Pre-eclampsia: pathophysiology, diagnosis, and management

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Abstract: The incidence of pre-eclampsia ranges from 3% to 7% for nulliparas and 1% to 3% for multiparas. Pre-eclampsia is a major cause of maternal mortality and morbidity, preterm birth, perinatal death, and intrauterine growth restriction. Unfortunately, the pathophysiology of this multisystem disorder, characterized by abnormal vascular response to placentation, is still unclear. Despite great polymorphism of the disease, the criteria for pre-eclampsia have not changed over the past decade (systolic blood pressure ≥140 mmHg or diastolic blood pressure ≥90 mmHg and 24-hour proteinuria ≥0.3 g). Clinical features and laboratory abnormalities define and determine the severity of pre-eclampsia. Delivery is the only curative treatment for pre-eclampsia. Multidisciplinary management, involving an obstetrician, anesthetist, and pediatrician, is carried out with consideration of the maternal risks due to continued pregnancy and the fetal risks associated with induced preterm delivery. Screening women at high risk and preventing recurrences are key issues in the management of pre-eclampsia.

Keywords: pre-eclampsia, epidemiology, pathophysiology, therapeutic management

Introduction

The criteria that define pre-eclampsia have not changed over the past decade. These are: onset at >20 weeks' gestational age of 24-hour proteinuria ≥30 mg/day or, if not available, a protein concentration ≥30 mg (≥1+ on dipstick) in a minimum of two random urine samples collected at least 4–6 hours but no more than 7 days apart, a systolic blood pressure >140 mmHg or diastolic blood pressure ≥90 mmHg and 24-hour proteinuria ≥0.3 g). Clinical features and laboratory abnormalities define and determine the severity of pre-eclampsia. Delivery is the only curative treatment for pre-eclampsia. Multidisciplinary management, involving an obstetrician, anesthetist, and pediatrician, is carried out with consideration of the maternal risks due to continued pregnancy and the fetal risks associated with induced preterm delivery. Screening women at high risk and preventing recurrences are key issues in the management of pre-eclampsia.

Epidemiology

Pre-eclampsia is a multisystem disorder that complicates 3%–8% of pregnancies in Western countries and constitutes a major source of morbidity and mortality worldwide. Overall, 10%–15% of maternal deaths are directly associated with pre-eclampsia and eclampsia. Some epidemiological findings support the hypothesis of a genetic and immunological etiology. The risk of pre-eclampsia is 2-fold to 5-fold higher in pregnant women with a maternal history of this disorder. Depending on ethnicity, the incidence of pre-eclampsia ranges from 3% to 7% in healthy nulliparas and 1% to 3%
in multiparas. Moreover, nullparity and a new partner have been shown to be important risk factors (Table 1).

Other risk factors have been identified, including a medical history of chronic hypertension, kidney disease, diabetes, obesity, birthplace in Africa, age ≥35 years, and pregnancy characteristics, such as twin or molar pregnancy, previous pre-eclampsia, or fetal congenital abnormality. High altitude has also been shown to increase the incidence of pre-eclampsia, and is attributed to greater placental hypoxia, smaller uterine artery diameter, and lower uterine artery blood flow.

Pre-eclampsia may be life-threatening for both mother and child, increasing both fetal and maternal morbidity and mortality. In the mother, pre-eclampsia may cause premature cardiovascular disease, such as chronic hypertension, ischemic heart disease, and stroke, later in life, while children born after pre-eclamptic pregnancies and who are relatively small at birth, have an increased risk of stroke, coronary heart disease, and metabolic syndrome in adult life.

The sole curative treatment being delivery, management must continuously balance the risk–benefit ratio of induced preterm delivery and maternal–fetal complications. Screening women at high risk and preventing recurrences are also key issues in the management of pre-eclampsia.

Pathophysiology
During normal pregnancy, the villous cytotrophoblast invades into the inner third of the myometrium, and spiral arteries lose their endothelium and most of their muscle fibers. These structural modifications are associated with functional alterations, such that spiral arteries become low-resistance vessels, and thus less sensitive, or even insensitive, to vasoconstrictive substances.

