INTRODUCTION
Poisoning is a significant global public health problem. According to World Health Organisation data, an estimated 350,000 people died from unintentional poisoning in 2002. Accidental poisoning occurs most often in the age group 1-5 years, although less than one per cent of poisoning in children is serious. More than 94% of fatal poisonings occur in low- and middle-income countries. It remains a challenge to identify those at risk at an early stage.

Education of the general public and also the provision of child-proof containers for household chemicals and medicines play a key role in prevention.

(Toxic plants, insect and snake bites are not discussed in this article, but see www.rch.org.au/poisons for more information)

ACCIDENTAL POISONING IN CHILDREN: GENERAL PRINCIPLES
A history of poisoning usually makes the diagnosis obvious but poisoning should be considered in any child presenting with altered consciousness, cardiorespiratory depression, hypothermia, seizures or gastrointestinal symptoms. The possibility of more than one poison should always be considered.

1. Ascertain the nature and time of poisoning.
2. Contact the local Poison Centre for advice (if available). Sources on information are common and include internet-based information databases such as the International Programme on Chemical Safety (www.inchem.org), the UK-based TOXBASE (www.spib.axl.co.uk), or in the USA, TOXNET (toxnet.nlm.nih.gov).
3. Assess the patient following the principles of 'ABC' (Airway, Breathing, and Circulation). If airway protection is impaired, the patient requires intubation.
4. Induced emesis is no longer recommended and is contraindicated with volatile substances.
5. Consider gastric emptying and administration of activated charcoal. Gastric lavage is recommended if life-threatening poisoning has occurred within the previous hour, but contra-indicated if the airway cannot be protected. Gastric lavage is also contra-indicated in the ingestion of hydrocarbons (risk of aspiration and chemical pneumonitis) and corrosives. Multiple-dose activated charcoal (1g/kg every four hours) is used for substances with a long half life (anticonvulsants, digoxin, and theophylline). The complications of gastric lavage include aspiration pneumonia, hypoxia, mechanical injury to the gut and the induction of hypo-or hypernatraemia. It may also induce charcoal bezoars that may cause intestinal obstruction. Activated charcoal can cause serious complications if aspirated.
6. Remove contaminated clothing and wash patient with soap and water.
7. Send samples for lab investigation (urea, electrolytes, blood glucose.) Urine and gastric aspirates should be saved for later toxicology analysis (where available).
8. Measure relevant drug levels (paracetamol and salicylate and others if available).
9. ECG – important in cases of tricyclic antidepressants and unknown poisons. QRS prolongation is an early sign of cardiovascular involvement
10. Specific antidotes should be given as per instruction by Poison Centre.

SPECIFIC POISONS
Household chemicals
Children frequently ingest household substances and most of these are nontoxic. However, some products contain alcohol which may cause seizures and hypoxia.

Household solutions of bleach contain approximately 10% hypochlorite. They are rarely ingested as they are extremely unpalatable. Commonly, they cause nausea, vomiting and diarrhoea. Less than 100ml of household bleach is unlikely to cause serious problems. Fluids should be encouraged, particularly milk. Oesophageal
damage occurs rarely and is associated with concentrated solutions. There is a worrying recent increase in oesophageal injury associated with dishwasher tablets. Removal of these under anaesthesia is a medical emergency.

Ingestion of small quantities of strong alkalis such as drain cleaner containing sodium hydroxide may cause devastating injuries. Oesophageal injury is most common and evolves over the course of a few days. Ingestion causes immediate burning pain, swelling of the lips, and depending on the quantity ingested, salivation, haematemeses, dyspnoea, stridor or shock.

The burn injury is classified by endoscopy:

- mucosal (first degree burn)
  - usually resolves after 2-3 days
- submucosal (second degree burn)
  - requires total parenteral nutrition
- full thickness (third degree burn)
  - shocked, will require intensive care.

Third degree burns may lead to gastrointestinal perforation or airway complications or long term, severe oesophageal stricture causing absolute dysphagia which evolves over a period of several months.

