### Combined intramuscular (IM) and intravenous (IV) regimen

**Loading dose:**
- Add 8ml 50% MgSO₄ (4g) in 100ml 0.9% saline or 5% glucose
  → administer IV over 20min
- OR, if you have syringe-driver pump
  - Add 8ml 50% MgSO₄ (4g) to 12ml 0.9% saline or 5% glucose and
  → infuse IV over 20min at 60ml.h⁻¹
- AND
- Give 2.5g MgSO₄ IM into each buttock
  (Total initial dose 4g IV + 2 x 2.5g IM = 9g)

**If the convulsions do not stop:**
Administer a further 2g MgSO₄
- Draw 4ml (2g) of 50% MgSO₄ into 10ml syringe and add 6ml 0.9% saline or 5% glucose
  → inject over 2min (5ml.min⁻¹)

**Do not exceed 9g total IV dose of MgSO₄ during the first hour**
- If convulsions still continue, consult medical staff and consider diazepam 5mg or lorazepam 1mg (IV or IM)
- Be aware of risk of respiratory depression

**Maintenance**
- 2.5 MgSO₄ IM 4 hourly using alternate buttocks if there are no signs of MgSO₄ overdose
- Check reflexes before giving MgSO₄
- Continue for 24 hours after the last convulsion or delivery

### Intravenous (IV) regimen

**Loading dose:**
- Fill a paediatric infusion burette set with 22ml 5% glucose
  - Add 8ml 50% MgSO₄ (4g)
  → administer at 60ml.h⁻¹ - the total will run over 30min

**If the convulsions do not stop:**
As above for combined IV/IM regime

**Maintenance**
- Fill a paediatric infusion burette with 112ml 5% glucose
  - Add 8ml 50% MgSO₄ (4g)
  → administer at 30ml.h⁻¹ - the total will run over 4 hours (1g.h⁻¹)
- Repeat the same management every 4hr for at least 24 hours after the last convulsion or delivery

**For recurrent siezures**
- Administer a second loading dose or increase the infusion to 1.5 or 2g.h⁻¹

### Adverse effects of MgSO₄
- hypotension, arrhythmias
- respiratory depression
- flushing, nausea/vomiting
- drowsiness, slurred speech, double vision
Management of severe pre-eclampsia and eclampsia

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INTRODUCTION
Administration of magnesium sulphate is one of the prime treatment modalities in eclampsia and severe pre-eclampsia. Figure 1 is a guideline for administration of magnesium sulphate to treat severe pre-eclampsia and eclampsia. The guideline has been drawn up with reference to the guidance of the Royal College of Obstetricians and Gynaecologists,1 with additional guidance for those working in settings where infusion pumps are unavailable and intramuscular administration of magnesium sulphate is more practical.2 Use of magnesium is described, along with other modes of treatment, in the context of a case history.

DEFINITIONS
Pre-eclampsia is the most likely diagnosis. Pre-eclampsia is a multisystem disorder which occurs after 20 weeks of pregnancy with variable features, severity and rates of progression. There are a number of definitions of hypertension in pregnancy which lack consistency and can be confusing.

In essence, high blood pressure in pregnancy can be:
• Pre-existing hypertension
• Pregnancy induced hypertension
• Pre-eclampsia.

Most definitions of hypertension in pregnancy are based on a diastolic BP >90mmHg on two occasions or diastolic >110mmHg on one occasion. Failure to record and treat systolic hypertension in women with severe pre-eclampsia was highlighted as a common problem in the most recent Confidential Enquiry into Maternal and Child Health (CEMACH). Treatment was recommended if the systolic BP was >160mmHg, on two consecutive readings at least 4 hours apart.

