#### **GUIDELINE**

## Hypoglycaemia

| Scope (Staff): | Nursing and Medical Staff                          |
|----------------|--|
| Scope (Area):  | KEMH postnatal wards, NICU KEMH, NICU PCH, NETS WA |

#### **Child Safe Organisation Statement of Commitment**

CAHS commits to being a child safe organisation by applying the National Principles for Child Safe Organisations. This is a commitment to a strong culture supported by robust policies and procedures to reduce the likelihood of harm to children and young people.

#### This document should be read in conjunction with this disclaimer

#### **Contents**

| Aim  | . 2 |
|--|-----|
| Risk   | . 2 |
| Background   | . 2 |
| Cause /Risk Factors for Hypoglycaemia  | . 2 |
| Infants at Risk of Hypoglycaemia   | . 3 |
| Infants at risk of hypoglycaemia that require early energy provision and BGL monitoring: | 3   |
| Early Energy Provision - Within 1-2 Hours of Birth                                       | . 3 |
| Management of Hypoglycaemia  | . 4 |
| Enteral Feeding  | . 4 |
| Parenteral Supplementation   | . 4 |
| Investigation of Neonatal Hypoglycaemia – "Hypoglycaemia Screen"                         | 6   |
| Persistent Hyperinsulinaemic Hypoglycaemia of Infancy (PHHI)                             | 6   |
| Infant with PPHI requiring short term diazoxide:   | 6   |
| Congenital Hyperinsulinism of Infancy (CHI)  | . 7 |
| Follow-up for Infants with Evidence of Hypoglycaemia                                     | . 7 |
| QUICK REFERENCE GUIDE FOR AT – RISK INFANT   | 8   |
| Appendix 1:  | 11  |
| Centile Chart for Hypoglycaemia  | 11  |

#### **Aim**

To facilitate early recognition and management of hypoglycaemia in infants at risk.

#### Risk

Symptomatic hypoglycaemia is a medical emergency and requires intravenous treatment. Failure to recognise and manage hypoglycaemia early in the newborn period may increase morbidity and mortality.

## **Background**

Asymptomatic hypoglycaemia is a common transient problem in most neonates. Symptomatic hypoglycaemia is an emergency and requires intravenous treatment. Symptoms include:

- CNS excitation: irritability, jitteriness, seizures.
- CNS depression: Hypotonia, lethargy, poor feeding, apnoea's.
- Non-specific: temperature instability, sweating, tachycardia.

The fetus under normal conditions derives all its glucose from the mother. At birth all infants must initiate glucose production and absorption. Most can mobilise glycogen, initiate gluconeogenesis and produce glucose at a rate of 4 - 6 mg/kg/min. This is usually adequate to maintain euglycaemia - normal blood glucose.

The definition used at KEMH and PCH for hypoglycaemia is a blood glucose of < 2.6mmol/L.

## Cause /Risk Factors for Hypoglycaemia

The cause/risk factors for hypoglycaemia can be divided into:

| Inadequate supply or reduced glycogen stores | Increased utilisation                                     | Hormone/metabolism imbalance  |  |
|--|---|---|--|
| <ul> <li>Prematurity</li> </ul>              | <ul><li>Infection</li></ul>                               | Infant of a diabetic mother   |  |
| Small for gestational age                    | <ul> <li>Respiratory Distress<br/>Syndrome</li> </ul>     | <ul> <li>Persistent hyperinsulinaemic<br/>hypoglycaemia of infancy</li> </ul> |  |
| <ul> <li>Poor feeding</li> </ul>             | <ul> <li>Hypothermia</li> </ul>                           | Inborn errors of metabolism   |  |
| <ul> <li>Peripheral IV tissued</li> </ul>    | <ul><li>Perinatal Asphyxia</li><li>Hyperthermia</li></ul> | Syndrome: Beckwith-<br>Wiedemann  |  |
|  | Erythroblastosis Fetalis                                  | Pancreatic tumour   |  |
|  |   | Congenital adrenal hyperplasia  |  |
|  |   | Hypopituitarism   |  |

Page 2 of 11 Neonatal Guideline

Persistent or recurrent hypoglycaemia (≥ 2 episodes of hypoglycaemia) warrants further investigation. It is commonly caused by hyperinsulinism secondary to maternal diabetes however other differentials should be considered such as CAH, syndromes, and inborn errors of metabolism.