Pre-eclampsia has a complex pathophysiology, the primary cause being abnormal placentation. Defective invasion of the spiral arteries by cytotrophoblast cells is observed during pre-eclampsia. Recent studies have shown that cytotrophoblast invasion of the uterus is actually a unique differentiation pathway in which the fetal cells adopt certain attributes of the maternal endothelium they normally replace. In pre-eclampsia, this differentiation process goes awry. The abnormalities may be related to the nitric oxide pathway, which contributes substantially to the control of vascular tone. Moreover, inhibition of maternal synthesis of nitric oxide prevents embryo implantation.

Increased uterine arterial resistance induces higher sensitivity to vasoconstriction and thus chronic placental ischemia and oxidative stress. This chronic placental ischemia causes fetal complications, including intrauterine growth retardation and intrauterine death. In parallel, oxidative stress induces release into the maternal circulation of substances such as free radicals, oxidized lipids, cytokines, and serum soluble vascular endothelial growth factor 1. These abnormalities are responsible for endothelial dysfunction with vascular hyperpermeability, thrombophilia, and hypertension, so as to compensate for the decreased flow in the uterine arteries due to peripheral vasoconstriction.

Endothelial dysfunction is responsible for the clinical signs observed in the mother, ie, impairment of the hepatic endothelium contributing to onset of the HELLP (Hemolysis, Elevated Liver enzymes and Low Platelet count) syndrome, impairment of the cerebral endothelium inducing refractory neurological disorders, or even eclampsia. Depletion of vascular endothelial growth factor in the podocytes makes the endotheliosis more able to block the slit diaphragms in the basement membrane, adding to decreased glomerular filtration and causing proteinuria. Finally, endothelial dysfunction promotes microangiopathic hemolytic anemia, and vascular hyperpermeability associated with low serum albumin causes edema, particularly in the lower limbs or lungs.

The crucial issue to understand is that the prime mover of pre-eclampsia is abnormal placentation. Two common theories appear to be interlinked, ie, a genetic theory and an immunological theory. Several susceptibility genes may exist for pre-eclampsia. These genes probably interact in the hemostatic and cardiovascular systems, as well as in the inflammatory response. Some have been identified, and in candidate gene studies they have provided evidence of linkage to several genes, including angiotensinogen on 1q42–43 and eNOS on 7q36; other main important loci are 2p12, 2p25, 9p13, and 10q22.1.

Pre-eclampsia can be perceived as an impairment of the maternal immune system that prevents it from recognizing the fetoplacental unit. Excessive production of immune cells causes secretion of tumor necrosis factor alpha which induces apoptosis of the extravillous cytotrophoblast. The human leukocyte antigen (HLA) system also appears to play a role in the defective invasion of the spiral arteries, in that women with pre-eclampsia show reduced levels of HLA-G and HLA-E. During normal pregnancies, the interaction between these cells and the trophoblast is due to secretion of vascular endothelial growth factor and placental growth factor by natural killer cells. High levels of soluble fms-like tyrosine kinase 1 (sFlt-1), an antagonist of vascular endothelial growth factor and placental growth factor, have been found in women with pre-eclampsia.
Accordingly, assays of sFlt-1, placental growth factor, endoglin, and vascular endothelial growth factor, all of which increase 4–8 weeks before onset of the disease, may be useful predictors of pre-eclampsia. Recent data show the protective role of heme oxygenase 1 and its metabolite, carbon monoxide, in pregnancy, and identify this as a potential target in the treatment of pre-eclampsia.20

**Clinical presentation and workup findings**

Clinical and laboratory tests are intended to define and determine the severity of pre-eclampsia. Headaches, tinnitus, phosphene signals, visual disorders, brisk tendon reflexes, and vigilance disorders are related to cerebral edema; oliguria to acute renal failure; uterine contraction, vaginal bleeding to placental abruption; vomiting to HELLP syndrome; band-like epigastric pain to subcapsular hepatic hematoma; and dyspnea to cardiac failure. Eclampsia, the major neurological complication of pre-eclampsia, is defined as a convulsive episode or any other sign of altered consciousness arising in a setting of pre-eclampsia, and which cannot be attributed to a pre-existing neurological condition. Clinical examination should include resting blood pressure measurement using an appropriate cuff, and screening for weight gain, edema (including signs of acute pulmonary edema and cerebral edema), cardiomyopathy, and acute renal failure. The fetus should be assessed by electrocardiotocography. Laboratory tests include: a complete blood count with platelets, haptoglobin, and lactate dehydrogenase; a blood smear to test for schistocytes; bilirubin, aspartate transaminase, and alanine transaminase in order to identify potential HELPP syndrome; electrolyte, urea, and creatinine assessment to check for acute renal failure or uremia; 24-hour proteinuria; prothrombin, activated thrombin time, and fibrinogen (microangiopathic hemolytic anemia); blood group; and irregular antibody screening. Other examinations include fetal ultrasound with Doppler velocimetry of the umbilical, cerebral, and uterine arteries, estimation of fetal weight, assessment of fetal well-being by Manning score, and examination of the placenta.21

