

HYPERAMMONAEMIA PROTOCOL (New patient)

1 DEFINITION AND SYMPTOMS

Hyperammonaemia corresponds to a rise in the level of ammonia in the blood. Ammonia is a waste product coming from the metabolism of proteins, which are cleared by the liver via the urea cycle.

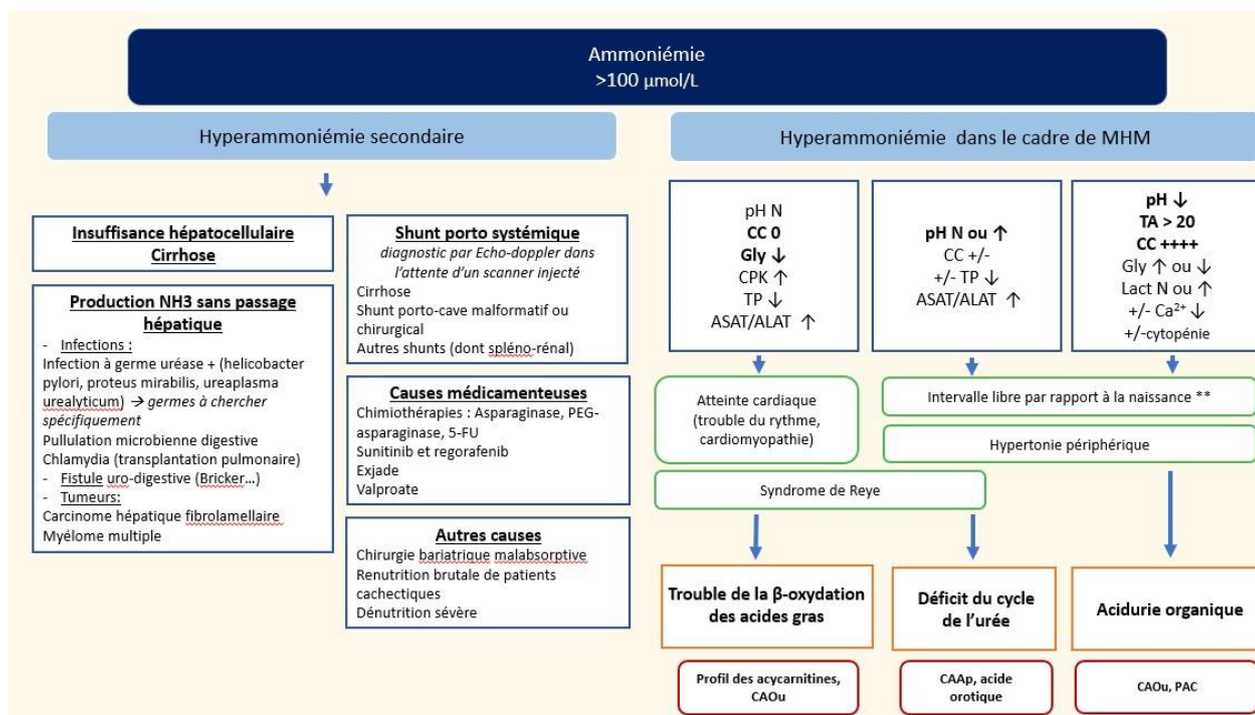
Ammonia is **toxic to the liver and the brain**. Therefore, treatment of hyperammonaemia is a **medical emergency**. Ammonia is an endogenous toxin which it is important to measure in all situations where an exogenous toxin assay is indicated, particularly where unexplained neuropsychiatric or hepatic/gastrointestinal symptoms are present.

The normal level of blood ammonia is $<100 \mu\text{mol/L}$ in a neonate, and $<50 \mu\text{mol/L}$ thereafter regardless of age. (conversion factor: $\text{mg/L} \times 58.7 = \mu\text{mol/L}$)

The level of ammonia in the blood reflects the total ammonia load and the distribution volume. Hence, the seriousness of hyperammonaemia at $100 \mu\text{mol/L}$ will vary depending on the patient's age (the risk being higher in older patients than in infants).

The causes of hyperammonaemia are linked to:

- A primary dysfunction of the urea cycle (urea cycle disorders: OTC, CPS, NAGS, citrullinaemia, arginosuccinic aciduria, arginase deficiency)
- A dysfunction of the urea cycle secondary to another inherited metabolic disease (IMD): deficiency of fatty acid beta-oxidation, organic aciduria, some defects of ketogenesis or ketolysis, some mitochondrial cytopathies.
- Secondary causes (see figure): liver failure, insufficient passage through the liver due to a portosystemic shunt (*increased serum bile acids*), massive endogenous toxicity (infection with a urease-positive bacterium, gastrointestinal or urinary tract microbial overgrowth etc.), exogenous drugs affecting the urea cycle (chemotherapy, other treatments), tumour-related causes.



3 TREATMENT TO BE STARTED URGENTLY

Immediately check blood ammonia levels and start treatment without waiting for the results.

In an adult, if the initial diagnosis points to a secondary cause of hyperammonaemia (particularly hepatic encephalopathy related to cirrhosis or a shunt) refer to paragraph C.

A. Baseline infusion

- NO amino acids IV, no oral proteins: **stop feeding**
- **Infusion of 10% glucose (G10)** with standard electrolyte additions* (not pure 10% glucose), via a peripheral line.