ASSESSMENT

Table 1. Symptoms and signs of pre-eclampsia

<table>
<thead>
<tr>
<th>Symptoms</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiovascular system</td>
<td>Hypertension, vasoconstriction leading to cool peripheries, peripheral oedema</td>
</tr>
<tr>
<td>Respiratory system</td>
<td>Pulmonary oedema, facial and laryngeal oedema, acute respiratory distress syndrome (ARDS)</td>
</tr>
<tr>
<td>Renal system</td>
<td>Proteinuria, oliguria, acute renal failure</td>
</tr>
<tr>
<td>Central nervous system</td>
<td>Hyperreflexia, clonus, cerebral haemorrhage, convulsions (eclampsia), papilloedema, coma</td>
</tr>
<tr>
<td>Others</td>
<td>HELLP (Haemolysis, Elevated Liver Enzymes and Low Platelets), thrombocytopenia, DIC (disseminated intravascular coagulopathy)</td>
</tr>
<tr>
<td>Foetal signs</td>
<td>CardioTocoGraphy (CTG) abnormalities, pre-term labour, and intrauterine growth retardation.</td>
</tr>
</tbody>
</table>
Some definitions of hypertension rely on a rise in BP, rather than an absolute value, e.g. a rise in systolic BP of 30mmHg, or diastolic BP of 25mmHg above the earliest BP taken in pregnancy. It is essential to monitor blood pressure closely throughout pregnancy and to identify other signs and symptoms suggestive of pre-eclampsia. The distinction between pregnancy induced hypertension and pre-eclampsia is important since pre-eclampsia is associated with worse outcomes. Pregnancy induced hypertension, like pre-eclampsia, occurs in the second half of pregnancy, but without proteinuria or other signs of pre-eclampsia.

**Urine**

More recently, urine protein: creatinine ratio (PCR): a PCR >30 is significant. Assessment of urine output is important.

**Blood tests**

Consider full blood count, urea and electrolytes, clotting, uric acid, liver function tests, magnesium, serum calcium, group and save.

**Cardiotocogram**

Regular assessment of fetal well-being using CTG, ultrasound scan to assess fetal growth and uterine artery Doppler blood flow to assess placental blood flow.

**CASE HISTORY**

A 28 year old woman in her first pregnancy is admitted to the labour ward at 38 weeks of gestation. She has no past medical history of significance. Her blood pressure when pregnancy was first confirmed at 8 weeks was 120/70. Today she presents with a mild frontal headache and increasing swelling of her ankles. Blood pressure is 170/120, urine dip stick testing shows 3+ of protein and there is oedema of both ankles to the mid-calf.

- What is the most likely reason for these clinical signs?
- What other symptoms and signs should be looked for?
- What investigations should be performed?

**Severe pre-eclampsia**

Clinical features of severe pre-eclampsia (in addition to hypertension and proteinuria) are:

- symptoms of severe headache
- visual disturbance
- signs of clonus
- papilloedema
- epigastric pain and/or vomiting
- abnormal liver enzymes (ALT or AST rising to above 70IU.l⁻¹)
- liver tenderness
- HELLP syndrome
- platelet count falling to below 100x10⁶.l⁻¹

**INVESTIGATIONS**

Proteinuria demonstrated by urine dipstick and 24hr collection of urine:

<table>
<thead>
<tr>
<th>Level</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1+</td>
<td>0.3g.l⁻¹</td>
</tr>
<tr>
<td>2+</td>
<td>1g.l⁻¹</td>
</tr>
<tr>
<td>3+</td>
<td>3g.l⁻¹</td>
</tr>
</tbody>
</table>

**Labetalol**

Control of acute hypertension in pre-eclampsia may be achieved by:

- Labetalol bolus – 25mg IV bolus (5ml of 5mg.ml⁻¹ neat solution) over at least 1 minute
- Repeat above at 15 minute intervals to a maximum dose of 200mg until blood pressure is controlled and then start infusion:
- Labetalol maintenance infusion – dilute 200mg (40ml of 5mg.ml⁻¹ neat labetalol with 10ml 0.9% sodium chloride gives a final concentration of 4mg.ml⁻¹)
- Commence infusion at 5ml.hr⁻¹ (20mg.hr⁻¹)
- Double infusion rate every 30 min to maximum 40ml.hr⁻¹ (160mg.hr⁻¹)
- Titrate to keep diastolic between 90 – 100mmHg

Labetalol should be avoided in women with asthma.
If the woman remains hypertensive on maximum rate or labetalol is contraindicated or causing side effects, add / replace with a hydralazine infusion.

**Hydralazine bolus**
- Dilute 40mg hydralazine in 40ml 0.9% saline to give a concentration of 1mg.ml⁻¹
- Give 5ml (5mg) slowly (e.g. over 15 minutes using an infusion pump set at 20ml.hr⁻¹)
- Check BP. After 20 minutes, if diastolic >100mmHg give further 5ml (5mg) over 15 minutes (pump rate 20ml.hr⁻¹)
- When diastolic 90–100mmHg commence maintenance infusion.

**Hydralazine infusion**
- Using same concentration as above set pump rate to 5ml.hr⁻¹ (5mg.hr⁻¹)
- Titrate to keep diastolic 90-100mmHg and systolic 140 - 150mmHg
- Usual maintenance dose is 2-3ml.hr⁻¹ (2-3mg.hr⁻¹)
- Maximum dose 18ml.hr⁻¹ (18mg.hr⁻¹)
- Reduce if significant side effects (see below) or maternal tachycardia >130bpm.

