The role of magnesium in the emergency department

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Magnesium has been advocated for the treatment of a variety of conditions seen in emergency medicine. The authors present a systematic review and advice on appropriate indications for its use. Evidence supports its use in severe asthma, eclampsia, and torsade de pointes. There is insufficient evidence to justify its routine use in other emergencies.

Magnesium is the second most abundant intracellular cation and the fourth most abundant cation in the body. It is an essential transmembrane and intracellular modulator of cellular electrical activity. Possible therapeutic uses are broad but until recently, few studies had been undertaken. Numerous clinical trials and other studies are now available in a wide range of conditions. We examine the role of magnesium in the emergency department.

MYOCARDIAL INFARCTION

There has been considerable interest in magnesium as a treatment to limit myocardial damage in myocardial infarction (MI). Experimentally it has been shown to have a role in myocardial salvage, possibly by inhibiting calcium influx to ischaemic myocytes and/or by reducing coronary tone. It has also been shown to increase the threshold for depolarisation of cardiac myocytes, theoretically reducing the risk of malignant arrhythmia. In healthy humans it can reduce peripheral vascular resistance and increase cardiac output with no effect on cardiac work.1

Prior to 1995 a number of small studies and one large study had all produced positive outcomes for magnesium intervention in acute MI. The LIMIT-2 study, a randomised controlled trial (RCT) with 2316 subjects, demonstrated a statistically significant 16% reduction in all cause mortality for magnesium compared with placebo (95% CI 2% to 29%) mainly because of a reduction in early left ventricular failure.2

However, the ISIS-4 study3 with 58 050 subjects, which assessed the effects on mortality and major morbidity of the addition of intravenous magnesium to standard treatments for acute MI found no benefit. There was no effect on mortality at five weeks and subgroup analysis found no benefit in those patients not receiving thrombolysis or in those presenting within six hours of onset of symptoms. Slightly fewer patients had ventricular fibrillation but more had some other form of cardiac arrest. Overall no difference in risk was seen. In addition, there was a small but significant increase in heart failure, cardiogenic shock and deaths attributed to cardiogenic shock (1.62% v 1.26% 2p<0.001) though the cause of the excess was unclear.

Further small studies subsequent to ISIS-4 have continued to suggest a positive benefit from magnesium in acute MI.4 5 The most recent by Gymlani et al6 demonstrated a significant reduction in deaths from arrhythmia (4% v 20%) and from pump failure (4% v 14%). The investigators suggested the timing of the treatment was crucial. However, a placebo controlled RCT of intravenous magnesium in 150 patients undergoing primary percutaneous transluminal coronary angioplasty, demonstrated that infusion before, during, or after reperfusion resulted in no improvement in short-term clinical outcome.7 At present a further large RCT, the MAGIC trial8 is in progress to specifically address the question of the role of magnesium in the management of acute MI, focusing on early use.

In summary, despite the theoretical benefits magnesium therapy may offer in the management of acute MI there is no good evidence to support its use. The results of the MAGIC trial are awaited with interest.

CARDIAC FAILURE AND MALIGNANT ARRHYTHMIAS

Hypomagnesaemia is common in left ventricular and biventricular failure, secondary to increased magnesuria mediated by loop diuretics. Positive results in animal experiments led to suggestions that magnesium should form part of the acute treatment protocol for malignant arrhythmias in patients with significant ventricular dysfunction.9 10

Bashir et al11 performed a randomised, double blind crossover trial of long term oral magnesium supplementation in congestive heart failure. They demonstrated a reduction in non-sustained monomorphic ventricular tachycardia (mVT) of 24% (p<0.01). The study only contained 21 patients. Similar results were obtained by Sueta et al12 in patients with symptomatic heart failure treated acutely with magnesium or placebo. However, only 30 patients were involved, none had a history of symptomatic arrhythmias, and only asymptomatic non-sustained mVT was suppressed. Ceremuzynski et al13 demonstrated hypomagnesaemia in 38% and excessive urinary magnesium in 72% of 78 patients with congestive heart failure (ejection fraction <40%) on admission. They demonstrated that magnesium treatment reduced episodes of non-sustained mVT.