## Infants at Risk of Hypoglycaemia

It is important to explain to the parents of at-risk infants that their infant is more likely than others to develop hypoglycaemia, and that their infant will need close monitoring of blood glucose. Refer to Quick reference guide for management.

**Infants at risk of hypoglycaemia** that require early energy provision and BGL monitoring:

- Infants of mothers with diabetes (insulin-dependent, type 2 DM or GDM).
- Infants who are small for gestational age (< 10th percentile) refer to <u>Appendix 1</u>
- Preterm infants (<37 weeks gestation).</li>
- Infants large for gestational age (>4.5kg or >97th centile) refer to Appendix 1
- Infants of mothers who received antenatal corticosteroids >34 weeks gestation.
- Infants of mothers who received beta blockers in the 3<sup>rd</sup> trimester.

## **Early Energy Provision - Within 1-2 Hours of Birth**

- Assist thermoregulation by offering early skin to skin under warm blankets.
- Within the first hour after birth offer a breast feed followed by 3 hrly feeds/more frequently if demanding.
- If **first** breastfeed is ineffective, consider supplemented enteral feeding (if not contraindicated.) Offer EBM/Term formula at 60ml/kg/day (or 80ml/kg/day, if <37 weeks gestation), immediately following the ineffective breast feed.
- If not breast feeding offer Term formula 7.5ml/kg within the first hour of birth followed by 3 hrly feeds 60ml/kg/day (or 80ml/kg/day, if <37 weeks gestation.)
- If enteral feeding is not possible then admit to Special Care Nursery and give IV 10% Glucose. Start at 60mL/kg/day (providing 4.2 mg/kg/min of glucose).

Page 3 of 11 Neonatal Guideline

#### **Glucose Monitoring of at-Risk Infants**

- Whole blood glucose (blood gas analyser) or plasma glucose (biochemistry lab) should be performed. Reagent strips should not be used for neonatal PGL monitoring.
- For at risk infants, first sample taken pre-second feed (3-4 hours of age).
- If infant feeding well and PGL ≥ 2.6mmol then repeat PGL 6 hourly (pre-feed).
  - o If 2 consecutive PGLs are ≥ 2.6mmol/L, then stop regular monitoring and test only if infant becomes symptomatic.

## Management of Hypoglycaemia

#### Asymptomatic Infants with PGL 1.5-2.5mmol/L

Needs paediatric RMO/ registrar review - consider "hypoglycaemia screen" (see below) and need for admission to SCN.

#### **Enteral Feeding**

- Start enteral feeding at 60-80mL/kg/day if no contra-indications.
- If there is insufficient breast milk and parents decline formula, escalate rapidly to senior neonatal staff to avoid delays in treatment.
- If persistent or recurrent hypoglycaemia, then increase feed volume to 12.5mL/kg/feed (100ml/kg/day).
- Consider more regular feeds (2 hourly)
- Admit to SCN for IV dextrose if:
  - o PGL remains between 1.5-2.5mmol/L despite the increased feeds.
  - o Infant is symptomatic (lethargic with inadequate feeds, seizure).

#### **Parenteral Supplementation**

- If unable to obtain IV access, consider glucagon (IM 100 micrograms/kg) and consider siting a UVC.
- Commence IV supplementation with 10% dextrose at 80-100mL/kg/day (5.6-7mg/kg/min).
  - Consider bolus of 2mL/kg of 10% dextrose.
- Monitoring
  - Repeat PGL after 30 minutes of IV glucose if normal then check again at 3 hours.
  - If 30 minute and 3-hour PGL is normal, monitor PGL 3-6 hourly.

Page 4 of 11 Neonatal Guideline

#### Asymptomatic Infants with PGL < 1.5mmol/L

Admit to SCN immediately for IV supplementation.

- Take hypoglycaemia screen (see below) if it does not delay treatment significantly.
- Commence IV supplementation with 10% dextrose at 100mL/kg/day (7mg/kg/min).
  - o If unable to obtain IV access, consider glucagon (IM 100 micrograms/kg).
  - o Consider bolus of 2mL/kg of 10% dextrose.
  - If hypoglycaemia continues, then aim to increase GIR by 2-3mg/kg/min (Increase total fluids by 20-30mL/kg/day or increase dextrose concentration by 2.5-5%).
  - o If needing > 12.5% dextrose, then central access (UVC) is required.
- Monitoring
  - o Recheck PGL at 30 minutely intervals until PGL is ≥ 2.6mmol/L.
  - o Once PGL is ≥ 2.6mmol/L then check 3 hourly.
  - o If 2 consecutive 3 hourly PGLs are normal, monitor PGL 6 hourly.

