

**Summary** This Guideline provides a framework for early identification of neonates ≥ 32 weeks gestation at risk of jaundice and provides guidance for appropriate care and management across the state. The Guideline assists clinicians to differentiate between pathological neonatal jaundice and those neonates with benign physiological jaundice and the appropriate treatment.

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**Distributed to** Public Health System, Divisions of General Practice, Government Medical Officers, NSW Ambulance Service, Ministry of Health, Private Hospitals and Day Procedure Centres, Tertiary Education Institutes

**Audience** Staff of Maternity; Paediatric; Neonatal; Emergency; Biomedical; Pathology; Community Health; GPs



# NEONATAL - JAUNDICE IDENTIFICATION AND MANAGEMENT IN NEONATES ≥ 32 WEEKS GESTATION

### **PURPOSE**

This Guideline provides a framework for the early identification and management of jaundice in neonates ≥ 32 weeks gestation. Approximately 60% of neonates born at term and 85% of preterm neonates will develop jaundice. Many of these neonates will develop 'physiological jaundice', which is usually benign. However, when unconjugated serum bilirubin levels are too high, bilirubin can cross the blood brain barrier. Bilirubin is neurotoxic, particularly to the auditory nerve and basal ganglia, which can result in brain injury and lifelong disability. It is important therefore, to identify those neonates at risk of acute bilirubin encephalopathy and kernicterus.

### **KEY PRINCIPLES**

This Guideline applies to all NSW Public Health Organisations providing care for neonates ≥ 32 weeks gestation which should include:

- The identification at birth of neonates with risk factors for neonatal jaundice
- Regular visual assessment from birth of all neonates
- Management of neonatal jaundice identified in the first 24 hours of age
- Management of neonatal jaundice identified ≥ 24 hours of age
- Follow-up care for neonates discharged at less than 3 days of age with risk factors for jaundice or jaundice at discharge
- Assessment and escalation of care for neonates with prolonged jaundice > 14 days of age in a term neonate, and beyond 21 days in a preterm neonate.

### **USE OF THE GUIDELINE**

The Chief Executives of all NSW Local Health Districts are responsible for the implementation of this guideline within their services / facilities to ensure:

- Local processes and operating procedures are developed in line with this document to manage neonates ≥ 32 weeks gestation to ensure:
  - Prompt appropriate identification, management and escalation of neonatal jaundice
  - Equipment is used, maintained and its effectiveness is monitored
  - o Discharge is planned and follow up processes are in place
  - Assessment and appropriate escalation of care for neonatal jaundice > 14 days of age in a term neonate and beyond 21 days in a preterm neonate.
- The Directors of Clinical Governance inform relevant staff in maternity, neonatal services and biomedical departments of this new Guideline



 Morbidity and mortality associated with neonatal jaundice is monitored and reviewed.

# **REVISION HISTORY**

Version	Approved by	Amendment notes
November 2016	Deputy Secretary, Strategy and	New policy
(GL2016_027)	Resources	

### **ATTACHMENTS**

1. Neonatal - Jaundice Identification and Management in Neonates ≥ 32 weeks Gestation: Guideline.

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### 1. BACKGROUND

Approximately 60% of neonates born at term and 85% of preterm neonates will develop jaundice<sup>1,2</sup>. Many of these neonates will develop 'physiological jaundice', which presents on day 3, peaks between 5 to 7 days of age and resolves by 14 days of age<sup>2</sup>. Physiological jaundice is usually benign. However, when unconjugated serum bilirubin (SBR) level is too high, bilirubin can cross the blood brain barrier. Bilirubin is neurotoxic, particularly to the auditory nerve and basal ganglia, which can result in brain injury and lifelong disability. It is important therefore, to identify those neonates at risk of acute bilirubin encephalopathy and kernicterus<sup>1,2,3</sup>.

The clinical challenge is to differentiate the minority of neonates ≥ 32 weeks with pathological neonatal jaundice from the majority with benign physiological jaundice.

### 1.1 Scope

This document provides guidance to all clinicians responsible for the care of neonates who are born at ≥ 32 weeks gestation. This Guideline does not apply to neonates born at < 32 weeks who require neonatal specialist care.

### 1.2 Key definitions

### 1.2.1 Terminology

**Should** - indicates a recommended action that should be followed unless there are sound documented reasons for taking a different course of action.

**Neonate** - any baby from time of birth up to and including 28 days of age.

**Preterm** - a baby born before 37<sup>+0</sup> weeks gestation.

**Late preterm** - a baby born between 34<sup>+0</sup> and 36<sup>+6</sup> weeks gestation.

**Well neonate** - a neonate, whose assessments are within normal range on the standard neonatal observation chart (SNOC).

**Local paediatric-specific Clinical Emergency Response System (CERS)** - a local **paediatric-specific** CERS protocol should be in place to define the process to escalate and access a senior medical officer or specialist paediatrician who has the care of the neonate incorporated in their scope of practice, and if required, specialty paediatric / neonatal expertise as per <u>PD2013 049 Recognition and management of patients who are clinically deteriorating.</u>

**Urgent medical review** - a bedside review by the most senior medical officer or specialist paediatrician responsible, as per local paediatric-specific CERS. Initial consultation may be by telephone to enable treatment to commence, however, a physical examination should occur as soon as possible.

**Medical review** - a bedside review should occur within six hours by the most senior medical officer responsible. Initial consultation may be by telephone to enable treatment to commence, however, a physical examination should occur within this time frame.



#### 1.2.2 Jaundice

**Jaundice** - a yellowish staining of the skin and sclera.

**Physiological jaundice** - a common condition caused by the breakdown of fetal red blood cells combined with an immature liver that cannot effectively metabolise bilirubin and prepare it for excretion. Usually presents on day 3, peaks between days 5 to 7 and has resolved by 14 days of age<sup>2</sup>.

