Pregnant Women With Past DVT/PE

- Prophylaxis for 6 weeks after delivery (Grade 2B) for all

- Moderate/high recurrence risk should consider taking prophylactic or intermediate-dose LMWH during pregnancy (Grade 2C). This includes women with a prior DVT or PE that was unprovoked, or related to pregnancy or estrogen use.

- Low risk for recurrence (because their VTE was provoked by a transient risk factor unrelated to pregnancy or estrogen use) and their doctors should watch carefully for signs or symptoms of DVT or PE during pregnancy, but ACCP suggests they not use LMWH as prophylaxis (Grade 2C).
VTE prophylaxis during pregnancy
ACCP 2012

• 1 episode of VTE with transient risk factor and no thrombophilia
  → surveillance antepartum and prophylaxis postpartum

• 1 episode of idiopathic VTE without long term anticoagulation and no thrombophilia
  → surveillance or prophylaxis antepartum and prophylaxis postpartum

• 1 episode of VTE with thrombophilia or multiple episodes of VTE
  → prophylaxis antepartum and prophylaxis postpartum

• Prior VTE and long-term anticoagulation
  → intermediate or full dose LMWH antepartum
VTE prophylaxis during pregnancy
ACCP 2012

- No prior VTE but severe thrombophilia (homozygous) or combined thrombophilia and familial history of VTE
  → prophylaxis antepartum and prophylaxis postpartum (grade 2B)

- No prior VTE and other thrombophilia
  → surveillance antepartum and prophylaxis postpartum (grade 2C)

- Prior VTE and long-term anticoaguation
  → intermediate or full dose LMWH antepartum
Epidural analgesia and LMWH

- Insertion of the needle when the anticoagulant effect is minimal (8-12 hours after last prophylactic injection and 24 hours after last therapeutic injection)

- Prophylactic dose should be given at least 4-6 hours after insertion/removal of catheter
Thrombophilia

- Inherited or acquired predisposition towards thrombosis
- Primarily associated with an increased risk of venous thromboembolic disease
- Can be associated with adverse pregnancy outcomes (APL)
Thrombophilia screening: recommendations

- Avoid pregnancy (< 12 weeks) and hormonal treatment
- Has a screening already been performed in the past?
- Known thrombophilic trait in the family??
- Informed consent and appropriate information regarding personal and family implications (insurance)
- Is screening useful and indicated?
- Eligibility for re-imbursement (cost > 400-800 euros)
Who should be screened for thrombophilia

- Family history of VTE
- Idiopathic VTE (or after a trivial provocation)
- VTE < 60 years old
- Recurrent VTE
- Unusual VTE
- Suspicion of APS

- A screening makes sense only if the care of the patient or her/his family will be modified
Thrombophilia and risk of DVT during pregnancy

- Basal risk: 1/1500
- FV Leiden: 1/500 – 1/437
- G20210A FII mutation: 1/200
- FVL + FII mutations: 4.6/100
- Protein C deficiency: 1/113
- Antithrombin deficiency (quantitative defect): 1 / 2.8
- Antithrombin deficiency (qualitative defect): 1 / 42
Ovarian stimulation

- Thromboprophylaxis with LMWH (3800 à 5000 anti/Xa units/day):
  - Past history of DVT/PE
  - Risk factors for DVT-PE
  - Known thrombophilia
  - Ovarian hyperstimulation
Cerebral venous sinus thrombosis

- Incidence: 4/1,000,000 /yr
- Common causes: trauma, auto-immune, pregnancy, puerperium, OC
- Risk increased by OC (20 x)
- Risk = two-fold risk with 3rd vs 2nd generation pills
- Risk higher in women with thrombophilia
  - Factor V Leiden: X 34
  - Prothrombin mutation: X 149.3 (31-711)
Prosthetic heart valves

- Anticoagulation is required throughout pregnancy in women with mechanical heart valve(s)

- Treatment should be individualised after careful counselling (valve type, position, VTE history)

- Options of anticoagulation in pregnant women with prosthetic heart valve(s)
  - Oral anticoagulants (VKA) throughout pregnancy (risk of embryopathy)
  - Replace VKA with Heparins (weeks 6-12) (UFH or LMWH)
  - Heparins throughout pregnancy (very few studies on the study and efficacy of LMWH in this setting)
Arterial thromboembolism in pregnancy
<table>
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<tr>
<th></th>
<th>Venous thrombosis</th>
<th>Arterial thrombosis</th>
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<tbody>
<tr>
<td><strong>Stasis</strong></td>
<td>+++</td>
<td>+</td>
</tr>
<tr>
<td>↑ Blood coagulability</td>
<td>+++</td>
<td>+</td>
</tr>
<tr>
<td>Altered vessel wall</td>
<td>+</td>
<td>+++</td>
</tr>
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</table>
Ischaemic stroke

- Pregnancy is associated with an increased risk of stroke
- No clear consensus about the incidence
- 5 cases / 100,000 deliveries
- Infarctions > haemorrhages
- Arterial > venous infarctions
- Risk higher in the post-partum
- Most common cause = eclampsia and pre-eclampsia
Thrombophilia and fetal loss / gestational vascular complications
Adverse pregnancy outcomes potentially related to thrombophilia

- Recurrent unexplained miscarriages (3 or >)
- Pre-eclampsia
- Unexplained IUGR (intra-uterine growth retardation)
- Unexplained IUD (intra-uterine fetal death)
- Abruptio placentae
## Placenta vasculopathy and thrombophilia

<table>
<thead>
<tr>
<th>Thrombophilia</th>
<th>Miscarriages</th>
<th>IUFD</th>
<th>Pre-eclampsia</th>
<th>HELLP</th>
<th>Abruptio placentae</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antithrombin deficiency</td>
<td>++</td>
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<tr>
<td>Protein C deficiency</td>
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<tr>
<td>Protein S deficiency</td>
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<tr>
<td>Dysfibrinogenemia</td>
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<tr>
<td>APC-resistance</td>
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<tr>
<td>Factor V Leiden</td>
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<tr>
<td>MTHFR C677T</td>
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<tr>
<td>Hyperhomocysteinaemia</td>
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<td>Prothrombin mutation</td>
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<tr>
<td>Antiphospholipid syndrome</td>
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<tr>
<td>Combined defects</td>
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<td>++</td>
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</tbody>
</table>
Thrombophilia and complicated pregnancy: evidence and controversies

• With the exception of Anti-Phospholipid antibodies, there is little evidence to support systematic screening for thrombophilia in women with complicated pregnancy.

• In our experience, combined treatment with LMWH and aspirin with close monitoring although not validated is safe and efficient in most cases to prevent recurrent complications.

• For women with past history of eclampsia and/or pre-eclampsia and no thrombophilia, low dose aspirin is recommended.
Antithrombotic therapy in women with complicated pregnancy and thrombophilia

- Low-dose enteric coated aspirin + LMWH (prophylactic dose)
- Dosis: 3800-4000 anti-Xa units / day (injection in the morning in order to allow monitoring with the anti-Xa assay)
- Stop aspirin at week 34
- Continuation of LMWH until delivery and 4 to 6 weeks post-partum
- For women with past history of eclampsia and/or pre-eclampsia and no thrombophilia, low dose aspirin is recommended
Thank you for your attention