The Toxicology and Toxinology guidelines have been extensively revised and expanded. The updated treatment recommendations are based on available evidence and the consensus of our expert group. The guidelines are targeted towards hospital doctors, especially emergency department staff, as well as primary care providers (first responders, general practitioners, rural and remote practitioners) and poisons information specialists.

### What is covered in Toxicology and Toxinology?

The guidelines provide practical advice on managing a wide range of poisonings, exposures to toxic agents and envenomings. Toxicology and Toxinology now comprises:

- **14 topics on the general approach to managing poisoning**
- **7 topics on specific toxidromes (toxic syndromes)**
- **117 monographs about individual drug poisonings, envenomings and exposures to other toxic agents.**

Dozens of new monographs have been added to encompass a larger number of rare but highly toxic drugs and other agents (e.g., *Amanita phalloides* mushroom poisoning, button battery ingestion, specific herbicide poisonings including Paraquat, Glyphosate and Chlorophenoxy herbicides).

Other new poisoning monographs include Apixaban and rivaroxaban, Dabigatran, Buprenorphine, Cocaine, Insecticides, Isoniazid, Local anaesthetics, and Monoamine oxidase inhibitors. Some topics have been split to address the different treatments for acute poisoning versus chronic accumulation (e.g., Digoxin, Lithium).

Some drug classes include both a general monograph for the class, and specific monographs for individual drugs with specific issues in poisoning. For example:

- **Antidiabetic drug poisoning (general), plus specific monographs for Insulin, Metformin and Sulfonylurea poisonings**
- **Antiepileptic drug poisoning (general), plus specific monographs for Barbiturates, Benzodiazepines, Carbamazepine and oxcarbazepine, Lamotrigine, Phenytoin, Pregabalin and gabapentin, Sodium valproate, Tiagabine and Topiramate poisonings**
- **Antipsychotic drug poisoning (general), plus specific monographs for Amsulpride, Clozapine, Olanzapine and Quetiapine poisonings**
- **Hydrocarbon poisoning (general), plus specific monographs for Ingestion, Inhalation and Injection of hydrocarbons**
- **Novel psychoactive drug poisoning (introduction), plus specific monographs for Novel stimulant, Novel hallucinogen and Synthetic cannabinoid receptor agonist poisonings**
- **Poisoning from plants (introduction), plus eight specific monographs for various plant toxins.**

### How do I use Toxicology and Toxinology?

The guidelines include content on the general approach to poisonings, and specific toxidromes (e.g., Cholinergic toxidrome, Serotonergic toxidrome), followed by the grouped and individual poisoning monographs, listed alphabetically.

Most of the poisoning monographs have the following consistent structure for ease of navigation:

- **Management overview**—drug(s)/agent(s) and preparations available; salient points about toxicity and urgency of management; key management priorities and pitfalls
- **Risk assessment**—toxic dose and toxic concentration (if available); clinical presentation; key investigations
- **Treatment**—resuscitation, including interventions for airway, breathing and circulation; decontamination and enhanced elimination techniques; specific treatments (e.g., antidotes, sedation, seizure control)
- **Observation and patient disposition**—optimal duration for observation; advice for admission to hospital; special considerations for discharge.

Throughout the guidelines, advice on risk assessment and treatment in specific monographs often links back to the general approach and toxidrome topics. Common interventions in the management of poisonings that are covered in the general topics include:

- **Inotropic support**: first-line therapy, including adrenaline (epinephrine) and noradrenaline (norepinephrine)
- **Inotropic support**: high-dose insulin euglycaemia therapy (HIET), which may be recommended for second-line inotropic therapy
- **Cooling for drug-associated hyperthermia**, depending on availability, practicality and expertise
- **Decontamination for poisonings**, including detailed discussion on the risks of harm versus benefit of gastrointestinal decontamination with activated charcoal
- **Sedation in poisoning**, which may require higher than usual doses
- **Seizure control in poisoning**, which may require antiepileptic drugs other than benzodiazepines or phenytoin
- **Multiple-dose activated charcoal or extracorporeal elimination techniques** (including haemodialysis).