**Pathological jaundice** - when non-physiological causes result in jaundice of the neonate, most commonly due to blood group incompatibility (ABO or rhesus blood group incompatibility). Other causes include sepsis, bruising, metabolic disorders or obstruction<sup>2</sup>. High conjugated fraction (> 20 micromol per litre (micromol/L) or > 20% of total SBR) is always pathological and should be investigated urgently.

**Prolonged jaundice** - jaundice persisting beyond 14 days of age in a term neonate and 21 days in a preterm neonate. It is more common in breast fed neonates<sup>2</sup>.

**Hyperbilirubinaemia** - SBR measurement above that which requires treatment<sup>2</sup> to prevent encephalopathy and kernicterus.

**Severe hyperbilirubinaemia** - SBR measurement above exchange transfusion threshold line.

#### 1.2.3 Bilirubin

**Bilirubin** - yellow pigment created in the body during the normal breakdown of red blood cells which leads to the production of unconjugated bilirubin<sup>2</sup>.

**Unconjugated bilirubin** - the lipid-soluble form of bilirubin that binds to albumin and metabolised in the liver to form conjugated bilirubin<sup>2</sup>. Unconjugated bilirubin can cross the blood brain barrier in neonates and is potentially toxic to neural tissue. The measurement at which unconjugated bilirubin becomes toxic varies between neonates but certain risk factors increase the risk of acute bilirubin encephalopathy<sup>2</sup>.

**Conjugated bilirubin** - unconjugated bilirubin is taken up by the liver cells and conjugated to form water-soluble bilirubin diglucuronide. This then passes through the gut and is excreted in the stools. Bilirubin can be reabsorbed from the stools remaining in the gut<sup>2</sup>. High conjugated fraction (> 20 micromol/L or > 20% of total SBR) is always pathological and should be investigated urgently.

**Serum Bilirubin (SBR)** - the measurement of the total conjugated and unconjugated bilirubin in the blood.

### 1.2.4 Bilirubin encephalopathy and kernicterus

**Bilirubin encephalopathy** - short or long term neurologic dysfunction caused by toxic unconjugated bilirubin crossing the blood-brain barrier. Signs and symptoms include: lethargy; hypotonia; poor suck; irritability; apnoea; abnormal posture (opisthotonos - rigid with back arched and retrocollis - head tilted backwards); high pitched cry; seizures and coma.

**Kernicterus** - the yellow staining caused by bilirubin deposited in the globus pallidus of the deep grey matter of the brain. It is a rare condition<sup>2</sup>.



### 1.2.5 Phototherapy

**Phototherapy** - light energy used to convert bilirubin in the skin to a water soluble isomer that is excreted.

**Fibre optic phototherapy** - comprises a light generator, a fibre optic cable carrying light to a flexible light pad or blanket placed under or around the neonate.

**Light emitting diode (LED) phototherapy** - emits high intensity light in a narrow wavelength spectrum and produces minimal heat<sup>4</sup>.

**BiliBed** - fluorescent tube, single light source positioned below the neonate in the cot while the neonate is wrapped in a therapy suit that exposes the back of the neonate to the light source (not recommended by the manufacturers for use in humidicribs).

**Conventional phototherapy** - a single fluorescent blue light unit positioned above the neonate<sup>2</sup>.

**Single light phototherapy** (15 μW/nm/cm² to 30 μW/nm/cm²) - one unit of phototherapy light; either fluorescent, LED or fibre optic phototherapy.

**Multiple light phototherapy** (>  $30 \mu W/nm/cm^2$ ) - more than one light source used simultaneously.



# 2. IDENTIFICATION, MEASUREMENT AND INVESTIGATION OF NEONATAL JAUNDICE

### 2.1 Identification and assessment

### 2.1.1 Universal surveillance and timing of visual assessments

Universal surveillance and timing of visual assessments (see Flowchart 1), is the responsibility of all clinical staff and includes:

- Identification at birth of neonates with risk factors for neonatal jaundice (see Table 1) who require planned, increased visual assessment, at least 3 times per day (recommended) for the first 24 to 48 hours. Visual assessment includes assessment of blanched skin (useful in all skin tones)<sup>2</sup> sclera and gums
- Regular visual assessment from birth of all neonates for jaundice at least daily as part of the newborn wellbeing assessment to identify neonates who become jaundiced
- Neonates who are jaundiced should be monitored for adequacy of oral intake.
   Providing lactation advice and support of breast feeding mothers is an important risk reduction strategy for hyperbilirubinaemia
- Neonates who are jaundiced ≤ 24 hours of age should have bilirubin measurement and urgent medical review in line with <u>PD2013\_049\_Recognition</u> <u>and management of patients who are clinically deteriorating</u> and the SNOC. Concerns should escalated as per local CERS
- Neonates identified as jaundiced ≥ 24 hours of age should have a medical review (see section 2.2 Measurement) and an SBR if their transcutaneous bilirubinometer (TcB) reading is ≥ 250 micromol/L or if staff or parents are concerned
- As neonatal jaundice usually peaks between 5 and 7 days of age, it is advised that all neonates are assessed regularly during this period. For those neonates discharged less than 3 days of age, guidance for timing of follow-up of neonates, with or without risk factors, is provided in Table 11 (see section 5.1 <u>Timing of follow-up</u>)
- Neonates with prolonged jaundice > 14 days of age require urgent medical review and bilirubin measurement
- All jaundiced neonates should be monitored for the presence of signs suggestive of early bilirubin encephalopathy.