Abbreviations: MI, myocardial infarction; mVT, monomorphic ventricular tachycardia; AVN, atrioventricular nodal; AF, atrial fibrillation; TdP, torsade de pointes; EAD, early after depolarisation; HF, hydrofluoric acid
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significantly but again demonstrated no change in the risk of adverse events or death. The PROMISE study enrolled 1068 patients with New York Class III/IV heart failure for a RCT of milrinone. All patients had magnesium levels checked during the trial. Serum magnesium was shown not to be an independent risk factor for sudden death or all cause death.

In conclusion there is little evidence to support the routine use of magnesium in patients with ventricular dysfunction presenting with a malignant arrhythmia. Further studies are necessary in view of the biological plausibility of this intervention. Until they are completed it should be reserved for those with proven hypomagnesaemia.

**PRIMARY VENTRICULAR TACHYCARDIA**

There is some experimental evidence but no clinical evidence to support the use of magnesium in monomorphic ventricular tachycardia either as treatment or post-event prophylaxis.13

**CARDIAC ARREST**

Magnesium has been investigated as a first line drug in cardiac arrest in addition to the standard interventions as per Advanced Life Support (ALS) guidelines. No benefit was found in either out of hospital or in hospital cardiac arrest.15

**ATRIAL FIBRILLATION**

Magnesium has no effect on sinus node function but does delay atrioventricular nodal (AVN) conduction and prolong the atrial refractory period. Only one small trial on cardioversion of acute atrial fibrillation (AF) has shown any benefit but this included patients with AVN tachycardia and compared the use of magnesium with verapamil rather than placebo.16 Another small study demonstrated some benefit in ventricular rate control though this was no better than digoxin. This was disputed by Frick et al who demonstrated no effect on heart rate or rate variability in patients with chronic AF.17

Magnesium has been shown to act as an indirect antagonist of digoxin at the sarcolemma Na+K+ATPase pump and has shown benefit in reducing the incidence of ventricular arrhythmias associated with digoxin toxicity.18 The gold standard however remains Fab antibodies.

**TORSADE DE POINTES**

Torsade de points (TdP) is a form of polymorphic ventricular proarrhythmia associated with QT interval prolongation and the presence of prominent U waves on the resting electrocardiogram (ECG). These ECG changes represent prolonged repolarisation and the development of early after depolarisation (EAD). It may degenerate to ventricular fibrillation.

In myocardial cells repolarisation occurs when the efflux of positive ions (mainly potassium) exceeds the declining influx of sodium and calcium ions. In the long QT syndromes potassium ion channel dysfunction causes an intracellular surplus of positive charge, so delaying ventricular repolarisation. Delay of inactivation of calcium ion channels results in late inflow producing EADs. These may reach threshold amplitude so triggering a ventricular arrhythmia. As the deep subendocardium is most susceptible to prolonged repolarisation and the development of EAD the heterogeneous state of the myocar-dium results in a specific type of re-entrant arrhythmia. This is reflected in the TdP pattern on ECG.

The long QT syndrome with risk of development of TdP can be congenital but is usually iatrogenic. It is associated with:

- Antiarrhythmics especially class Ia and III
- Phenothiazines and butyrophenones
- Tricyclic antidepressants
- Non-sedative antihistamines
- Some antibiotics especially macrolides
- Antifungals
- Organophosphates
- Cocaine

It is also associated with starvation, bradycardias, and subarachnoid haemorrhage.20–21

The aims of treatment in TdP are (a) to shorten the QT interval and (b) to alter the after depolarisation effect. There have been no RCTs of TdP therapy. Present management is based on theoretical concepts supported by experimental evidence and the evidence of small clinical trials and case series.22–24