Although the definition of severe pre-eclampsia varies,1,21,22 several components of this definition are usually accepted: maternal systolic blood pressure ≥160 mmHg or diastolic blood pressure ≥110 mmHg; maternal neurological disorders such as persistent headaches, phosphene signals, tinnitus, and brisk, diffuse, polykinetic tendon reflexes, eclampsia, acute pulmonary edema, proteinuria ≥5 g/day, oliguria <500 cc/day, creatinine >120 μmol/L, HELLP syndrome, thrombocytopenia <100,000/mm³, and fetal criteria, especially intrauterine growth retardation, oligohydramnios, or fetal death in utero. Mild pre-eclampsia is defined as diastolic blood pressure ≥90 mmHg measured on two occasions at least 6 hours apart, combined with proteinuria (two or more occurrences of protein on dipstick, >300 mg total protein in a 24-hour urine collection, or a protein creatinine ratio >30 mg/mmol).1

**Immediate emergency management**

Delivery is the only curative treatment for pre-eclampsia.1 Management is multidisciplinary, involving an obstetrician, an anesthetist, and a pediatrician. In some cases consultation of maternal fetal medicine and hypertension or nephrology subspecialists may be required. Management decisions must balance the maternal risks of continued pregnancy against the fetal risks associated with induced preterm delivery.2 The criteria for delivery are based on two often interrelated factors, ie, gestational age at diagnosis (estimated fetal weight) and severity of pre-eclampsia.

Severe pre-eclampsia requires treatment with a dual aim, ie, preventing the harmful effects of elevated maternal blood pressure and preventing eclampsia. Management of severe pre-eclampsia begins with transfer of the mother in a fully equipped ambulance or helicopter to a maternity ward providing an appropriate level of care for both mother and child.2 At admission and daily thereafter, clinical, cardiotocographic, laboratory, and ultrasound testing are required to detect the severity of pre-eclampsia and tailor management accordingly.22

Regardless of the severity of pre-eclampsia, there is no advantage in continuing the pregnancy when pre-eclampsia is discovered after 36–37 weeks.23–25 Nor is expectant management justified for severe pre-eclampsia before 24 weeks, in view of the high risk of maternal complications and the poor neonatal prognosis.26–28 The obstetric team must then discuss with the parents the possibility of a medical interruption of pregnancy. Prolongation of pregnancy in the event of mild pre-eclampsia can be discussed and re-evaluated on a regular basis. At 34–37 weeks, management depends on the severity of the pre-eclampsia. Expectant management is possible for mild pre-eclampsia to limit the risk of induced preterm delivery, but for severe pre-eclampsia, delivery remains the rule due to the increased risk of maternal and fetal complications.2,22

Similarly, at 24–34 weeks, management depends on the severity of pre-eclampsia. The presence of one or
more of the following signs indicates the need for imme-
diate delivery: uncontrolled severe hypertension (not
responsive to dual therapy), eclampsia, acute pulmonary
edema, abruptio placenta, subcapsular hepatic hematoma,
or thrombocytopenia <50,000/mm³. Delivery after
corticosteroid therapy for pulmonary maturation is necessary
if any of the following criteria is present: persistent epigastric
pain, signs of imminent eclampsia (headaches or persistent
visual disorders), de novo creatinine >120 µmol/L, oliguria
below 20 mL/hour, progressive HELLP syndrome, prolonged
or severe variable decelerations with short-term variability
less than 3 milliseconds. When emergency delivery is not
required, labor can be induced by cervical ripening.22

Antihypertensive treatment is useful only in severe pre-
eclampsia because the sole proven benefit of such management
is to diminish the risk of maternal complications (cerebral
hemorrhage, eclampsia, or acute pulmonary edema).29 There
is no international consensus concerning antihypertensive treat-
ment in pre-eclampsia. The four drugs authorized for the treat-
ment of hypertension in severe pre-eclampsia in France are
nicardipine, labetalol, clonidine, and dihydralazine.29 There
is no ideal target blood pressure value, and too aggressive a
reduction in blood pressure is harmful to the fetus.30 Therapy
with a single agent is advised as first-line treatment, followed
by combination treatment when appropriate. The algorithm
for antihypertensive treatment proposed by French experts22
is shown in Figure 1.