# 2.1.2 Risk factors and causes of neonatal jaundice

Table 1: Risk Factors and Causes of	Neonatal Jaundice		
Jaundice < 24 hours of age - Suspect haemolysis until proven otherwise			
Jaundice due to haemolysis	Immune - e.g. ABO blood group incompatibility, Rhesus disease, Kell, Duffy, anti-E (see section 3.4) Non-immune - e.g. Glucose-6-phosphate dehydrogenase deficiency (G6PD)		
Individual neonatal risk factors	<ul> <li>Prematurity</li> <li>Asphyxia</li> <li>Apgar &lt; 7 at 5 minutes, acidosis pH &lt; 7 or base excess ≤ 12 mEq/L</li> <li>Low serum albumin &lt; 30g/L</li> <li>Sepsis or congenital infections</li> <li>Maternal diabetes</li> <li>Cephalohaematoma / bruising</li> <li>History of sibling who was jaundiced as a neonate</li> <li>G6PD risk - family history or with exposure to trigger (see below)</li> </ul>		
Jaundice in the first 7 days of age - I	Investigate high SBR and possible underlying causes		
Typical neonatal jaundice	<ul> <li>Physiological jaundice</li> <li>Neonates with delayed (versus early) cord clamping, may have a higher haematocrit and therefore an increased incidence of jaundice requiring phototherapy<sup>5</sup></li> </ul>		
Breast feeding jaundice	<ul> <li>Early breast feeding jaundice. Develops within 2 to 4 days of birth and is thought to relate to infrequent breast feeding with a limited fluid intake</li> <li>Possible increased reabsorption of bilirubin from the bowel</li> </ul>		
Breakdown of extravasated blood	<ul><li>Significant bruising</li><li>Cephalohaematoma</li><li>Intracranial haemorrhage</li></ul>		
Increased enterohepatic circulation	Delayed passage of stool or gut obstruction		
Red cell membrane defects	<ul><li>Spherocytosis</li><li>Elliptocytosis</li></ul>		
Prolonged jaundice after 2 weeks of	age - should be investigated measuring total and conjugated SBR.		
Unconjugated hyperbilirubinaemia	<ul> <li>Breast milk jaundice (rare - can last up to 12 weeks)</li> <li>Sepsis</li> <li>Hypothyroidism (thyroid agenesis/dysplasia or hypopituitarism)</li> <li>G6PD</li> <li>Rarely, inborn deficiency of UDP-glucuronyltransferase enzyme in Crigler-Najjar Syndrome and related disorders</li> </ul>		
Conjugated hyperbilirubinaemia	<ul> <li>Idiopathic neonatal hepatitis</li> <li>Infections (Hepatitis B, sepsis, non-bacterial congenital infection)</li> <li>Congenital biliary tract obstruction (biliary atresia, choledochal cyst, bile duct stenosis)</li> <li>Metabolic disorders (galactosaemia, hereditary fructose intolerance, Alpha-1 antitrypsin deficiency, tyrosinemia, glycogen storage disease type IV, hypothyroidism)</li> </ul>		
Onset at any time			
Secondary to sepsis	<ul> <li>Can occur following onset of sepsis (both early and late onset)</li> <li>May have both raised unconjugated and conjugated bilirubin components of SBR</li> </ul>		
Glucose-6-phosphate dehydrogenase deficiency (G6PD)	Can occur any time following exposure to a trigger such as naphthalene (moth balls), fava beans, sepsis and hypothermia.		



#### 2.2 Measurement

### 2.2.1 Non-invasive transcutaneous bilirubinometer (TcB) measurement

The main goal of TcB measurement is to identify more accurately those jaundiced neonates who need an SBR<sup>6,7</sup> and to reduce the number of invasive tests required. In the first instance a TcB measurement should be used if possible for the well neonate who is jaundiced at:

- ≥ 35<sup>+0</sup> weeks gestation at birth and
- ≥ 24 hours of age.

#### 2.2.2 Serum bilirubin measurement

SBR measurement remains the 'gold standard' for jaundice treatment decisions<sup>8</sup>.

An SBR should be measured if:

- A TcB is not available<sup>2</sup>
- The TcB measurement is ≥ 250 micromol/L, or the result is on, or within 20 micromol/L of the phototherapy threshold line for gestation at birth (see section 2.2.3 Plotting bilirubin measurement and assessment for treatment)
- The neonate is:
  - Unwell
  - < 35 weeks gestation at birth
    </p>
  - < 24 hours of age (see section 2.3 Investigation)</p>
  - Undergoing phototherapy or has undergone phototherapy (there is insufficient evidence to recommend the use of TcB after phototherapy)<sup>9</sup>.

It is essential to follow up bilirubin results in a timely way or ensure clinical handover of requirement to follow up.

Both venous and capillary total SBR results should be considered equivalent measures<sup>1,2</sup>. The total SBR should be used to determine appropriate treatment<sup>1,2</sup> rather than the unconjugated fraction of bilirubin.

If the SBR is < 50 micromol/L below the phototherapy treatment threshold line repeat the SBR within 12 to 24 hours.

### 2.2.3 Plotting bilirubin measurement and assessment for treatment

Accurate data entry of the TcB or total SBR measurement, plotted on **the appropriate** Neonatal Jaundice Treatment Threshold Graph for gestational age at birth<sup>10</sup> (see <u>attachments 1-7</u>) is essential to:

- Monitor the progression of neonatal jaundice
- Identify hyperbilirubinaemia and support decision to treat
- Monitor the effect of treatment and inform clinical decision making.

The appropriate jaundice treatment threshold graph for gestational age at birth should not be changed for the corrected gestation.