Magnesium, at a dose of 2 g (25–50 mg/kg in children) magnesium sulphate intravenously over one to two minutes, is used to suppress EADs in the emergency situation. In vitro studies have shown it reduces the amplitude of EADs to sub-threshold levels by blocking calcium influx. This should be accompanied by correction of hypokalaemia to a serum K+ concentration of >4.5 mmol/L.25 Lignocaine (lidocaine) indirectly suppresses the development of triggered potentials by shortening the action potential duration. However, this effect is inconsistent with a reported success rate of only 50%.26

Magnesium with/without potassium infusion should be accompanied by acceleration of the basic heart rate, as there is an inverse relation between rate and the repolarisation duration.23

In summary, magnesium is a safe and simple intervention in TdP. It should be the first line drug therapy in TdP followed by attempts to accelerate the heart rate.

**BRONCHIAL ASTHMA**

Magnesium acts as a smooth muscle relaxant by altering extracellular calcium influx and intracellular phosphorylation reactions. It may also attenuate the neutrophilic burst associated with inflammatory bronchoconstriction by attenuating mast cell degranulation. The principal trigger for this degranulation is a rise in intracellular calcium, which is antagonised by magnesium.27 It has been shown experimentally that degranulation is a cause of bronchial asthma. Magnesium acts to suppress EADs in the emergency situation.28–29

A number of small RCTs have examined intravenous magnesium use in bronchial asthma. Skobeleff et al reported significant improvements in peak expiratory flow rates and in numbers discharged from the emergency department in adults with severe asthma. This was supported by further studies in adults and children with moderate to severe exacerbations using 2 g in adults and 20–25 mg/kg in children.30–32

There was no evidence for its efficacy in unselected groups with asthma exacerbation.33 One study however showed no benefit in moderate and severe exacerbations in adults. A Cochrane systematic review in 1999 supported the use of intravenous magnesium sulphate in a subgroup of patients with severe acute asthma. Since their meta-analysis three further small RCTs of intravenous magnesium in acute severe childhood asthma have been published two positive and one negative.34

The possibility of using isotonic magnesium as a vehicle for nebulised β agonists has also been investigated with some positive results.35

At present, intravenous magnesium seems to be both safe and effective in acute severe asthma in all age groups. A large prospective RCT is necessary to confirm this. Until this is available magnesium should be used as a safe, easy to administer and effective second line agent in acute severe asthma. Nebulised magnesium remains experimental.

**ECLAMPSIA**

Eclampsia is the occurrence of one or more epileptiform fits in association with the syndrome of pre-eclampsia. Pre-eclampsia is a multisystem disorder of pregnancy usually
associated with hypertension and proteinuria. Eclampsia occurs in one in 2000 deliveries in the developed world and is a factor in 10% of maternal deaths in the UK.

Magnesium sulphate was first suggested as an anticonvulsant in 1906 and has been widely used in the United States since the 1930s. The suggested mechanism of action is cerebral vasodilatation and/or prevention of ischaemic neuronal damage by blockade of n-methyl-d-aspartate (NDMA) receptors.60

The Eclampsia Trial Collaborative Group organised a large double blind RCT with two arms comparing magnesium with phenytoin and diazepam.40 In the comparison of magnesium and diazepam (910 women) there were significantly fewer recurrent convulsions with magnesium. Treatment with magnesium resulted in 15 fewer women per 100 having further convulsions. There was no significant difference in maternal mortality between the treatments though the trend was favourable for magnesium therapy. There was no effect on the baby’s outcome. In the magnesium and phenytoin arm of the study (777 women) there were significantly fewer recurrent convulsions with magnesium. There was also a reduced risk of the need for anaesthesia, an 8% reduction in ITU admissions, a reduced risk of perinatal death and a reduced risk of neonatal special care for more than seven days. There was a favourable trend for overall maternal outcome with the mortality rate 5.1% in the diazepam group compared with 3% in the magnesium group though this failed to reach significant levels.

Two Cochrane Systematic Reviews produced by the Cochrane Pregnancy and Childbirth Group 41 42 support these findings.