Age	0 - 24 months	2 - 4 years	4 - 14 years	> 14 years / adult	MAX. FLOW RATE
G10% + added electrolytes*	6ml/kg/hr (10mg/kg/min)	5ml/kg/hr (8mg/kg/min)	3.5ml/kg/hr (6mg/kg/min)	2.5ml/kg/hr (4mg/kg/min)	120ml/hr (3L/24 hrs)

*e.g.: Polyionique, Bionolyte, B45, Glucidion etc. to which must be added 2 g/L NaCl to obtain 6 g/L in total or, if pre-made solutions not available, 10% glucose in water + 6 g/L NaCl and 2g/L KCl

- No lipids initially, **until impaired fatty acid oxidation has been excluded**. If indicated, 20% lipids given via a Y-set

Age	0 - 24 months	> 2 years	MAX. FLOW RATE
20% lipids (if needed)	0.4 ml/kg/hr (2g/kg/day)	0.3ml/kg/hr (1.5g/kg/day)	20ml/hr (500ml/24 hrs)

B. Treatments to clear ammonia: **Start sodium benzoate straight away and add other detoxifying agents if available:**

- **Sodium benzoate** by continuous IV infusion Start with a **loading dose** of 250 mg/kg over 2 hours (**max. 8g over 2 hours**), then 250 to 500 mg/kg/24 hrs (**max. 12g/24 hrs**) (give orally or via NG tube if no venous access available). Check blood ammonia at the end of the loading dose. SODIUM BENZOATE IV: [Sodium benzoate AP-HP 1g-10mL]; ampoule 1g = 10ml, to be diluted 1:1 by volume in 10% glucose. Contains 7 meq of sodium per gram of benzoate. Status = hospital preparation
- Ammonaps orally only, possible as continuous administration via NG tube or in 4 to 6 doses: loading dose 250 mg/kg (max. 8g) then 250 to 500 mg/kg/24 hrs (**Max. 12g/24 hrs**). Combine with sodium benzoate.
- **Carbaglu Carbaglu®** (N-carbamyl-glutamate) if available: oral loading dose 50-100 mg/kg then maintenance dose 50 mg/kg/6 hours orally or via NG tube (Max. 8g over 24hrs).

C. Treatment of secondary causes

In addition to chelating treatment, start specific treatment for the secondary causes:

- Gastrointestinal microbial overgrowth: Tixtar (rifaxymin) orally 500mg x2/day (adult) or 250 mgx2/day (child): in an adult only following advice from liver specialist
- Urinary microbial overgrowth: Flagyl (usual doses)
- Infection with a urease-positive bacterium: antibiotic treatment targeting intracellular bacteria
- 5-FU and PEG-asparaginase: consider a fruit and vegetable diet on a case-by-case basis, with the possibility of accompanying further treatment with chelator treatment.
- Seek liver specialist opinion for managing cirrhosis and/or shunt and/or liver failure
- Hepatic encephalopathy in an adult (while waiting for the advice of a liver specialist): Lactulose (enema or retention enema, then switch to oral administration following the dosage recommended in Vidal).

4 SEVERITY SIGNS / MANAGEMENT IN ICU

- Coma or lack of neurological improvement 3 hrs after starting treatment
- and/or severe hyperammonaemia (Infant >200 µmol/L - Child & adult >150 µmol/L)
- and/or severe liver failure
 - Start Ammonul® (250 mg/kg/day), ideally via a central line, max.12g/24 hrs (stop sodium benzoate and phenylbutyrate)
 - In the meantime, option of giving an additional loading dose of Ammonaps: 250mg/kg orally (max. 10g).
 - Consider haemodialysis
 - **Preferably use a central line as soon as possible to concentrate the infusion** (risk of cerebral oedema) while maintaining glucose and sodium supply [example: 30% glucose (enough to provide the same glucose intake as above), NaCl 6 g/L (100meq/L), potassium and calcium according to serum electrolyte results + physiological saline (NaCl 0.9%) in parallel via a Y-set, to give a total intake of **1.5 L/m²/day** (*Body surface area = (4 x W + 7) / (W + 90)*)
 - In ICU: Measures for neuroprotection and avoiding secondary insults to the brain of systemic origin

5 MONITORING

- Follow-up checks (NH₃, PT, serum electrolytes): at the end of the benzoate loading dose, then every 4 to 6 hours depending on progression. Correction of potential electrolyte abnormalities (in particular hypokalaemia in arginosuccinic aciduria)
- Capillary blood glucose every 4 hrs: target 1 to 1.8g/L. If blood glucose >2g/L and glycosuria, consider insulin 0.01IU/kg/hr, adjust in line with blood glucose checks. Consider reducing sugar intake (25% to 50%) if, despite insulin therapy, hyperglycaemia persists. at 0.05 IU/kg/hr and/or appearance of hyperlactataemia > 3mmol/L.

NUMBERS AND MEDICAL SPECIALISTS

On-call telephone numbers for metabolic emergencies:

At night, only medical teams can call in emergency situations, and only if the emergency certificate has not been understood or if the clinical state or test results are worrying. As far as possible, make calls before nighttime.

Secretarial issues must be dealt with via the medical secretariat during the week, or by email addressed to the patient's metabolic medicine specialist.