### Flowchart 1: Identification and Investigation of Neonatal Jaundice

#### Universal surveillance of all neonates comprises: Identification of risk factors that increase the risk of hyperbilirubinaemia in individual neonates Jaundice within first 24 hours of age Sepsis A sibling who was jaundiced as a neonate · Cephalohaematoma / bruising ABO blood type incompatibility or Rh incompatibility • East Asian or Mediterranean descent Non-optimal sucking at the breast Preterm G6PD deficiency or other red cell abnormalities • Diabetes (maternal) Regular visual assessment of skin colour for the first 4 days of life for all neonates • Blanched skin (useful in all skin tones) • Sclera Gums Risk factors? AND / Visible jaundice? **OR** YES ΝO NO YES Regular visual assessment for the first 4 days of age More frequent visual assessment should occur in the first 24 to 48 hours of age Monitor and record assessments (SNOC if in hospital) Term neonate > 14 days of age or preterm beyond 21 days Well neonate of age should have Neonate < 24 hours ≥ 24 hours of age and urgent medical review by a senior medical officer or specialist of age ≥ 35 weeks gestation? paediatrician for possible signs of obstructive jaundice NO YES Initiate urgent medical **SBR Pathway TcB Pathway** review Measure SBR · Plot SBR result Initiate <u>medical review</u> TcB should be used if available. As per urgent · Measure SBR as requested Plot on treatment threshold graph medical review Plot SBR result Document in clinical record Consider starting · Document in clinical record If no TCB available an SBR should phototherapy at a be taken and follow SBR pathway lower SBR if the neonate is unwell or has risk factors for As per medical review If SBR > 50 jaundice Consider starting If SBR < 50 micromol/L Consider (see phototherapy at a lower micromol/L below 2.3.3) Additional SBR, if the neonate is below If TcB If TcB treatment investigations unwell or has risk factors treatment measurement is measurement threshold line Measure SBR for jaundice threshold line ≥ 250 micromol/L < 250 continue 6 hourly until SBR is SBR at or above the repeat SBR in micromol/L or or the both below visual treatment threshold line 12 - 24 hours measurement is below the assessment treatment threshold commence phototherapy ≤ 20 micromol/L treatment line and stable or · Additional investigations below the threshold line falling then Measure SBR treatment for gestation 12-24 hourly for the After 6 hours if SBR is threshold line for continue duration of treatment stable or falling then gestation follow visual Consider transfer to 12 - 24 hourly for SBR pathway assessment a higher level facility duration of treatment

A neonate who has severe hyperbilirubinaemia or whose SBR is rapidly rising (> 8.5 micromol/L/hour) or who has any signs and symptoms of bilirubin encephalopathy is considered a medical emergency and should have an <u>urgent medical review</u> as per local paediatric-specific CERS.



# 2.3 Investigation

### 2.3.1 Urgent investigation of the neonate with visible jaundice < 24 hours of age

Table 2 outlines steps to identify and investigate the neonate with visible jaundice < 24 hours of age (see also Flowchart 1: Identification and Investigation of Neonatal Jaundice).

Table	Table 2: Urgent Investigation of the Neonate with Visible Jaundice < 24 Hours of Age			
Step	Action			
1	Initiate <u>urgent medical review</u> by the most senior medical officer or specialist paediatrician responsible as per local CERS protocol. The initial consultation may be by telephone to order investigations and enable treatment to commence, however, a medical review at the bedside should occur as soon as possible			
2	Measure and plot the SBR as per section 2.2 Measurement <sup>2</sup> and Flowchart 1: Identification and Investigation of Neonatal Jaundice			
3	Measure and record any Additional Investigations recommended in section 2.3.3			
4	Commence phototherapy			
5	Measure the SBR at least every 6 hours until the SBR is both <sup>2</sup>			
	<ul><li>Below the treatment threshold and</li><li>Stable and / or falling</li></ul>			
6	Measure and record SBR every 12 - 24 hours for the duration of phototherapy			
	Consider starting phototherapy at a lower SBR, if the neonate is < 24 hours of age, has risk factors of neonatal jaundice or is unwell			

### 2.3.2 Investigation of a neonate with visible jaundice ≥ 24 hours of age

Table 3 outlines the steps to identify and investigate the neonate with visible jaundice ≥ 24 hours of age (see also Flowchart 1 Identification and Investigation of Neonatal Jaundice).

	Table 3: Investigation of a Neonate with Visible Jaundice ≥ 24 Hours of Age			
Step	Action			
1	Where possible and if appropriate, the non-invasive TcB measurement should be used to determine if an SBR is required. If a TcB is not available an SBR should be taken <sup>2</sup>			
2	Measure and plot the neonatal bilirubin measurement as per section 2.2 Measurement <sup>2</sup> and Flowchart 1: Identification and Investigation of Neonatal Jaundice			
	If total SBR is at or above phototherapy treatment threshold	If total SBR is rapidly rising	If total SBR is at or above exchange transfusion threshold	
3	Initiate a <u>medical review</u> Initiate <u>urgent medical review</u> as per local CERS protocol			
	NOTE: Initial consultation may be by telephone to enable phototherapy treatment to commence, however, a bedside medical review should occur within 6 hours	NOTE: Initial consultation may be by telephone to enable phototherapy treatment to commence, however, a bedside medical review should occur as soon as possible		
4	Commence phototherapy and arrange Additional Investigations as recommended in section 2.3.3			
5	Measure the SBR at 6 hours to ensure the SBR is stable or falling			
6	When SBR is stable or falling, measure and record SBR every 12 - 24 hours for the duration of phototherapy			



# 2.3.3 Additional investigations to be considered in particular clinical situations

Table 4 outlines additional investigations to be considered in particular clinical situations

Table 4 Additional Investigations		
Clinical Feature	Investigation	
Neonate of Rhesus negative mother	Blood group Direct Antiglobulin Test (DAT) An immediate SBR is required if the DAT is positive and the SBR is unknown	
Neonate with jaundice within the first 24 hours of age OR  Neonate with a rapidly rising total SBR (> 8.5 micromol/L/hour)  OR  Neonate with a total SBR above the phototherapy threshold	Full blood count (FBC) and film with reticulocyte count Blood group DAT Septic screen including blood and urine culture and sensitivity if sepsis suspected A G6PD screen if There is a family history This is a male neonate from a high risk ethnic origin/geographic area; African, Asian, Mediterranean and Middle Eastern <sup>2</sup> descent	
The maternal blood group should	d be known and considered with the above investigations	
Neonate with a total SBR approaching exchange transfusion thresholds	<ul><li>Serum albumin level</li><li>Liver function tests</li><li>Conjugated bilirubin</li></ul>	
medical review by the most senior i	ted bilirubin > 20 micromol/L or > 20% of the total SBR <sup>1</sup> , should have a medical officer or specialist paediatrician <sup>11</sup> (same day) and not	

discharged from hospital unless the cause is identified and treatment commenced.