Pulmonary maturation using corticosteroids must be con-
sidered, taking gestational age into account. Betamethasone
remains the gold standard at a dosage of two injections of
12 mg 24 hours apart; this treatment reduces the risk of hyaline
membrane disease, intraventricular hemorrhage, and
neonatal mortality.31

Magnesium sulfate (MgSO4), may be part of the
therapeutic armamentarium for severe pre-eclampsia. It
is indicated in the treatment of eclamptic convulsions as
well as for secondary prevention of eclampsia, thus replac-
ing treatment by diazepam, phenytoin, or the combination
of chlorpromazine, promethazine, and pethidine.32 The
efficacy of MgSO4 in the reduction of maternal and neo-
natal complications of eclampsia is well established. It is
administered intravenously, first at a loading dose of 4 g
over 15–20 minutes, which can be repeated at a half dose
(2 g) if convolution recurs, and then at a maintenance dose
of 1 g/hour for 24 hours.32 MgSO4 treatment must be
monitored in the intensive care unit because organ failure
may occur. This monitoring is based on repeated check-
ning for a Glasgow score of 15, tendon reflexes, respiratory
frequency >12 per minute, and diuresis >30 mL/hour. Any
manifestation of overdose requires stopping the infusion,
considering injection of calcium gluconate, and measuring
blood magnesium levels.22

Eclampsia is generally considered an indication for
emergency cesarean section. Nonetheless, a decision to
delay a cesarean, albeit rare, may be based on fetal status
and justified if the mother’s condition is stable and reassuring
after treatment.33

Management following delivery
Although delivery is the only effective treatment for
pre-eclampsia, and despite the fact that clinical symptoms
and laboratory abnormalities usually regress in the hours
afterwards, the risk of complications persists for some time
following delivery.6 Pre-eclampsia is associated with long-
term morbidity and mortality. Approximately 20% of women
with pre-eclampsia develop hypertension or microalbuminu-
ria during long-term follow-up, and the risk of subsequent
cardiovascular and cerebrovascular disease is doubled in
women with pre-eclampsia and gestational hypertension
compared with age-matched controls.34 A recent prospective
epidemiological study with a median follow-up duration of
30 years provides evidence that pre-eclampsia is a marker of
increased mortality from cardiovascular disease.35

Hemodynamic, neurological, and laboratory monitoring
is necessary following delivery for patients with severe pre-
eclampsia.36 Hemodynamic monitoring includes frequent
blood pressure measurements to enable adjustment of anti-
hypertensive treatment and frequent monitoring of diuresis
and weight according to intake (oliguria should prompt
progressive fluid resuscitation and sometimes diuretic use).
Neurological monitoring consists of checking for signs of
imminent eclampsia, including headaches, phosphene sig-
nals, tinnitus, and brisk tendon reflexes. Clinical monitor-
ing must be done several times daily during the week after
delivery, a period considered at high risk for complications.
If necessary, monitoring can be performed in an intensive
care unit.

Laboratory monitoring should be done several times daily
in the first 72 hours after delivery and thereafter adapted
according to progress of the indices. It must include a
complete blood count, liver function tests, and measurement
of lactate dehydrogenase. Discharge from hospital cannot
be considered until all clinical and laboratory indices have
returned to normal, and regular monitoring by the patient’s
general practitioner as necessary if treatment for hypertension
is to be continued after discharge.
The risk of recurrence of pre-eclampsia during a subsequent pregnancy has to be considered. This risk is estimated to be less than 10% for all cases of pre-eclampsia, but is greater when pre-eclampsia is discovered before 28 weeks. The relative risk is 15 if pre-eclampsia occurs at 20–33 weeks, 10 at 33–36 weeks, and 8 after 37 weeks.

