Updated guidelines for the management of paracetamol poisoning in Australia and New Zealand

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Paracetamol poisoning is the commonest cause of severe acute liver injury in Western countries.1,2 It is also the most common reason for calls to Poisons Information Centres in Australia and New Zealand.3 Not only is it one of the commonest medications involved in deliberate self-poisoning, it is also involved in a large proportion of accidental paediatric exposures and in overdoses with therapeutic intent when taken for symptoms such as pain or fever (repeated supratherapeutic ingestions). Since the publication of the previous guidelines in the Medical Journal of Australia in 2015, further research has emerged, particularly regarding acetyl cysteine regimens, massive paracetamol ingestions, and modified release paracetamol ingestion. These have led to a change in management of paracetamol poisoning, and the 2015 guidelines do not reflect the current practice recommended by clinical toxicologists. The key changes from the previous guidelines are acetyl cysteine regimen (two-bag regimen) and dosage, management of patients taking large or massive overdoses, staggered ingestions, modified release paracetamol ingestions and repeated supratherapeutic ingestions. The full guidelines are available online in the Supporting Information.

Methods

The Treatment of Paracetamol Poisoning Writing Group was comprised of clinical toxicologists and pharmacologists from Australia and New Zealand. All members completed a detailed literature review and critically appraised existing evidence, including reviewing the relevant chapters from the newly updated Australian Therapeutic Guidelines — Toxicology and toxicology.4 Drafts of evidence-based recommendations, practice points and a background manuscript were developed. We conducted a face-to-face meeting in May 2019 to draft the guideline. Further revisions were made via email and teleconference. The summary recommendations follow the National Health and Medical Research Council levels of evidence (https://www.mja.com.au/sites/default/files/NHMRC.levels.of.evidence.2008-09.pdf) and the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system (www.gradeworkinggroup.org) to determine the strength of the recommendations.

Recommendations

Acute deliberate self-poisoning, accidental paediatric exposure and inadvertent repeated supratherapeutic ingestions all require specific approaches to risk assessment and management. The initial approach focuses on risk assessment (Box 1). Key factors to consider for paracetamol poisoning are the formulation and dose ingested, time since ingestion, and serum paracetamol concentration (early), or clinical and laboratory features suggesting acute liver injury (late). Serum paracetamol concentration should be used to assess the need for acetyl cysteine administration in all patients presenting with deliberate self-poisoning with paracetamol, regardless of the stated dose. The paracetamol treatment nomogram (Box 2) can only be used in acute immediate release paracetamol ingestions with a known time of ingestion.

We have summarised with flow charts the management of acute immediate release paracetamol ingestion (Box 3), acute modified release paracetamol ingestion (Box 4), repeated supratherapeutic ingestion (Box 5), and a management flow chart for rural and remote centres with limited pathology facilities (Box 6).

Acetylcysteine infusions

Acetylcysteine should be administered as a two-bag regimen (Box 7) — this has changed from previous guidelines. The standard three-bag intravenous weight-based dosage regimen (150 mg/kg body weight over 15–60 min, then 50 mg/kg over 2 h) has similar efficacy but significantly reduced adverse reactions compared with the previous three-bag regimen. Massive paracetamol overdoses that result in high paracetamol concentrations more than double the nomogram line should be managed with an increased dose of acetylcysteine. All potentially toxic modified release paracetamol ingestions (≥ 10 g or ≥ 200 mg/kg, whichever is less) should receive a full course of acetylcysteine. Patients ingesting ≥ 30 g or ≥ 500 mg/kg should receive increased doses of acetylcysteine.

Abstract

Introduction: Paracetamol is a common agent taken in deliberate self-poisoning and in accidental overdose in adults and children. Paracetamol poisoning is the commonest cause of severe acute liver injury. Since the publication of the previous guidelines in 2015, several studies have changed practice. A working group of experts in the area, with representation from all Poisons Information Centres of Australia and New Zealand, were brought together to produce an updated evidence-based guidance.

Main recommendations (unchanged from previous guidelines):

1. The optimal management of most patients with paracetamol overdose is usually straightforward. Patients who present early should be given activated charcoal. Patients at risk of hepatotoxicity should receive intravenous acetylcysteine.
2. The paracetamol nomogram is used to assess the need for treatment in acute immediate release paracetamol ingestions with a known time of ingestion.
3. Cases that require different management include modified release paracetamol overdoses, large or massive overdoses, accidental liquid ingestion in children, and repeated supratherapeutic ingestions.