### 3. MANAGEMENT AND TREATMENT OF NEONATAL JAUNDICE

The decision to treat jaundice is based on:

- The bilirubin measurement plotted on the appropriate graph for gestational age and the proximity to:
  - The phototherapy treatment threshold line or
  - The exchange transfusion treatment threshold line
- The age at recognition of jaundice
- The clinical condition of the of the neonate
- Identified risk factors for jaundice.

### Consideration should be given to starting phototherapy at a lower SBR if the neonate is < 24 hours of age, has risk factors for neonatal jaundice or is unwell

Treatment options will vary according to the services available at each facility. Treatment should encompass general management by a clinician skilled in neonatal care to assess the neonate, monitor the effectiveness of phototherapy (see Section 3.1.3) and treat any underlying illnesses that may be causing jaundice e.g. sepsis. If appropriate treatment is not available locally, transfer may be required.

Consultation, escalation and / or transfer to a higher level facility may be required. In these circumstances clinicians should:

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- Follow local escalation processes in the first instance. This may involve contacting a specialist paediatrician and / or neonatologist in the Maternity and Neonatal Tiered Network with an appropriate service capability level
- Contact a neonatologist urgently either directly or via NETS (1300 36 2500) if the jaundice treatment required is an exchange transfusion.

Fully breast fed neonates with physiological jaundice can develop hyperbilirubinaemia associated with poor oral intake and/or dehydration<sup>1</sup>. By day 3 of life, 5-10% of fully breast fed neonates will lose 10% or more of their birth weight<sup>1</sup>. Ensuring adequate oral intake and appropriate lactation advice and support is therefore essential. It is preferable that expressed breast milk is given if additional feeds are required.

In some cases enteral or intravenous rehydration may be required for neonates under phototherapy with weight loss > 10% of birth weight and dehydration - see <u>GL2015\_008</u> Standards for Paediatric Intravenous Fluids

Sunlight is not a treatment option for jaundice.

### 3.1 Phototherapy

Phototherapy is the first line of treatment for neonatal jaundice and effectively reduces the SBR in most neonates. Clinical response to phototherapy depends on:

- The cause and severity of the hyperbilirubinaemia 12
- The balance between the neonate's rate of bilirubin production, enterohepatic circulation, bilirubin elimination and degree of tissue bilirubin deposition
- The rate of the photochemical reactions of bilirubin
- The skin surface area exposed to phototherapy
- Phototherapy device efficacy<sup>13,14</sup> which can be influenced by multiple factors see Appendix B: Maximising Phototherapy Efficacy.

Each facility should have written information and established processes, in line with manufacturer's recommendations (see Section 3.1.3 <u>Effectiveness of phototherapy</u> and Appendix B: <u>Maximising Phototherapy</u> Efficacy) to guide:

- Clinical staff to set up, use and maximise the effectiveness of phototherapy
- Biomedical departments to measure light intensity and maintain the effectiveness of phototherapy equipment.

**NOTE:** At the time of this guideline publication there was no available high-quality evidence to support or refute the use of home phototherapy for uncomplicated physiological neonatal jaundice<sup>13,15</sup>.

### 3.1.1 Contraindications to phototherapy

Contraindications for phototherapy include:

- Neonates with congenital porphyria
- Family history of porphyria
- Concurrent treatment with photosensitising drugs<sup>12,13</sup>.



### 3.1.2 Potential adverse effects of phototherapy

Concerns regarding possible long term effects on the reproductive system from continuous phototherapy have been raised but have not been substantiated in animal studies<sup>13</sup>. Prolonged phototherapy is associated with increased<sup>13</sup>:

- Oxidative stress
- Lipid peroxidation
- Riboflavin deficiency
- Retinal damage (if eye protection recommendations are not followed)
- Eye trauma from eye protecting covers.

Recent clinical reports of other adverse outcomes have yet to be validated but potentially include skin changes<sup>2</sup>. Neonates who are not within phototherapy range should therefore not be treated.

# **3.1.3 Effectiveness of phototherapy** (See <u>Appendix B: Maximising Phototherapy</u> <u>Efficacy</u>)

A review of the literature found that when used and maintained according to the manufacturer's instructions and when the light intensity is adequate:

- All modes of phototherapy are safe and effective as first-line medical treatment of hyperbilirubinaemia in preterm neonates<sup>2</sup>
- Conventional modes of phototherapy have been recommended for term neonates<sup>2</sup> however,
- Emerging evidence supports the use of LED phototherapy for term and near term neonates<sup>16</sup>
- The effect of fibre optic devices may be limited by the size of the device and the surface area of skin exposed<sup>2</sup>.

It is essential to monitor the effectiveness of phototherapy (see section 2.2 <u>Measurement</u>) as some neonates, despite treatment, may require further medical intervention<sup>2</sup>.

### 3.1.4 When to use single light phototherapy

Table 5 outlines appropriate clinical circumstances to use single light phototherapy.