Three months after delivery, screening for underlying renal or hypertensive disease may be requested by the patient’s primary physician. Such screening is intended to check for normalization of blood pressure values and disappearance of proteinuria, and if abnormalities persist, a referral should be made to a nephrologist or a hypertension expert to determine the cause. This examination is important because pre-eclampsia may unmask previously undiagnosed systemic or kidney disease or thrombophilia. It should include a specific set of questions, blood pressure measurement, a clinical examination looking for signs of autoimmune conditions, and a urinary dipstick test. Testing for antiphospholipid antibodies is recommended after severe pre-eclampsia. The search for hereditary thrombophilia by assays for protein C and S, antithrombin III, and a test for resistance to activated protein C is recommended in the case

Figure 1 Algorithm for antihypertensive treatment of pre-eclampsia.22
Note: MBP = [systolic BP + 2 × diastolic BP]/3.
Abbreviations: MBP, mean blood pressure; CI, contraindication; SBP, systolic blood pressure; DBP, diastolic blood pressure.
Table 1 Major risk factors for pre-eclampsia49

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>OR or RR (95% CI)</th>
</tr>
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<tbody>
<tr>
<td>Antiphospholipid antibody syndrome</td>
<td>9.7 (4.3–21.7)</td>
</tr>
<tr>
<td>Renal disease</td>
<td>7.8 (2.2–28.2)</td>
</tr>
<tr>
<td>Prior pre-eclampsia</td>
<td>7.2 (5.8–8.8)</td>
</tr>
<tr>
<td>Systemic lupus erythematosus</td>
<td>5.7 (2.0–16.2)</td>
</tr>
<tr>
<td>Nulliparity</td>
<td>5.4 (2.8–10.3)</td>
</tr>
<tr>
<td>Chronic hypertension</td>
<td>3.8 (3.4–4.3)</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>3.6 (2.5–5.0)</td>
</tr>
<tr>
<td>High altitude</td>
<td>3.6 (1.1–11.9)</td>
</tr>
<tr>
<td>Multiple gestations</td>
<td>3.5 (3.0–4.2)</td>
</tr>
<tr>
<td>Strong family history of CV disease</td>
<td>3.2 (1.4–7.7)</td>
</tr>
<tr>
<td>(heart disease or stroke in ≥2 first-degree relatives)</td>
<td></td>
</tr>
<tr>
<td>Obesity</td>
<td>2.5 (1.7–3.7)</td>
</tr>
<tr>
<td>Family history of pre-eclampsia in first-degree relative</td>
<td>2.3–2.6 (1.8–3.6)</td>
</tr>
<tr>
<td>Advanced maternal age (&gt;40 years)</td>
<td>1.68 (1.23–2.29)</td>
</tr>
<tr>
<td></td>
<td>1.96 (1.34–2.87)</td>
</tr>
</tbody>
</table>

Abbreviations: CI, confidence interval; OR, odds ratio; RR, relative risk; CV, cardiovascular.

of a personal or family history of venous thromboembolic disease, early pre-eclampsia, or pre-eclampsia with any intrauterine growth retardation, abruptio placentae, or in utero death.22,38 Percutaneous needle biopsy of the kidney should be performed only if kidney failure persists at three months postpartum or if signs of a systemic underlying condition or proteinuria persist at 6 months.23

Patients who have had severe pre-eclampsia may share predispositions with nonpregnant patients who have cardiovascular risk factors.39 Accordingly, long-term monitoring of cardiovascular, renal, and metabolic risk factors is recommended after severe pre-eclampsia.22

Prevention

Primary prevention of pre-eclampsia is based on the detection of modifiable risk factors. The literature is plentiful regarding the risk factors for pre-eclampsia, but should be interpreted with caution.4–8 Women at high risk are those with a personal history of severe pre-eclampsia, while those at low risk are defined as those who have never had pre-eclampsia but have at least one risk factor.9 There are numerous risk factors,4–8 including genetic risk factors, family history of pre-eclampsia, immunologic factors, nulliparity, a new partner, and demographic factors such as a maternal age >35 years, the woman’s own gestational age and birth weight (with elevated risks for women born before 34 weeks or weighing less than 2500 g at birth), factors related to the pregnancy, such as multiple pregnancy, congenital or chromosomal anomalies, a hydatidiform mole, or urinary infection, risk factors associated with maternal disease, including chronic hypertension, kidney disease, obesity, insulin resistance, and diabetes, as well as thrombophilia, and environmental factors such as living at a high altitude and stress. Although the search for these risk factors is important, they may not effectively predict this pre-eclampsia by themselves.