Major changes in management in the guidelines:

1. The new guidelines recommend a two-bag acetylcysteine infusion regimen (200 mg/kg over 4 h, then 100 mg/kg over 16 h). This has similar efficacy but significantly reduced adverse reactions compared with the previous three-bag regimen.
2. Massive paracetamol overdoses that result in high paracetamol concentrations more than double the nomogram line should be managed with an increased dose of acetylcysteine.
3. All potentially toxic modified release paracetamol ingestions (≥ 10 g or ≥ 200 mg/kg, whichever is less) should receive a full course of acetylcysteine. Patients ingesting ≥ 30 g or ≥ 500 mg/kg should receive increased doses of acetylcysteine.

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Immediate release paracetamol ingestion

The management of acute immediate release paracetamol ingestion — defined as any intentional or deliberate self-poisoning — is summarised in Box 3.

Recommendations on gastric decontamination have not changed since 2015. Fifty grams of activated charcoal should be administered to a cooperative, awake adult if they present within 2 hours of ingestion of a toxic dose (Box 1) of immediate release paracetamol, or within 4 hours of immediate release paracetamol overdoses greater than 30 g.13–15 GRADE: Strong; Evidence: Low.

The paracetamol treatment nomogram has been validated as an excellent predictor of risk but only for acute ingestions of immediate release paracetamol with a known time of ingestion. The current nomogram used in Australia and New Zealand has not changed (Box 3), except the units on the left and right axis have now been swapped. It is important to check the units used, with many laboratories recently changing from μmol/L (right axis) to mg/L (left axis). GRADE: Strong; Evidence: Strong.

Patients with a high initial paracetamol concentration (greater than double the nomogram line) are at increased risk of acute liver injury if given standard acetylcysteine regimens.13,16,17 Only a small percentage of paracetamol overdoses will have a paracetamol concentration greater than double the nomogram line and they typically have ingested 30 g or more of paracetamol. Those with an initial paracetamol concentration greater than double the nomogram line may benefit from an increased dose of acetylcysteine. The second bag in the two-bag acetylcysteine regimen should be doubled to 200 mg/kg intravenous acetylcysteine over 16 hours (instead of 100 mg/kg over 16 h). Patients with even higher concentrations (eg, ≥ triple the nomogram line) may benefit from even higher acetylcysteine doses. These patients should be discussed with a clinical toxicologist or a Poisons Information Centre. GRADE: Strong; Evidence: Low.

Near the completion of acetylcysteine (ie, 2 h before completion of the infusion), alanine aminotransferase (ALT) should be repeated in all patients. For patients with an initial paracetamol level greater than double the nomogram line, a paracetamol concentration should also be repeated. Acetylcysteine should be continued if the paracetamol concentration is greater than 10 mg/L (66 μmol/L) or ALT is elevated (> 50 U/L) and increasing (if baseline ALT > 50 U/L). The normal reference range for ALT varies between pathology laboratories and with patient age; an elevated ALT > 50 U/L is considered significant. Small fluctuations in ALT (eg, ± 20 U/L or ± 10%) are common and do not on their own indicate the need for ongoing acetylcysteine. ALT should be repeated in all cases as there is a small (< 1%) risk of developing acute liver injury despite treatment with acetylcysteine within 8 hours.5,17,19,20 GRADE: Strong; Evidence: Low.

Multiple or staggered immediate release paracetamol ingestions

Any multiple or staggered paracetamol ingestions over more than 2 hours for the purpose of deliberate self-harm are distinct from repeated supratherapeutic ingestions, which are ingestions of excessive paracetamol for therapeutic purposes (Box 5). Staggered ingestions should be treated as per acute immediate release paracetamol ingestion (Box 3) using the earliest time of ingestion for the paracetamol nomogram. Hence, if it is more than 8 hours since the first dose of paracetamol or the paracetamol concentration cannot be obtained within 8 hours, then commence treatment with acetylcysteine. If the first paracetamol concentration was measured within 2 hours of the last ingested paracetamol dose, it should be repeated after 2 hours to ensure there is no ongoing absorption. If either concentration
3 Acute immediate release paracetamol ingestion management flow chart

ALT = alanine aminotransferase. * Cooperative adult patients who have potentially ingested ≥ 10 g or ≥ 200 mg/kg (whichever is less). For paracetamol ingestions ≥ 30 g, activated charcoal should be offered until 4 hours after ingestion. † Baseline ALT measurement. ‡ If paracetamol concentration will not be available until ≥ 8 hours after ingestion, commence acetylcysteine while awaiting paracetamol concentration. § For acetylcysteine dosage, see Box 7. ¶ Patients should be advised that if they develop abdominal pain, nausea or vomiting, further assessment is required. For patients in rural or remote regions where pathology services are not available, see Box 6.
4 Acute ingestion modified release paracetamol management flow chart