Table 5: When to use Single Light Phototherapy (15μW/nm/cm² to 30μW/nm/cm²)

Use a single phototherapy light when

- The total SBR is at or above the phototherapy threshold as plotted on the appropriate <u>Jaundice</u> <u>Treatment Threshold Graph for gestational age (See Attachment 1-7)</u>
- The SBR is not rising rapidly
- The SBR is more than 50 micromol/L below the exchange transfusion threshold

Within 6 hours of commencing phototherapy the SBR should have decreased by 34 micromol/L in both the term and preterm neonate<sup>13</sup>



# 3.1.5 Clinical care of the neonate under single light phototherapy

Table 6 outlines clinical care considerations for neonates undergoing single light phototherapy.

Table 6 Clinical Care of the Neonate Under Single Light Phototherapy			
Step	Action		
Parent involvement	Parents are given clear information and are included in treatment and care planning decisions as well as care giving <sup>2,17</sup>		
Location	Postnatal ward, special care nursery/non tertiary facility or Neonatal Intensive Care Unit (NICU)		
Assessments	<ul> <li>Document input / output - loose stools are common (dark urine and or light stools may indicate obstructive causes of jaundice)</li> <li>Bare weigh as necessary</li> <li>Daily assessment of neonatal wellbeing should include assessment of skin integrity</li> <li>Observe and record assessments 3-6 hourly on the SNOC and in clinical record</li> </ul>		
Monitoring	Use a cardio respiratory monitor or continuous oximetry when the neonate  Is cared for in a humidicrib  Is cared for in a position other than supine  Is receiving blue light phototherapy Record appropriately		
Temperature	<ul> <li>Hourly for the first 3 to 4 hours and monitor and record on the SNOC</li> <li>Then measure 3 - 6 hourly</li> <li>Provide care in an environment that will maximise thermal stability and minimise energy expenditure taking into consideration the light source in use (e.g. LED phototherapy lights produce minimal heat). Consider using a humidicrib<sup>2</sup>.</li> </ul>		
SBR measurement	<ul> <li>Repeat SBR 6 hours after commencement of phototherapy (the total SBR should be decreased by 34 micromol/L in this time period for both term and preterm neonates<sup>13</sup>)</li> <li>Subsequent SBRs in line with neonatal age at recognition of jaundice see Table 2: <u>Urgent investigation of the neonate with visible jaundice &lt; 24 hours of age</u> or Table 3: <u>Investigation of the neonate with visible jaundice &gt; 24 hours of age</u></li> <li>If SBR is rapidly rising (&gt; 8.5 mmol/L per hour) or continuing to rise under single light phototherapy consider changing to multiple light sources and earlier repeat of SBR</li> <li>Repeat SBR 24 hours after phototherapy ceases</li> </ul>		
Feeding and hydration	<ul> <li>Demand breast feeding (maximum of 4 hour between feeds)</li> <li>If formula feeding, recommend 3 - 4 hourly feeding</li> <li>Phototherapy may be interrupted for feeding</li> </ul>		
Positioning	Place the neonate in a supine position unless other clinical conditions prevent this		
Skin care	Lotions or lubricants should not be used		
Eye Care	<ul> <li>Eye protective mask/patches are mandatory for conventional light therapy (check placement)</li> <li>If the neonate's eyes will not be directly exposed to BiliBed or fibre optic treatment lights eye protection is not required</li> <li>Remove eye masks at feeds and check for eye discharge and conjunctivitis.</li> </ul>		
Surface area exposed	<ul> <li>Position phototherapy device according to manufacturer's instructions</li> <li>Remove clothing but leave the nappy on for most single light and BiliBed phototherapy</li> <li>Some fibre optic devices may be positioned next to the neonates skin under the singlet</li> </ul> BiliBed in an hymidicrib (see manufacturer's recommendations)		

Do **NOT** use a BiliBed in an humidicrib (see manufacturer's recommendations)

Do NOT turn the humidicrib off during phototherapy (see manufacturer's recommendations)

Plastic heat shields are no longer recommended for use



# 3.1.6 Multiple light phototherapy

Evidence shows that multiple light phototherapy is more effective than conventional or single light phototherapy<sup>2</sup>, and that it may reduce the need for exchange transfusion and possibly reduce the severity of bilirubin neurotoxicity<sup>13</sup>. This approach consists of delivering high levels of irradiance to the maximum skin surface (see Appendix B Maximising Phototherapy efficacy). The surface area exposed can be increased by using additional light banks and by combining devices such as a conventional phototherapy light bank plus fibre optic pads or a light emitting diode (LED) devices<sup>13</sup>.

There is no evidence regarding the efficacy of intermittent phototherapy when multiple light phototherapy is required. Treatment should therefore not be interrupted for oral feeds<sup>2</sup> see Table 8 Clinical care of neonate under multiple light phototherapy.

Table 7 outlines the clinical circumstances in which to use multiple light phototherapy

### Table 7: When to use Multiple Light Phototherapy (> 30µW/nm/cm²)

Initiate multiple light phototherapy to treat all neonates if any of the following applies

- The SBR is rising rapidly (> 8.5 micromol/L per hour)
- The SBR is < 50 micromol/L below the exchange transfusion treatment threshold line
- The SBR fails to respond to single light phototherapy (that is, the SBR is static, continues to rise, within 6 hours of starting single light phototherapy)
- A rapid reduction in SBR is required

Multiple light phototherapy will usually cause a high SBR to fall when due to physiological jaundice. If the SBR falls during multiple light phototherapy to 50 micromol/L below the threshold for which exchange transfusion is indicated, a step down to single light phototherapy should be considered.