However, accurate prediction of pre-eclampsia would enable early and optimal management of women at high risk. Several predictive tests are being assessed currently. These include clinical tests, such as blood pressure measurement during the second trimester or 24-hour ambulatory blood pressure monitoring, but these lack sensitivity and specificity.40 Laboratory tests for oxidative response have been assessed, including assays for uric acid, urinary kallikrein, and fibronectin, but no evidence of their relevance has so far been found.40 Among the markers used to screen for trisomy 21 during the second trimester (beta human chorionic gonadotropin, alpha fetoprotein, and unconjugated estriol), elevated alpha fetoprotein is associated with a higher risk of pre-eclampsia (unless there are neural tube abnormalities, as when beta human chorionic gonadotropin is elevated). Frequent monitoring of women with elevated levels could be useful, but these tests may not be carried out for screening purposes due to their low negative predictive value.40 Serum markers for trisomy 21 in the first trimester (pregnancy-associated plasma protein A, inhibitin A, corticotropin-releasing hormone, and activin) have been tested, but their likelihood ratios seem to be insufficient.

Imaging tests have been evaluated, including uterine artery Doppler ultrasound.41,42 Uterine artery Doppler ultrasound is not advised during the first or second trimester in low-risk populations due to the excessive variability of likelihood ratios in this population, which allows for the prediction of only one-third of pre-eclampsia cases.43 In a high-risk population, the definition of which is often imprecise, uterine artery Doppler can be performed during the second trimester morphologic ultrasound examination and checked 1 month later in case of abnormal results (resistance index >0.58 or 90–95th percentile, unilateral or bilateral notch). The combination of a uterine artery Doppler examination during the first trimester and a three-dimensional ultrasound assessing placental volume may predict the risk of pre-eclampsia as early as the first trimester.44

In clinical practice, because no single marker effectively predicts the risk of pre-eclampsia, the current trend is to test a combination of markers. The most commonly used combination of markers assesses sFlt-1, placental growth
factor, endoglin, and vascular endothelial growth factor during the first or second trimester. Increased vascular endothelial growth factor and endoglin levels, combined with increased sFlt-1 and decreased placental growth factor during the first trimester, is associated with a significantly increased risk of pre-eclampsia.  

Improved prediction of pre-eclampsia has been noticed when serum markers are combined with Doppler indices. In a recent nested case-control study, second trimester maternal serum cystatin C, C-reactive protein, and uterine artery mean resistance index were observed to be independent predictors of pre-eclampsia.  

Secondary prevention is based on antiplatelet aspirin therapy, which reduces the risk of pre-eclampsia by 10% in women who have at least one risk factor. No study currently allows determination of the exact dosage or the best time for initiation of aspirin. However, aspirin should be initiated as early as possible, ie, before 12–14 weeks, which corresponds to the beginning of the first phase of trophoblast invasion. The efficacy of aspirin has been shown only in women with previous pre-eclampsia associated with intrauterine growth retardation and without thrombophilia. Low molecular weight heparin is indicated only in cases of complicated thrombophilia (history of thromboembolic complications or of pre-eclampsia). Calcium supplementation at a dosage of 1.5 g/day, beginning at 15 weeks and continued throughout the pregnancy, is recommended for prevention of pre-eclampsia in women with a daily calcium intake <600 mg/day. The statins, which stimulate HO-1 expression and inhibit sFlt-1 release, could have the potential to ameliorate early-onset pre-eclampsia. Other treatments, such as antioxidant treatment by vitamins C and E, oligoelements, and nitric oxide have no proven efficacy.

**Conclusion**

Pre-eclampsia is a rare pregnancy-related disease with an unpredictable course that can have serious consequences for both the mother and the fetus. The treatment is simple, ie, delivery. Nonetheless, induced preterm delivery requires careful weighing of both maternal and fetal risk–benefit. Accordingly, identifying delivery criteria in case of pre-eclampsia is crucial to optimal management. Current research focuses on the prediction of onset of pre-eclampsia or even severe pre-eclampsia so as to allow early management and improve the morbidity and mortality associated with this disease. Specific tools for secondary prevention must also be developed for recurrent pre-eclampsia.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


