Elevated plasma ammonia is a medical emergency

- The neurological outcome of affected neonates is directly related to the duration of hyperammonaemic coma.

DO NOT delay treatment whilst awaiting results of further investigations.

Definition

- Premature neonate > 150 μmol/L.
- Term neonate > 100 μmol/L.

Symptoms

- Symptoms are non-specific but include tachypnoea, seizures and encephalopathy. Consider hyperammonaemia in the differential diagnosis of any sick neonate.

Differential Diagnosis

- Spurious: Incorrect sampling (sample haemolysed, not collected on ice, or delayed separation).
- Hepatic: Liver failure / impairment.
- Metabolic Urea Cycle defects (UCD):
  - Organic acidaemias (OA).
  - Fatty acid oxidation defects (FAOD).
- Transient hyperammonaemia of the newborn (due to open ductus venosus).
- HIIHA - Hyperinsulinism/hyperammonaemia - hypoglycaemia with raised ammonia.
# Investigations

<table>
<thead>
<tr>
<th>ACUTE INVESTIGATIONS</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ammonia (x2)</td>
<td>Free flowing, consider arterial if difficult. Place on ice and transport urgently to laboratory.</td>
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<tr>
<td>Blood gas</td>
<td>Respiratory alkalosis in UCD, metabolic acidosis in OA.</td>
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<tr>
<td>Lactate</td>
<td>Raised in OA &amp; FAOD &amp; UCD with circulatory collapse.</td>
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<tr>
<td>Liver function</td>
<td>Deranged in liver failure, OA &amp; FAOD.</td>
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<tr>
<td>Clotting</td>
<td>Deranged in liver failure, OA &amp; FAOD.</td>
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<tr>
<td>Glucose</td>
<td>Low in FAOD &amp; OA.</td>
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<tr>
<td>Ketones (urine dipstick)</td>
<td>Low in FAOD &amp; UCD, raised in OA.</td>
</tr>
<tr>
<td>Amino acids (plasma)</td>
<td>Glucose and ketones are not necessarily reduced in OA’s and UCD’s.</td>
</tr>
<tr>
<td>Amino acids (urine)</td>
<td>Obtain baseline.</td>
</tr>
<tr>
<td>Organic acids/ Orotic acid (urine)</td>
<td>Obtain baseline.</td>
</tr>
<tr>
<td>Acylcarnitines</td>
<td>If Hiawatha’s is a consideration with associated hypoglycaemia - then insulin tests should be done additionally.</td>
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</table>

## Acute Management

1. **Stop all enteral feeds.** Promote anabolism. Start intravenous glucose to ensure glucose infusion of 6 mg/kg/minute, aiming for glucose 4-8 mmol/L.

2. Add insulin infusion 0.05 U/kg/hr if blood glucose > 10 mmol/L. DO NOT just turn down the rate of 10% glucose (remember that the aim is to stop catabolism and this can only be done by giving lots of calories).

3. If boluses of fluid are required, use 0.9% Saline. Remember that 4.5% human albumin solution and FFP contain protein.

4. If ammonia > 100µmol/L repeat in 4 hours.

5. If ammonia > 150 µmol/L, discuss with Metabolic Consultant, as urgent exclusion of possible ‘small molecule’ metabolic disease has to be considered and managed appropriately (see step 7).

6. The decision to administer ammonia scavengers (sodium benzoate and phenylbutyrate - both at 250 mg/kg) either as a loading dose (i.e. over ~ 2 hours) or maintenance over 24 hours depends on the level of elevation of ammonia and clinical presentation (i.e. encephalopathic etc).
   - **URGENTLY** obtain a rapid review of newborn screening results as well as urine metabolic screen, plasma amino acids, and acylcarnitine profile and liaise with the metabolic team.
   - Arrange cranial ultrasound to look for cerebral oedema.
   - Sodium benzoate and phenylbutyrate are stocked in NICU (hyperammonaemia kit). See Neonatal Medication Protocol Hyperammonaemia Medications.

**KEMH:** Hyperammonaemia kit is kept in SCN3 Medication imprest cupboard.
Hyperammonaemia

PCH: Medications are kept in the ADM as virtual kit: ‘Hyperammonaemia Kit’
Contact the on-call pharmacist urgently if there are any problems with
accessing the kit.
- In addition, consider arginine ~ 250 mg/kg/day as a continuous infusion but
depends on type of urea cycle defect etc. Hence, important to get metabolic
tests analysed urgently.

7. The ammonia level at which a decision needs to be made on haemodiafiltration
(usually if >300) depends on clinical presentation, and how soon the levels drop
post-loading dose of ammonia scavengers. If there is a need to prepare for
probable haemodiafiltration. Contact PCC Consultant to discuss. Remember that it
can take some time to get appropriate central access and commence
haemodiafiltration so ALL other measures to lower plasma ammonia (i.e. steps 1, 2,
3, 4 and 5) must be instituted as soon as possible.

8. In the event of imminent death, an ante mortem liver biopsy and skin biopsy should
be obtained, to assist with diagnosis. Contact the biochemistry laboratory to
arrange. 2 mLs EDTA whole blood should also be collected for potential genetic
studies and state “extracted DNA for storage” on lab form.

9. Finally, consider carefully whether to perform a post-mortem, even if liver and skin
biopsies have been taken.

Related CAHS internal policies, procedures and guidelines

<table>
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<th>Neonatology Medication Protocols</th>
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<td>• Hyperammonaemia Medications</td>
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formats on request for a person with a disability.

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