### 3.1.7 Clinical care of the neonate under multiple light phototherapy

Table 8 details the clinical care for neonates undergoing multiple light phototherapy

Table 8: Clinical Care of Neonate when Undergoing Multiple Light Phototherapy Treatment			
Step	Action		
Parent involvement	As for parents of neonates under single light phototherapy		
Location	Special care nursery / non tertiary facility or NICU		
Assessments	As for neonates under single light phototherapy		
	Assess for the presence of signs suggestive of early bilirubin encephalopathy		
Monitoring	As for neonates under single light phototherapy		
Temperature	As for neonates under single light phototherapy		
SBR measurement	<ul> <li>Repeat SBR 6 hours after commencement of phototherapy (the total SBR should be decreased by 34 micromol/L in this time period for both term and preterm neonates<sup>13</sup>)</li> </ul>		
	Subsequent SBRs in line with neonatal age at recognition of jaundice see		
	Table 2: <u>Urgent investigation of the neonate with visible jaundice &lt; 24 hours of age</u> Table 3: <u>Investigation of the neonate with visible jaundice &gt; 24 hours of age</u>		
<ul> <li>If the SBR is rapidly rising (&gt; 8.5mmol/L per hour) consider early repeat of S</li> </ul>			
	Repeat SBR 24 hours after phototherapy ceases		
Feeding and	Phototherapy should not be interrupted for breast / bottle feeding		
Hydration	Consider administration of intravenous or enteral feeds		
-	Expressed breast milk is the fluid of choice if additional fluids are required		
Positioning	As for neonates under single light phototherapy		
	If only one side of the neonate is exposed to phototherapy consider position change		
011.	every 3-4 hours to maximise skin exposure		
Skin care	Lotions or lubricants should not be used		



Eye care	As for single light phototherapy plus		
	Eye protective mask/patches are mandatory for multiple light phototherapy		
Surface area exposed	Position phototherapy device according to manufacturer's instructions  • Maximise skin surface area exposed to phototherapy (see Appendix B)  ° Remove head covers, clothes and nappy  ° Ensure phototherapy is not obstructed from reaching the neonate's skin, consider		
Diagtic baset	removal of tape, reposition chest leads.  shields are no longer recommended for use		

### 3.1.8 Ceasing phototherapy

The suggested total SBR measurement for ceasing phototherapy is  $\geq$  50 micromol/L below the phototherapy treatment line on the appropriate Jaundice Treatment Threshold Graph for gestational age at birth<sup>1,2</sup> (see <u>Attachments 1-7</u>).

A rebound in total SBR can occur after phototherapy is discontinued<sup>2</sup>. A clinically significant rebound is more likely in neonates who are < 37 weeks gestation, have known haemolytic disease or who have identified pathology. Check for rebound of hyperbilirubin by repeat SBR at 12 to 24 hours of age.

Neonates who do not have these risk factors do not need to delay discharge to assess for a rebound in total SBR. Instead consider follow-up SBR measurement within 12 to 24 hours after discharge (See Section 5: <u>Discharge planning</u>).

# 3.2 Adjunct therapy

The only adjunct therapy supported by evidence is the use of intravenous immunoglobulin in cases of Rhesus or ABO haemolytic disease <sup>2,16</sup>.

Pharmacologic options should always be discussed with a neonatologist prior to treatment as per *PD2010\_69 NSW Critical Care Tertiary Referral Networks (Perinatal)*.

# 3.2.1 Intravenous immunoglobulin (IVIG)

There is some evidence that intravenous immunoglobulin (IVIG) will reduce the need for exchange transfusions in neonates with immune haemolytic jaundice<sup>18</sup>. Consider using IVIG (500 mg/kg over 4 hours) as an adjunct to multiple light phototherapy in isoimmunised haemolytic disease when the SBR continues to rise by > 8.5 micromol/L per hour<sup>2</sup>.

### 3.2.2 Other agents

The use of albumin is not currently recommended as an intervention for jaundice treatment. There is insufficient evidence to support its routine use as an adjunct therapy prior to exchange transfusion <sup>16,19</sup>.

Agents such as metalloporphyrins, gammaglobulins, drugs (phenobarbitol, clofibrate, cholestyramine), agar, charcoal, suppositories, other rectal modes of treatment; and complementary or alternative medicines (e.g. Chinese herbal remedies such as Yinchen) are not recommended for the treatment of neonatal hyperbilirubinaemia<sup>2,16</sup>.



# 3.3 Exchange transfusion for severe hyperbilirubinaemia

A neonate who has severe hyperbilirubinaemia or whose SBR is rapidly rising or who has signs and symptoms of bilirubin encephalopathy is considered a medical emergency and should have an urgent medical review by the most senior medical officer or specialist paediatrician as per the local paediatric-specific CERS protocol.

### 3.3.1 When to undertake an exchange transfusion

Exchange transfusion may be appropriate in the following clinical circumstances and care should be escalated accordingly where:

- The total SBR is above the exchange transfusion threshold when plotted on the Jaundice Treatment Threshold Graph for gestational age see Attachments 1-7
- SBR is rising > 8.5 micromol/L per hour despite multiple light phototherapy in a neonate with known haemolysis OR
- There are signs of bilirubin encephalopathy (see section <u>1.2.4</u>).

### 3.3.2 Where to undertake an exchange transfusion

Table 9 provides information in relation to the most appropriate place to undertake an exchange transfusion and lists important considerations.

### **Table 9: Where to Undertake an Exchange Transfusion**

Exchange transfusions should be carried out at a level 5 or 6 NICU (see <u>GL2016\_018 NSW Maternity</u> and Neonatal Service Capability Framework.

When anticipated, antenatal referral is recommended for care planning including the appropriate place of birth.

If the neonate presents at the Emergency Department or at lower level neonatal facility consult a neonatologist within the Tiered Maternity and Neonatal Network <u>PD2010 069 NSW Critical Care Tertiary Referral Networks (Perinatal)</u> regarding treatment.

Consider the need for urgent transfer to a level 5 or 6 NICU facility if the baby is systemically unwell and contact NETS (Newborn and Paediatric Emergency Transport Service) 1300 36 2500.