#### **Symptomatic Infants – Seizures, Reduced Consciousness**

- If on the post-natal ward, access and use the hypoglycaemia kit in the emergency trolley
- Admit to NICU for urgent IV supplementation
- Take hypoglycaemia screen (see below) if it does not delay treatment significantly.
- Commence IV supplementation with 10% dextrose at 100mL/kg/day (7mg/kg/min).
  - If unable to obtain IV access, give glucagon (IM 100 micrograms/kg) and consider siting UVC.
  - o Give bolus of 2mL/kg of 10% dextrose; repeat until seizure has stopped.
- Monitoring
  - o Recheck PGL at 15-30 minutely intervals until PGL is ≥ 2.6mmol/L.
  - Once PGL is ≥ 2.6mmol/L then check 3 hourly.
- If 2 consecutive 3 hourly PGLs are normal, monitor PGL 6 hourly.

Page 5 of 11 Neonatal Guideline

# Investigation of Neonatal Hypoglycaemia – "Hypoglycaemia Screen"

If hypoglycaemia is persistent/recurrent (≥ 2 episodes), resistant to treatment, or glucose delivery rate is > 10mg/kg/min then investigate further with a hypoglycaemia screen.

#### Hypoglycaemia Screen

The critical blood samples **MUST** be collected at the time of hypoglycaemia, wherever safe, prior to commencing supplementation.

**DO NOT** administer sucrose before heel stab/ venepuncture.

- 1 mL of clotted blood and 1 mL of heparinised blood (2 small red top and 2 small green top tubes).
  - $\circ$  Request insulin, cortisol, growth hormone, glucose, ketones or β-hydroxybutyrate.
- Blood gas analysis: lactate.
- The NEXT urine passed is important (aim for 5 mL urine).
  - o Request ketones, amino acids, and organic acids.

# Persistent Hyperinsulinaemic Hypoglycaemia of Infancy (PHHI)

PHHI is commonly seen in infants born to a mother with gestational diabetes, however, can occur in mothers with a normal glucose tolerance test. It is diagnosed by finding an elevated insulin level during a period of hypoglycaemia. Infants with PHHI may require a significantly higher glucose delivery rate of up to 10-12mg/kg/min.

#### Infant with PPHI requiring short term diazoxide:

- If glucose delivery rate > 10mg/kg/min and has unstable PGLs then consider diazoxide.
- Please discuss with endocrinology if patient is to commence on diazoxide.
- Once infant is ready for discharge then a prolonged fast is required (6 hours).
  - o 3-hour pre-feed PGL, then hourly up to 6 hours.
  - If during the fast the PGL drops below 3.0mmol/L, then a hypoglycaemia screen should be sent, and endocrinology informed.
- Endocrinology should be informed of all babies that have received diazoxide and are being discharged home as they will organise a 6-week follow-up in their outpatient clinic.

Page 6 of 11 Neonatal Guideline

## **Congenital Hyperinsulinism of Infancy (CHI)**

Infants that cannot be weaned off diazoxide or have unstable PGLs on diazoxide may require further investigations to exclude CHI.