**MR paracetamol ingestion ț 10 g or ș 200 mg/kg (whichever is less)**

- Yes
  - ≤ 4 hours: Activated charcoal
  - ≥ 4 hours: Commence acetylcysteine infusion*

- No
  - Measure two paracetamol concentrations at least 4 hours after ingestion and 4 hours apart

**Dose ingested ș 30 g or ș 500 mg/kg**

- Yes
  - Complete acetylcysteine infusion with double dose second bag (200 mg/kg over 16 h) of acetylcysteine infusion
  - Measure two paracetamol concentrations at least 4 hours after ingestion and 4 hours apart to guide acetylcysteine dose and need for further decontamination

- No
  - Either paracetamol concentration over nomogram treatment line

**Either paracetamol concentration more than double the nomogram line?**

- No
  - Complete standard acetylcysteine infusion*

- Yes
  - Complete acetylcysteine infusion with double dose second bag (200 mg/kg over 16 h) of acetylcysteine infusion*

**ALT and paracetamol concentrations are required in all patients before ceasing acetylcysteine infusion**

Continue acetylcysteine treatment if:
- Paracetamol concentration > 10 mg/L (66 µmol/L), or
- ALT > 50 U/L and increasing (if baseline ALT > 50 U/L)

For criteria of when to cease acetylcysteine, see Box 8

**Recommendations of when to seek further advice from Poisons Information Centre**

- Very large overdoses: modified release paracetamol overdose of > 50 g or 1 g/kg (whichever is less)
- High paracetamol concentration, more than triple the nomogram line
- Serial paracetamol concentrations remain unchanged or increasing

These are situations where the risk of hepatotoxicity may be greater, the optimum advice is developing and where it is useful to seek advice

ALT = alanine aminotransferase; MR = modified release; * Patients should be advised that if they develop abdominal pain, nausea or vomiting, further assessment is required. † If paracetamol concentration is static or rising, a repeat dose of activated charcoal may be beneficial; please seek further advice. ‡ For acetylcysteine dosage, see Box 7. ◆
is above the nomogram line (using time from the earliest ingestion), start or continue treatment with acetylcysteine. **GRADE:** Weak; **Evidence:** Very low.

### Modified release paracetamol ingestions

Modified release paracetamol contains 69% modified release and 31% immediate release paracetamol in a 665 mg tablet. In the previous guidelines, management was very similar to that for immediate release paracetamol. However, evidence from case series from Australia and Europe has shown that this approach appears inadequate. **GRADE:** Strong; **Evidence:** Very low.

### Rural and remote centres

Many rural and remote health care facilities do not have access to 24-hour pathology or have very limited pathology services (e.g., point of care testing only). These facilities can still manage certain acute paracetamol poisoning cases, provided acetylcysteine is available and the patient is not at high risk of developing acute liver injury. **GRADE:** Weak; **Evidence:** Very low.
Paediatric liquid paracetamol ingestion

These recommendations are unchanged from the 2015 guidelines. In children under 6 years of age where ingestion of more than 200 mg/kg of liquid paracetamol is suspected, a serum paracetamol concentration should be measured at least 2 hours after ingestion.24 If the 2–4-hour concentration is below 150 mg/L (1000 μmol/L), acetylcysteine is not required. If the 2-hour paracetamol concentration is greater than 150 mg/L (1000 μmol/L), this should be repeated 4 hours after ingestion and acetylcysteine commenced if this is 150 mg/L or more (1000 μmol/L).

A 2-hour concentration should only be used in a well child under 6 years of age with isolated liquid paracetamol ingestion. In all other cases, a 4-hour concentration should be performed. Further, for children who present later than 4 hours after ingestion or in children older than 6 years of age, treatment is as per the adult acute paracetamol exposure guideline. GRADE: Strong; Evidence: Very low.