Hydralazine causes headache, tremor, nausea and tachycardia and may be less well tolerated than labetalol

A fluid bolus of 250ml should be considered before commencing IV antihypertensive therapy as there is some evidence that this may avoid the hypotension observed with initiation of vasodilator therapy.

**CASE HISTORY**
Given the severity of this woman's symptoms, measures to control BP with an IV agent should have been considered earlier in this case.

A bolus dose of labetalol is given, followed by a continuous infusion of labetalol and her BP starts to stabilise and her diastolic falls to 90mmHg. A litre of Hartmann's solution (Ringers lactate) is commenced at a rate of 85ml/hr, aiming for urine output of >100ml in 4 hours (excessive fluid administration is harmful in severe pre-eclampsia and fluids should be restricted).

Observations of BP, oxygen saturations, heart rate and respiratory rate are performed every 15 minutes along with continuous CTG monitoring. While considering a plan for delivery she has a grand mal fit.

**What is the most likely reason for the fit?**

**How should it be managed?**
The fit is most likely to be an eclamptic seizure. Other causes of a fit in this situation include:

- Epilepsy
- Intracranial event (e.g. subarachnoid haemorrhage, cerebrovascular accident)
- Vaso-vagal (may be caused by rapid fall in BP due to treatment)
- Hypoglycaemia.

**MANAGEMENT OF AN ECLAMPTIC SEIZURE**
- The patient should be turned to the left lateral position
- Call for help
- Assess and support Airway, Breathing and Circulation
- High flow oxygen by face mask
- Obtain IV access
- Treat with IV magnesium sulphate (see Figure 1)
- Monitor ECG, BP, respiratory rate and oxygen saturations
- Check blood sugar

It is beneficial for obstetric departments to set up an “eclamptic box” for use in this kind of emergency. Within the box is stored: magnesium sulphate, normal saline, syringes, needles and instructions for the correct dose and administration. This will improve the timely administration of magnesium and reduce errors in dosing and administration.

Would earlier administration of magnesium have reduced the risk of fitting in this particular case?

The drug of choice to manage eclampsia and reduce subsequent fits is magnesium sulphate. The use of magnesium to prevent seizures in women with pre-eclampsia is less clear. Guidelines on the management of pre-eclampsia by the Royal College of Obstetricians and Gynaecologists (RCOG) in 2006, recommended prophylactic magnesium sulphate should be considered for women with severe pre-eclampsia for whom there is concern about the risk of eclampsia. This is based on the Magpie Trial. This study showed that magnesium sulphate given to women with pre-eclampsia reduced the risk of an eclamptic seizure by around 58%. It should be noted however that not all women with pre-eclampsia will progress to eclampsia; only 1–2% of women in the UK with pre-eclampsia will fit in the absence of anticonvulsant treatment. The Magpie trial calculated that the number of pre-eclamptic women needed to be treated with magnesium to prevent one of them from fitting is 91 (i.e. of 91 pre-eclampsics receiving magnesium, 1 would gain benefit by not fitting while 90 will derive no benefit from its administration and may be at risk of its side effects.)

**CASE HISTORY**
A bolus dose of magnesium is administered and she is commenced on an infusion of magnesium at a rate of 1g.h⁻¹. Her respiratory rate and reflexes are checked regularly. She is nursed in the left lateral position.

The CTG now demonstrates repetitive and severe fetal heart rate decelerations.

She is reviewed by the senior obstetrician and a decision is made for an emergency delivery by Caesarean section (category 1). The anaesthetist attends to make a plan for anaesthesia.

**What is the most appropriate anaesthetic technique? GA or regional?**
Women with severe pre-eclampsia should be encouraged to have regional anaesthesia for caesarean section.

Regional anaesthesia is not contraindicated after an eclamptic fit if the mother has regained consciousness and treatment for seizures and BP control has been commenced. If an epidural has already been in place for labour, this can be topped up as long as it has been effective. If there is no epidural (as in this case) or it is considered that there is not enough time to top up the epidural, spinal anaesthesia should be provided.

The benefits of spinal anaesthesia include provision of a rapid, dense and predictable block suitable for surgery while avoiding general anaesthesia which has the risk of BP surges due to the pressor response of laryngoscopy, intubation and extubation.