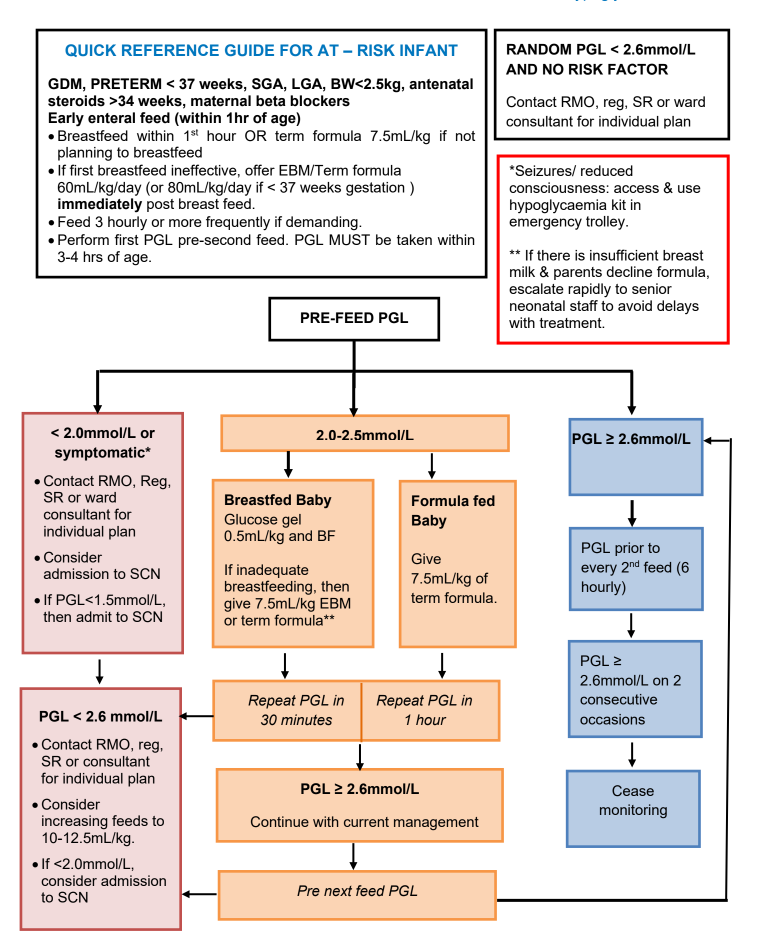
CHI is a clinically and genetically heterogeneous disease, is characterized by the unregulated secretion of insulin from pancreatic beta-cells. It is the commonest cause of PHHI

The most common and severe forms of CHI are caused by inactivating mutations in ABCC8 and KCNJ11 genes, encoding the two subunits of the pancreatic beta-cell ATP-sensitive potassium channel (KATP). Discuss further with endocrinology.

## Follow-up for Infants with Evidence of Hypoglycaemia

All infants who have been symptomatic or had persistent asymptomatic hypoglycaemia need follow up, the intensity of which needs to be graded to the severity. An MRI of the brain should be considered for all infants with severe and/or prolonged hypoglycaemia, or who developed hypoglycaemic seizures.

Page 7 of 11 Neonatal Guideline



Page 8 of 11 Neonatal Guideline

#### Related CAHS internal policies, procedures, and guidelines

#### **Neonatal Medication Protocols**

- Diazoxide
- Glucagon

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Page 9 of 11 Neonatal Guideline

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| Document Owner:  | Neonatology  |       |                                 |  |  |
|--|--|-------|---------------------------------|--|--|
| Reviewer / Team:   | Neonatology Coordinating Group   |       |                                 |  |  |
| Date First Issued:   | July 2006 Last Reviewed:   |       | October 2023                    |  |  |
| Amendment Dates:   | October 2023 Combined postnatal ward guideline with neonatology guideline.  February 2024 QRG wording changed to clarify when first PGL is to be taken.  Next Review Date: |       | October 2026                    |  |  |
| Approved by:   | Neonatology Coordinating Group   | Date: | O 4th O 1 1 1 2 2000            |  |  |
| Endorsed by:   | Neonatology Coordinating Group   | Date: | - 24 <sup>th</sup> October 2023 |  |  |
| Standards<br>Applicable:   | NSQHS Standards: 1,10  |       |                                 |  |  |
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Page 10 of 11 **Neonatal Guideline** 

# **Appendix 1:**

## **Centile Chart for Hypoglycaemia**

| Birth weight of term babies at the <b>10</b> <sup>th</sup> <b>centile</b> |                    | Gestation | Birth weight of term babies at the <b>97</b> <sup>th</sup> <b>centile</b> |                    |
|---|--------------------|-----------|---|--------------------|
| Male<br>(weight)  | Female<br>(weight) | (weeks)   | Male<br>(weight)  | Female<br>(weight) |
| 1900  | 1800               | 35        | 3280  | 3200               |
| 2170  | 2050               | 36        | 3550  | 3500               |
| 2400  | 2300               | 37        | 3800  | 3800               |
| 2600  | 2500               | 38        | 4020  | 4020               |
| 2800  | 2650               | 39        | 4280  | 4250               |
| 3000  | 2800               | 40        | 4500  | 4450               |
| 3200  | 3000               | 41        | 4750  | 4680               |
| 3400  | 3150               | 42        | 5020  | 4920               |

Page 11 of 11 Neonatal Guideline