Magnesium is the first line drug in eclampsia. It is substantially more effective than both phenytoin and diazepam. The most widely used regimen is a magnesium sulphate 4 g intravenous bolus followed by 1 g per hour intravenous infusion.

MIGRAINE

Interest in magnesium as a possible treatment in migraine stems from studies that demonstrate:

1. Ionised magnesium affects serotonin receptor activity.
2. An increased ionised calcium/ionised magnesium ratio promotes cerebral vasospasm.

A number of studies have suggested it may have a role in prophylaxis.40 Two small studies have also suggested a role in the treatment of acute migraine. Mauskop et al 44 demonstrated relief of headache within 15 minutes of intravenous magnesium in 32 of 40 patients with migraine, cluster headache, or tension headache. In 18 this was maintained for 24 hours and 16 of these had low serum magnesium levels. Demirkaya et al demonstrated relief of headache in 13 of 15 migraine patients and relief of associated symptoms in all patients compared to partial relief of headache in one patient in the placebo group.45

Further large prospective studies are necessary as this intervention in acute migraine is to date unproven though biologically plausible. It is potentially a safe and rapid intervention.

ALCOHOL WITHDRAWAL

Significant magnesium deficiency is a common, though not universal, finding in chronic alcoholism. This is attributable to alcohol induced urinary magnesium loss, malnutrition, gastrointestinal losses, and phosphate and vitamin D deficiency.46 A role for magnesium therapy in alcohol withdrawal was suggested by animal studies, which demonstrated that magnesium deficient mice developed ethanol withdrawal features without ethanol exposure. However, the only RCT of magnesium supplementation in the alcohol withdrawal syndrome in humans found no difference in symptoms or sedation requirement. A recent evidence based practice guideline from the American Society of Addiction Medicine on the pharmacological management of alcohol withdrawal states there is no evidence that magnesium supplementation reduces withdrawal severity or the frequency of delerium or seizures.46

Hydrofluoric acid burns

Hydrofluoric acid (HF) readily penetrates the skin and mucous membranes, causing deep tissue layer destruction. Exposure can produce hypocalcaemia, hypomagnesaemia, hyperkalaemia, cardiac arrhythmias, and death. Fluoride ion chelation is the treatment of choice as free fluoride ions are thought to be responsible for burn progression. Topical and/or parenteral calcium salts are the standard treatments for both dermal and systemic manifestations.46 However, magnesium is also an effective fluoride ion chelator. An animal study by Burkhart et al 44 compared topical preparations of magnesium and calcium in dermal exposure. Histological analysis of the burns demonstrated calcium treatment was associated with less severe and more superficial injury. However, further animal studies comparing intradermal calcium injection with intravenous magnesium suggest that magnesium therapy reduces the severity of HF burns.46

There have been no trials in humans and no comparisons of intravenous magnesium with intravenous or intra-arterial calcium. Magnesium in HF burns remains experimental unless treating proven systemic hypomagnesaemia.

ADVERSE EFFECTS

Magnesium has minimal side effects in usual therapeutic doses and has a large therapeutic index. At typical doses the main problems are transient facial warmth and flushing though transient hypotension may occur with over-rapid intravenous infusion. At a serum level of 12 mg/dl abnormal cardiac conduction, absent reflexes and muscle weakness with respiratory depression occur. Regular assessment of the patellar tendon reflex provides a quick and easy method of monitoring for developing toxicity.46

CONCLUSION

Magnesium has a clear role in the emergency management of a number of conditions.

• It should be used as first line therapy in eclampsia and torsade de pointes ventricular tachycardia.
• It has a clearly defined role as a second line therapy in acute severe bronchial asthma.
• Hypomagnesaemia should be considered in patients with biventricular failure presenting with malignant arrhythmias.
• Magnesium should be considered as a temporising measure in cases of severe digoxin toxicity while using Fab antibodies as the specific antidote.

Magnesium is safe and easy to use. In conclusion, magnesium should be available for immediate use in all emergency departments.

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REFERENCES

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