Treatment is supportive; emesis is contraindicated and nasogastric aspiration unhelpful as the injury is oesophageal. Patients with first degree burns may be discharged after 2-3 days once tolerating oral fluids. Patients with third degree burns will be in shock and will require intensive care: tracheostomy is indicated for patients with evidence of supraglottic/epiglottic burns. Steroids may reduce the risk of stricture formation if given within 12-24 hours but their use is controversial. Antibiotics are indicated for perforation. Long term gastrostomy or oesophageal replacement surgery (gastric pull-up) may be required for oesophageal stricture.

**Hydrocarbons (paraffin, white spirit)**

Most of the hydrocarbons are petroleum distillates, containing a variable amount of saturated and unsaturated aliphatic (open-chain) and aromatic (cyclic) hydrocarbons. The aliphatics are not readily absorbed from the gastrointestinal tract and therefore cause minimal systemic toxicity. Paraffin, petrol, thinner, diesel and benzene are low viscosity aliphatic-based petroleum distillates. The aromatic hydrocarbons are well absorbed, and therefore may cause systemic toxic effects but are less inclined to aspiration-related complications. The main hazard of accidental ingestion of the aliphatic hydrocarbons (paraffin) is that of chemical pneumonitis characterised by ventilation/perfusion imbalance and hypoxia. The aspirated hydrocarbons inhibit surfactant and also cause direct broncho-alveolar injury. This can occur even in the absence of vomiting or impaired consciousness and as little as 1ml aspirated hydrocarbon can result in a chemical pneumonitis.

Paraffin is often sold or stored in unlabeled containers which may be within children's reach. Volumes ingested tend to be small because of a burning sensation in the mouth and the bad taste of the liquid. Nausea and vomiting are common. Aspiration pneumonitis occurs in 12-40% of patients. Signs and symptoms appear 30 minutes after ingestion but may be delayed for 8 hours. A non-productive cough, tachypnoea and tachycardia are signs of a developing chemical pneumonitis. There may be wheeze, coarse crackles as well as signs of respiratory distress (intercostals and subcostal retraction). Hypoxia and cyanosis may be present in severe cases. Patients may be lethargic or irritable. A fever may also be present. The clinical picture generally deteriorates over the first 24 hours. A temperature that persists or only develops after 24 hours usually suggests a secondary infection.

Abnormalities on the chest film may be seen within 30 minutes after aspiration even in the absence of clinical signs or symptoms. Common radiological findings of chemical pneumonitis include bilateral perihilar infiltrates which progress to form patchy infiltrates and later become large areas of consolidation.

**Management of hydrocarbon poisoning**

- ABC principles.
- Observe all asymptomatic patients for at least 8 hours. Carefully examine patients for the development of respiratory signs. A chest film at 6 hours is recommended.
- If asymptomatic after 8 hours and the chest film is normal, the patient may be discharged.
- Symptomatic patients should be Xrayed on admission.
- Emesis and gastric lavage is contra-indicated as it increases the risk of aspiration. The use of milk is not recommended, but clear fluids are not contra-indicated.
- Oxygen should be administered to all patients with respiratory signs or symptoms. Other respiratory support modalities should be instigated as required, according to standard indications.
- Corticosteroid therapy may increase the risk of secondary bacterial infection and has not been shown to be of any benefit.
- Antibiotics should be given if there are suggestions of a secondary infection (persistent fever or a fever that develops after 24 hours.)
- Prophylactic antibiotics have not been shown to prevent secondary infection and are not advocated.

**PESTICIDES**

**Paraquat**

Paraquat ingestion remains a problem in a large number of countries. Most of these cases are intentional suicide attempts (73% in a Malaysian study) or occupational exposure and therefore not that frequently seen in the paediatric population. It is the most toxic herbicide known, producing multi-organ failure. After oral ingestion, patients develop a severe gastroenteritis with oral, oesophageal and gastric ulceration. Depending on the dose ingested, multi-organ failure may develop within 48-72 hours. As little as 10ml may be fatal. Those who survive the initial phase develop pulmonary fibrosis
as a response to the acute alveolitis in the first phase. This leads to respiratory failure and patients die of anoxia.