Repeated supratherapeutic ingestion

Patients who ingest excessive paracetamol for a therapeutic purpose (eg, pain, viral illness) or ingest therapeutic doses of paracetamol and have symptoms of acute liver injury (eg, abdominal pain, nausea and vomiting) are managed as per repeated supratherapeutic ingestion (Box 5). If the ingestion is deliberate or intentional, they should be managed as per acute intentional ingestion. There is little evidence to guide risk assessment for repeated ingestion of high doses of paracetamol. The margin of safety has for many years been assumed to be high.25 Minor subclinical elevations of serum ALT are quite common with prolonged therapy.26 Hepatotoxicity has been reported at doses within the therapeutic range of paracetamol (in some cases at doses less than the recommended 4 g/day). The reasons why certain individuals are at greater risk of toxicity are unclear,27 but toxicity could be...
influenced by age, comorbidities, alcohol use, nutritional status (eg, prolonged fasting), concurrent medicine use, and genetics. Some patients are likely to be at increased risk for acute liver injury after repeated supratherapeutic ingestion due to glutathione depletion or cytochrome P450 (CYP450) induction. Clinical flags would include pregnancy, prolonged fasting, chronic alcoholism, febrile illness and chronic use of CYP450-inducing drugs, such as carbamazepine. Hence, the threshold for a potentially toxic dose has been made deliberately and conservatively low in these and previous guidelines. Patients who meet the criteria for supratherapeutic ingestion (Box 1) should have the paracetamol concentration and ALT measured. There is evidence that the combination of a low paracetamol concentration and normal ALT at any time indicates there is minimal risk of subsequent hepatotoxicity. If the paracetamol concentration is greater than 20 mg/L (132 μmol/L) or ALT is greater than 50 U/L, then acetylcysteine is commenced, and pathology repeated 8 hours after the initial sampling. Acetylcysteine can be ceased at this stage if the paracetamol concentration is less than 10 mg/L and ALT is less than 50 U/L or static. Patients with significant acute liver injury secondary to paracetamol will have a very high and/or rapidly rising ALT. Small fluctuations in ALT (eg, ± 20 U/L or ± 10%) are common and do not on their own indicate the need for ongoing acetylcysteine. All patients with an initial ALT greater than 1000 U/L should receive at least a full 20-hour course of intravenous acetylcysteine. GRADE: Strong; Evidence: Very low.

Cessation of acetylcysteine

Some patients will require ongoing treatment with acetylcysteine if they have a persistently high paracetamol concentration greater than 10 mg/L (66 μmol/L) or ALT greater than 50 U/L and increasing (if baseline ALT > 50 U/L) — small fluctuations in ALT (eg, ± 20 U/L or ± 10%) are common and do not on their own indicate the need for ongoing acetylcysteine. There is very little evidence to guide when is the optimum time to cease acetylcysteine, with no published data on what is an acceptable rate of decline in transaminases. In patients requiring ongoing acetylcysteine, treatment should be continued until the patient has met all the criteria outlined in Box 8, with international normalised ratio (INR) below 2 and the patient being clinically well likely more important than the ALT/aspartate aminotransferase (AST) decline. Acetylcysteine is generally continued at the rate of the second infusion (eg, 100 mg/kg over 16 h) (Box 7). Higher infusion rates may be warranted for massive paracetamol ingestions, especially if the paracetamol concentration is 100 mg/L or more (660 μmol/L) at the completion of the initial acetylcysteine infusion — a clinical toxicologist should be consulted in such cases. Regular clinical review and blood tests at least every 12 hours are recommended for patients requiring prolonged treatment. GRADE: Strong; Evidence: Low.

Hepatotoxicity and subsequent liver failure

Only a small proportion of patients develop hepatotoxicity (ALT > 1000 U/L). Early symptoms include nausea, vomiting, abdominal pain, and right upper quadrant tenderness. Of these, only a minority will develop fulminant hepatic failure, and most patients recover fully with standard treatments. Typically, in patients with paracetamol-induced acute liver injury, ALT and AST will rise for 3–4 days before recovering. Acetylcysteine is continued until the criteria in Box 8 are met. Investigations that monitor liver function and guide prognosis should be performed regularly in all patients with hepatotoxicity, including electrolytes, urea, creatinine, liver function tests, INR, blood sugar, phosphate, and venous blood gas (looking at the pH and lactate levels). A liver transplant unit should be consulted if any of the following criteria are met:

- INR greater than 3.0 at 48 hours or greater than 4.5 at any time;
- oliguria or creatinine greater than 200 μmol/L;
- persistent acidosis (pH < 7.3) or arterial lactate greater than 3 mmol/L;
- systolic hypotension with blood pressure below 80 mmHg, despite resuscitation;
- hypoglycaemia, severe thrombocytopenia, or encephalopathy of any degree; or
- any alteration of consciousness (Glasgow Coma Score < 15) not associated with sedative co-ingestions.

Do not give clotting factors unless the patient is bleeding or after discussion with a liver transplant unit. GRADE: Strong; Evidence: Strong.

Seeking advice from a Poisons Information Centre

It is recommended to seek advice from a Poisons Information Centre in the following situations:...
结论

This is a summary of the updated guidelines for the management of paracetamol poisoning in Australia and New Zealand, for more detailed information please access the full guidelines, available in the online Supporting Information.

If there are any concerns regarding the management of paracetamol ingestion, advice can always be sought from a clinical toxicologist or a Poisons Information Centre (dialling 131126, in Australia, or 0800 764766, in New Zealand).

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This article is a summary of the full guidelines, available online in the Supporting Information.

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Supporting Information

Additional Supporting Information is included with the online version of this article.