Flowcharts to aid in clinical decision-making have been updated based on the most recent evidence. Flowcharts available as printable figures include:

- **Management flowchart for a single ingestion of 10 grams (or 200 mg/kg in patients under 50 kg) or more of immediate-release paracetamol**
- **Management flowchart for modified-release paracetamol poisoning**
- **Summary of the acute management of suspected snake bite in Australia.**

### Is Toxicology and Toxinology evidence based?

Many of the poisonings, envenomings and other exposures covered in the guideline are uncommon, and evidence in this field is often limited. The guidelines are based on the available evidence supported by the consensus of the expert group. Every monograph includes evidence-based treatment recommendations, including children’s doses (if available) and maximum doses.

Specific recommendations for pregnant or breastfeeding women are not included because management for poisoning in this group is no different to the general population.
What practice-changing updates should I be aware of in Toxicology and Toxinology?

- **Paracetamol poisoning**
  It is now recommended to measure alanine aminotransferase (ALT) concentrations in addition to serum paracetamol concentrations in all patients, at least 4 hours after ingestion.

  Changes to recommendations for acetylcysteine as an antidote for paracetamol poisoning include:
  - a standard two-bag (20-hour) acetylcysteine protocol to replace the former three-bag protocol
  - varying acetylcysteine protocols for poisonings due to immediate-release, liquid, and modified-release paracetamol preparations
  - recommendation of high-dose (double-dose) acetylcysteine in the second infusion for high risk poisonings and for serum paracetamol concentrations that are more than double the paracetamol treatment nomogram line
  - addition of specific treatment for unintentional paracetamol poisonings (also known as repeated supratherapeutic ingestion [RSTI])
  - recommendation of extended acetylcysteine therapy for patients with evidence of hepatotoxicity.

- **Acetylcysteine therapy for hepatoprotection in other poisonings**
  Acetylcysteine is recommended for hepatoprotection in a standard two-bag protocol for:
  - *Amanita phalloides* mushroom poisoning
  - arsenic poisoning
  - paraquat poisoning
  - some hydrocarbon poisonings, including benzene, carbon tetrachloride and chloroform
  - some essential oil poisonings, including clove oil and pennyroyal oil.

- **Snake bite**
  The critical steps in the management of snake bite in Australia are:
  - determine if the patient has evidence of systemic envenoming
  - if so, determine if antivenom is indicated
  - if so, determine the snake group(s) likely to be responsible for the bite, to guide the choice of antivenom(s).

  These decisions are supported by a new flowchart of management of suspected snake bite in Australia.

  In most places, treatment of snake bite comprises combinations of single-dose monovalent antivenoms (eg tiger snake plus brown snake antivenom). Testing with a venom detection kit is no longer recommended.

- **Red-back spider bite**
  Red-back spider antivenom is not recommended. Analgesia is the mainstay of therapy.

- **Major box jellyfish (*Chironex fleckeri* species) sting**
  Intramuscular administration of box jellyfish antivenom is not recommended. Intravenous administration in hospital is recommended.

- **Button battery ingestion**
  If there is any suggestion of a button battery ingestion, a time-critical X-ray of the neck, chest and abdomen is urgently recommended, as well as referral for endoscopic removal if a battery is seen.

- **Digoxin poisoning**
  Titrated digoxin-specific immune antibody fragments (digoxin immune Fab) is now recommended for acute digoxin poisoning with life-threatening hyperkalaemia or arrhythmia, unless the patient is in cardiac arrest. Lower doses, also titrated, are used for the same complications in patients with chronic digoxin accumulation.

- **Colchicine poisoning**
  Updated treatment recommendations rely on the dose of colchicine ingested to guide interventions, including gastrointestinal decontamination and enhanced elimination using activated charcoal. Patients at higher risk of developing colchicine toxicity have a lower dose threshold for these interventions.

- **Gastrointestinal decontamination**
  Activated charcoal for gastrointestinal decontamination can be used to treat significant overdoses (after assessing the risk of harm versus benefit):
  - up to 2 hours after ingestion of immediate-release preparations
  - 4 hours or more after ingestion of modified-release preparations.

  Whole bowel irrigation can be considered:
  - up to 4 hours after ingestion of significantly toxic drugs (eg metals, modified-release preparations)
  - for ‘body packers’ and ‘body stuffers’ (concealed illicit drugs).

- **Maximum doses**
  Specific maximum total doses have been added for some drugs; for example:
  - the maximum total dose of intravenous sodium bicarbonate 8.4% for serum alkalinisation is 6 mmol/kg
  - the maximum total dose of calcium gluconate 10% for cardiac stabilisation in hyperkalaemia is 60 mL.

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