**Management of paraquat poisoning**

- Any history of oral exposure should be considered potentially fatal. Treatment strategies are aimed at prevention of absorption, enhancement of elimination and prevention of pulmonary damage.
- Treat immediately with Fuller’s earth (Fuller’s earth 300g and magnesium sulphate 50g in a litre of water). Give 5–10ml.kg⁻¹ orally every 2–3 hours for 2–3 days.
- Activated charcoal may be used if Fuller’s earth is not available. Mix with saline and a laxative such as lactulose to prevent constipation. Repeat every 4 hours until paraquat is no longer detected in the urine.
- If neither is available, careful gastric emptying may be done but be aware of possible pharyngeal and oesophageal ulceration and perforation.
- Haemoperfusion or dialysis may be used (if available).
- Appropriate management of respiratory complications includes PEEP/CPAP. Low FiO₂ should be used as high concentrations of oxygen lead to worse pulmonary toxicity. Paraquat accumulates in the lung where it generates superoxide anions through the reaction with oxygen.

**Organophosphate poisoning**

Organophosphate poisoning (OP) remains an issue in developing countries. In one study, 35% of OP poisoned victims were children. The organophosphates and carbamates are cholinesterase inhibitors, thereby eliciting cholinergic signs and symptoms. They can be absorbed by ingestion, inhalation or via the skin. The organophosphates form an irreversible complex with cholinesterase but the carbamyl-enzyme complex is reversible. Thus a less severe clinical picture of shorter duration is presented. It also causes fewer CNS effects because of less penetration of the blood brain barrier.

OP poisoning causes: SLUDGE; Salivation, Urination, Lacrimation, Defication, Gastric, Emesis and may progress quickly to CNS symptoms with seizures, especially in children.

- Muscarinic effects – hypersecretion, vomiting, diarrhoea, constricted pupils, bronchoconstriction and urinary incontinence
- Nicotinic effects – muscular weakness, fasciculations (rarer in children) and respiratory muscle weakness (may override muscarinic effect and cause tachycardia, hypertension and mydriasis)
- CNS effects – irritability, seizures, coma (and accompanying respiratory depression).

**Management of organophosphate poisoning**

- ABC
- Patient may require intubation (avoid suxamethonium because of prolonged effect)
- Gastric lavage, activated charcoal
- Remove all contaminated clothes (take care not to get exposed yourself)
- Wash patient with soap and water (mild alkaline soap helps deactivate organophosphates)
- Start intravenous atropine administration as soon as possible. Initial test dose of 0.01mg.kg⁻¹. Then 0.05mg.kg⁻¹ every 15 minutes until full atropinisaton is reached. Maintenance therapy is usually around 0.05mg.kg⁻¹.h⁻¹. Adequate therapy is achieved when there is control of excessive bronchial and oral secretions. Don’t worry about the tachycardia – it is well tolerated. Eye signs aren’t a good marked of progress. When the patient improves, the dose should be tapered slowly over 24 hours. Atropine should never be stopped abruptly. Close observation is required as rebound organophosphate toxicity may occur due to their lipid solubility.
- Watch out for chemical aspiration pneumonia as a lot of the organophosphate pesticides have a petroleum base.

**INHALED POISONS**

**Carbon monoxide**

Carbon monoxide is produced through incomplete combustion processes. It is colourless, tasteless and non-irritant. The commonest sources of carbon monoxide poisoning are smoke inhalation, poorly maintained domestic gas appliances, and deliberate inhalation of car exhaust fumes (the latter less common in children). Carbon monoxide causes tissue hypoxia by the interruption of electron transport in the mitochondria. It also reduces oxygen delivery by competing with O₂ for binding to Hb and altering the shape of the HbO₂ dissociation curve (making the curve less sigmoidal, a shift to the left). Its affinity for Hb is >200-fold that of O₂.