There have been concerns with spinal anaesthesia and severe pre-eclampsia due to the fear of causing a sudden and significant drop in BP. This fear appears unfounded because women with pre-eclampsia have high levels of circulating catecholamines which may protect them against a fall in BP as a result of a spinal induced sympathetic block. If a spinal technique is chosen there should be cautious use of vasopressors as an exaggerated hypertensive response to vasopressors may be observed.

Sometimes a regional block may be contraindicated, for example: maternal refusal, coagulopathy, thrombocytopenia and poorly controlled seizures or there is no time because of severe foetal distress. In these cases general anaesthesia will have to be undertaken.

The factors which make general anaesthesia in pre-eclampsia particularly hazardous include:
- the increased risk of difficult airway and intubation
- marked pressor response at laryngoscopy, intubation and extubation resulting in dangerous surges in blood pressure

There is a significant risk of intracranial haemorrhage secondary to uncontrolled severe hypertension at induction of general anaesthesia. Therefore in patients who require general anaesthesia, BP and convulsions should be maximally controlled and ideally, invasive monitoring inserted prior to induction of general anaesthesia. A senior anaesthetist should be present to manage these challenging cases.

CASE HISTORY

The blood tests which were sent earlier are now available and demonstrate a platelet count of 55x10^9/l. She is assessed by the anaesthetist and consented for a general anaesthetic. A thorough assessment of her airway is performed and she denies any history of stridor, voice change or hoarseness. She is transferred to theatre and full monitoring is attached to the patient. An arterial line is inserted prior to induction. She is pre-oxygenated in the left tilted position and a rapid sequence induction is performed with thiopentone 450mg and suxamethonium 100mg. No other drugs are administered prior to laryngoscopy. The trachea is intubated after 45 seconds with a size 7 endotracheal tube at which point her BP surges to 240/140. Anaesthesia is maintained with isoflurane in nitrous oxide and oxygen, she is paralysed with atracurium 40mg and prophylactic antibiotics are given. After delivery of the baby she is given morphine 10mg.

At the end of the caesarean section she is slow to wake up.

How can you obtund the pressor response to laryngoscopy and intubation?

What might be the cause of her slow recovery?

The response to laryngoscopy can be obtunded by the following:
- A short acting opiate bolus (e.g. alfentanil 10-20mcg.kg^-1 or remifentanil 1mcg.kg^-1)
- A bolus dose of labetalol 10-20mg IV
- Bolus of magnesium 40mg.kg^-1 IV
- Bolus of lidocaine 1.5mg.kg^-1 IV 3-5 min before induction

If opiates are given prior to delivery, inform the neonatal team of the possibility of neonatal respiratory depression.

There are several reasons why she is slow to wake up and these include:
- Effect of excess anaesthetic agents
- Effect of excess opiates
- Inadequate reversal of neuromuscular block: magnesium potentiates non-depolarising neuromuscular blocking drugs
- Respiratory depression due to magnesium toxicity
- Hypoglycaemia.

The most concerning possibility is that she has experienced an intracranial event due to excessive hypertension during intubation. This should be diagnosed by pupil examination and response and emergency CT scan.

Fortunately, over the next 15 minutes she slowly wakes and makes an uneventful recovery from her general anaesthetic.

What on-going management of her pre-eclampsia should be continued?

After her caesarean section this lady should go to a high dependency area for close observation and blood tests.

It is important she also has:
- Effective postoperative analgesia as this reduces the stress response and consequent hypertension caused by poorly controlled pain; however NSAIDs should be avoided until proteinuria has resolved and there is no evidence of impairment of renal or platelet function.
• Ongoing anti-hypertensive therapy. This should be continued after the delivery as dictated by her blood pressure. Although blood pressure usually falls initially after delivery it may rise again around 24 hours postpartum. IV antihypertensive therapy should be converted to oral therapy with further reductions made in a stepwise fashion.

• Close monitoring must continue as she is at risk of further eclamptic seizures. Magnesium therapy should be continued until 24 hrs after her delivery (or the last convulsion, whichever is the later).

• Cautious fluid intake. She should continue with a total of 85 ml h⁻¹ of fluids (subtracting oral intake from IV prescription). Fluid overload must be avoided and transient rises in plasma urea and creatinine concentrations are acceptable in the short term as a spontaneous diuresis is usual within 1-2 days of delivery.

Most women with severe pre-eclampsia or eclampsia will need inpatient care for 4 days or more following delivery.

SUMMARY
Administration of magnesium sulphate is indicated to treat eclamptic seizures and also in the prevention of seizures in women with severe pre-eclampsia. This treatment should considered in the context of pharmacological control of the woman’s blood pressure and provision of safe anaesthesia to allow delivery of the baby.

FURTHER READING