Carbon monoxide causes injury by hypoxia, with symptoms referable to tissues with greatest oxygen consumption, notably the heart and brain. Patients present with hypoxia without cyanosis. Skin and mucosal surfaces may appear cherry red. Symptoms correlate roughly with the COHb level:

<table>
<thead>
<tr>
<th>Carboxyhaemoglobin (COHb) level</th>
<th>Symptoms</th>
</tr>
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<tbody>
<tr>
<td>&lt;30%</td>
<td>dizziness and headache</td>
</tr>
<tr>
<td>50-60%</td>
<td>syncope, tachypnoea, tachycardia and fits</td>
</tr>
<tr>
<td>&gt;60%</td>
<td>increased risk of cardiorespiratory failure and death</td>
</tr>
</tbody>
</table>

Anaemia, increased metabolic rate (e.g. children) and underlying ischaemic heart disease all increase susceptibility to CO. Neurological recovery depends on the duration of hypoxic coma.
Management of carbon monoxide poisoning

- Pulse saturation monitoring does not distinguish between HbO₂ and COHb and will read falsely high.
- Arterial blood gases should be done if available. PaO₂ may be normal but any evidence of metabolic acidosis indicates serious poisoning (useful even in the absence of COHb measuring facilities.) COHb measurements are diagnostic but not always available.
- Treat according to ABC principles.
- Apply tight-fitting mask with 100% oxygen. This reduces the half life of COHb from 320 mins (in room air) to 80 minutes. Intubation and ventilation may be necessary in severe cases.
- Hyperbaric oxygen is useful to further reduce the COHb half life, but access and transfer times to a hyperbaric chamber make it an impractical option.
- Check 12 lead ECG – if there are signs of myocardial ischaemia, the patient needs intubation/ventilation.
- Supportive care should include continuous cardiac monitoring, treatment of arrhythmias and correction of acid/base and electrolyte abnormalities.
- Lab bloods should be done for creatinine and electrolytes, FBC, creatine kinase and cardiac enzymes.
- Brain hypoxia leads to cerebral oedema and fits. Diazepam 5-10mg iv is used to control the fits.
- Initial chest Xray may be normal even in severe smoke inhalation.

Patients need to be followed up as some of the neuropsychiatric complications may take weeks to develop.

Combustion of household materials can also generate a number of other toxic substances such as sulphur dioxide, nitrogen dioxide, acrolein, cyanide and various acids from PVC and polyurethane. This may cause direct lung, skin and conjunctival injury.

Cyanide poisoning

Most commonly seen in victims of smoke inhalation as a combustion product of polyurethane foams. Cyanide derivates are also used in industrial processes and fertilizers. Children may ingest amygdalin, a cyanogenic glycoside contained in kernels of almonds and cherries. Cyanide acts by irreversibly blocking mitochondrial electron transport.

HCN gas can lead to cardiorespiratory collapse and arrest within a few minutes. Patients surviving to reach hospital are unlikely to have suffered significant poisoning. Early signs include dizziness, chest tightness, dyspnoea, confusion and paralysis, followed by cardiovascular collapse, apnoea and seizures.

Management of cyanide poisoning

- Do not attempt mouth to mouth resuscitation as the skin will be contaminated.
- Apply 100% O₂ with a tight fitting mask. Treatment is supportive.
- Decontaminate the skin with soap and water.
- Gastric lavage could be attempted where there are no signs of cyanide poisoning.
- Dicobalt ededate, sodium thiosulphate and sodium nitrate are used as antidotes.

MEDICINES

Children frequently ingest medicines. Dangerous substances for children include salicylates, paracetamol, iron, theophylline and tricyclic antidepressants.

Aspirin / salicylate poisoning

Commonly ingested by children. Also probably the commonest drug to be ingested deliberately in overdose. Oil of wintergreen is 98% methyl salicylate. Its primary toxic effect is by uncoupling of oxidative phosphorylation.

Patients present with restlessness, hyperventilation, tinnitus, deafness, tachycardia, nausea, vomiting, sweating, hyperthermia and dehydration. Pulmonary oedema, acute renal failure, hypokalaemia, hypoglycaemia and hypothermia may also develop. In adults there is an early increase in respiration rate causing a respiratory alkalosis that precedes the later development of metabolic acidosis. However this is not seen in children.

Management of aspirin / salicylate poisoning

- Gastric lavage should be attempted, even up to 12 hours or longer post ingestion. In overdose, salicylates may form concretions in the stomach and delay absorption.
- Activated charcoal orally every four hours.
- Severity of the poisoning can be determined by measuring blood salicylate levels 6 or more hours after ingestion but may be misleading in severe acidosis. Ideally should be done on admission and 4 hours later to assess continued absorption. Therapeutic levels of salicylate are generally less than 300mg.l⁻¹. Levels more than 750mg.l⁻¹ represents severe poisoning.
- Check electrolytes. Blood glucose monitoring should be done every 2 hours.
- Dehydration, acidosis, hypoglycaemia and electrolyte disturbances should be corrected. Pay particular attention to potassium. Care should be taken to avoid fluid overload, and renal function should be closely monitored.
- Alkalisation of the urine (pH 7.5-8.5) by administering sodium bicarbonate (either orally or by infusion) is recommended to increase excretion of salicylates.
- Haemodialysis or charcoal haemoperfusion may be required in severe poisoning (levels > 1000mg.l⁻¹ or 7.25mmol.l⁻¹) or in patients with decreased urine output, pulmonary oedema or progressive deterioration.
Paracetamol overdose
Paracetamol ingestion is common but seldom leads to severe toxicity in children due to the diluted concentration of the paediatric syrup formulation. The children at risk are those with glutathione depletion, such as children with cystic fibrosis, adolescents with eating disorders and those also on enzyme inducing agents such as anticonvulsants. The liver is the main target organ in paracetamol poisoning. Patients are generally asymptomatic up to 24 hours post ingestion. Mild nausea, vomiting and anorexia may occur. Hepatic necrosis becomes apparent in 24-36 hours with right subchondral pain and tenderness, jaundice, vomiting and acute liver failure. Confusion and encephalopathy develop over 36-72 hours. Oliguria and renal failure may occur from acute tubular necrosis in the absence of liver failure. Lactic acidosis may be seen early or late.

Management of paracetamol poisoning
• Patients presenting within 4 hours of ingestion should undergo gastric lavage.
• Activated charcoal should not be given orally if the specific antidote is also given orally.
• Measure paracetamol levels at least 4 hours post ingestion andplot on ‘plasma paracetamol concentration – time graph’ to determine treatment line.
  - If the initial levels indicate no treatment, repeat after 4 hours to check for delayed absorption.
  - All patients on or above the ‘Normal treatment line’ should be given N-acetylcysteine.
  - Patients on enzyme-inducing drugs should be treated if above the ‘Enhanced risk treatment line’.
• Where levels are not available, assume liver damage after ingestion of a single dose of more than 150mg.kg⁻¹ paracetamol, or a staggered overdose (spread over several hours) of the same amount.
• Acetylcysteine is the antidote of choice and is usually given iv. Oral methionine may be used if the patient is allergic to acetylcysteine and is also a suitable alternative in remote areas if vomiting is not a problem. Oral acetylcysteine and carbocysteine have also been used.
• Monitor urea and electrolytes, PT and LFT.
• Give vitamin K but avoid giving fresh frozen plasma unless there is active bleeding. The PT is the best indicator of the severity of liver failure.
• All patients with encephalopathy or a rapidly rising PT should be referred to a liver unit (where available).
• Supportive treatment and treatment of liver and renal failure as indicated.

Iron poisoning
Ingestion of more than 20mg.kg⁻¹ of elemental iron is considered potentially toxic, and the lethal dose is estimated at about 180mg. kg⁻¹ Iron is extremely irritant. The clinical features can be divided into three phases (not always very clear). Patients present with vomiting, diarrhoea, abdominal pain, haematemesis and rectal bleeding in the early phase (0-2 hours). This is followed by a period of stabilisation (up to 12 hours) during which a deceptive recovery occurs. This is followed by a life-threatening period during which coma, fits, jaundice, hepatic failure renal failure, clotting abnormalities, hypoglycaemia and cardiovascular collapse may occur. Patients alive 72 hours after ingestion usually make a full recovery. Late complications of gut stricture, gastric fibrosis and pyloric obstruction have been reported.

Management of iron overdose
• The stomach should be emptied with gastric lavage.
• Blood samples for FBC, glucose, UE, iron levels and ABG where available.
• Serum iron only peaks four hours after ingestion.
• An abdominal X-ray helps to determine the number of tablets ingested and also the success or failure of gastrointestinal evacuation.
• A patient with serum iron >90mmol.l⁻¹ may be treated with a chelating agent.
• If <20mg.kg⁻¹ are ingested, treatment is supportive. Patients with iron level below 54mmol.l⁻¹ and who remains asymptomatic 6 hours post ingestion would not be expected to develop significant toxicity and require no active treatment.
• For ingestion of between 20-60mg.kg⁻¹ elemental iron, gastric emptying may be considered if within one hour of ingestion. Whole bowel irrigation may be used in patients with ingestion of more than 60mg.kg⁻¹ and more than one hour post ingestion. It is especially useful if a slow release preparation has been ingested. Charcoal is of no benefit as iron does not bind to it.
• Desferrioxamine chelates iron and is the recommended treatment.
  - Give 1 gram im every 6-12 hours for children (2g for adolescents). (100mg of desferrioxamine binds 8.5mg of elemental iron)
  - If the patient is hypotensive, give desferrioxamine iv at a rate of 15mg.kg⁻¹.hr⁻¹, until the serum iron falls (maximum daily dose 80mg.kg⁻¹)
• Haemodialysis is indicated in very high serum iron levels that respond poorly to chelation therapy or if the urine output is not maintained. (the iron-chelate is excreted entirely in the urine)

Theophylline poisoning
This type of poisoning is rare but serious. Most preparations are slow release so that problems develop 12-24 hours after ingestion. Features of acute ingestion reflect the local irritant GI effects – nausea, vomiting, haematemesis and diarrhoea. The child may be hyperactive with dilated pupils, hyperreflexia, hypotonia and myoclonus. There may also be severe hypokalaemia, arrhythmias, metabolic acidosis, hyperglycaemia, hypotension and seizures.
Management of theophylline poisoning

- Gastric lavage should be considered if ingestion occurred within 1-2 hours.
- Repeated administration of activated charcoal to prevent further absorption and enhance systemic clearance. (May be difficult in the presence of nausea and vomiting).
- Whole bowel irrigation is considered if a slow-release preparation is ingested.
- Haemoperfusion should also be considered.
- Seizures are treated with diazepam.
- Assess hydration and correct hypokalaemia and electrolyte disturbances.
- Cardiac monitoring to detect arrhythmias.
- Verapamil and propanolol may be required to treat supraventricular and ventricular tachyarrhythmias. Lignocaine appears to have little effect on ventricular ectopy.

Tricyclic antidepressants

Cardiovascular toxicity is the main cause of death in tricyclic antidepressant overdose. It is caused by the blockade of noradrenaline uptake as well as the anticholinergic, membrane-stabilising and alpha-blocking effects. Nervous system toxicity includes drowsiness, agitation, hallucinations, hyperreflexia, myoclonus, rigidity, convulsions, respiratory depression and coma. Anticholinergic effects include flushing, dry mouth, dilated pupils, hyperpyrexia and bladder/bowel paralysis. Cardiovascular toxicity includes sinus tachycardia, hypotension, conduction abnormalities and arrhythmias. Respiratory complications include respiratory depression, aspiration pneumonia, ARDS and pulmonary oedema.

Management of tricyclic overdose

- Gastric emptying is delayed by tricyclic antidepressants, and therefore gastric lavage should be attempted as late as 12 hours post ingestion.
- Due to CNS depression and inadequate ventilation, intubation may be required.
- Continue cardiac monitoring for up to 48 hours.
- Inotropic support should be given as necessary.
- Correct metabolic acidosis to enhance protein binding of the drug.
- Control seizures with diazepam.

SUMMARY

Prevention of childhood poisoning is vital. There must be adequate supervision, safe placement of medications, child safe cabinets and containers, blister packaging and education. Most paediatric cases are not severe. Recognition of potentially life-threatening ingestions is important so that appropriate early treatment can be instituted. Aggressive supportive care is vital.


USEFUL WEBSITES

- World Health Organisation – The International Programme on Chemical Safety : Poisoning Prevention and Management (www.inchem.org)
- TOXBASE.org www.spib.axl.co.uk (NHS only, requires registration)
- Agency for toxic substances & disease registry: (no password or registration required) http://www.atsdr.cdc.gov/substances/toxsubstance.asp?toxid=19
- TOXNET (toxnet.nlm.nih.gov)
- Royal Children’s Hospital Melbourne: Victorian Poisons Information Centre (www.rch.org.au/poisons)