A systemically unwell neonate	The risks of exchange transfusion are much higher in a systemically unwell neonate Consider transfer to a NICU in this situation
Staff availability	A minimum of two staff members (nurse/midwife and doctor) are required to remain at the bedside for the duration of the exchange transfusion
Staff capacity	A clinician with the skills to perform an exchange transfusion should be available.  This includes the capacity to insert an umbilical venous catheter (UVC). If the skills are not available locally, the regional paediatric- specialist may be able to attend
Availability of blood	As exchange transfusion is a medical emergency, low titre O negative blood is used but only in consultation with a neonatologist



### 3.3.3 Clinical care of the neonate undergoing exchange transfusion

Table 10 outlines clinical care required by the neonate undergoing exchange transfusion.

**Table 10: Clinical Care of Neonate Undergoing Exchange Transfusion** 

**NOTE:** Multiple light phototherapy treatment (if not in progress) should be commenced immediately and continue throughout the exchange transfusion.

#### Clinical care

#### **Tertiary facility (NICU)**

Complete all clinical care and assessments as per local guidelines

#### Non tertiary facility

Follow Neonatal Exchange Transfusion in a Non-Tertiary Hospital - How to guide available on the NSW Pregnancy and Newborn Services Network website <a href="https://www.psn.org.au">www.psn.org.au</a> for all recommended clinical care and assessments during exchange transfusion

The insertion of a UVC remains a suitable option for up to 7-10 days post birth<sup>20</sup> If insertion of a UVC is not possible then a peripheral line will need to be inserted

# Post exchange transfusion

- Maintain multiple light or high intensity phototherapy
- Measure SBR within 2 hours of completion of the exchange transfusion
- Continue under the care of a neonatal specialist.

### **Tertiary facility (NICU)**

Complete all clinical care and assessments as per local guidelines post exchange transfusion.

#### Non tertiary facility

Follow Neonatal Exchange Transfusion in a Non-Tertiary Hospital - How to guide available at the NSW Pregnancy and newborn Services Network website <a href="https://www.psn.org.au">www.psn.org.au</a> for all recommended clinical care post exchange transfusion

# 3.4 Management of neonates with known in utero rhesus sensitisation

All neonates with known isoimmunisation prior to birth should be birthed at a tertiary facility with an NICU. Transfer if birth has occurred at a lower level facility.

The following investigations should be completed at birth on cord blood:

- Blood group
- FBC
- DAT
- SBR
- If affected, have blood ready for exchange transfusion.

#### 3.4.1 Without in-utero transfusion

For neonates who have <u>not</u> received an in-utero transfusion the threshold for a rapidly rising total SBR remains > 8.5 micromol/L per hour.

If initial SBR result is  $\geq$  80 micromol/L commence single light phototherapy. Consider multiple light phototherapy, immunoglobulin and early exchange transfusion<sup>1</sup>.



#### 3.4.2 With in-utero transfusion

For neonates who have had an in-utero transfusion the criteria for an exchange transfusion should be decided on a case by case basis by a neonatologist experienced in the management of neonates with known Rhesus disease.

### 4. PROLONGED JAUNDICE

Jaundice persisting beyond the first 14 days of life in a term neonate or beyond 21 days of life in a preterm neonate should have an urgent medical review by the most senior medical or specialist paediatrician for signs of obstructive jaundice.

The initial investigations should include:

- Assess stool colour look for acholic pale chalky stools
- Assess urine look for dark urine that stains the nappy
- Complete the following tests:
  - Total bilirubin
    - Conjugated bilirubin
  - FBC to exclude a red cell structural problem (e.g. spherocytosis)
  - Blood group (if not already done)
  - Confirm maternal blood group
  - DAT (if not already done) and interpret the result of the DAT taking account of the strength of reaction, and whether or not the mother received prophylactic anti-D immunoglobulin during pregnancy
  - Urine culture
  - Thyroid function tests including TSH and Free T4.

Conjugated bilirubin < 20 micromol/L is usually benign breast milk jaundice, however specific investigations may be considered e.g. metabolic screen; G6PD screen

Conjugated bilirubin > 20 micromol/L or > 20% of the total SBR is always pathological and should be investigated for intra-hepatic and obstructive causes.

Delay in diagnosis of biliary atresia is an important prognostic factor. Early discussion with a gastroenterologist is essential. Where local services are not available, it is important to refer the neonate to a tertiary paediatric critical care centre able to investigate and in particular, to exclude biliary atresia.

See NSW Health Policy Directive <u>PD2010\_69 NSW Critical Care Tertiary Referral Networks (Perinatal)</u> and <u>PD2010\_030 Critical Care Tertiary Referral Networks (Paediatrics)</u>



### 5. DISCHARGE PLANNING

Hyperbilirubinaemia is a potentially preventable cause of 35% of early readmissions of neonates, with higher rates among late preterm neonates<sup>21,22</sup>.

Individual evaluation of each mother-neonate dyad to determine the optimal time of discharge and the follow-up required is essential. LHDs are responsible for the development of local process for follow up assessment of the neonate including providing the location of that service and a process for escalation of concerns about the ongoing care of the neonate with jaundice. Midwives, early childhood nurses and general practitioners should be aware that jaundice in the term neonate peaks between 5 to 7 days of age and if discharged prior to this time may require further assessment after discharge.

A recent study in NSW found that birth at 37 and 38 weeks gestation with a length of stay (LOS) of 0 to 2 days increased the risk of readmission for treatment of hyperbilirubinaemia compared with birth at 39 weeks gestation and LOS of 3 to 4 days<sup>22</sup>. Significant factors related to neonatal readmission for hyperbilirubinaemia include neonates discharged 0 to 2 days of age, vaginal birth, being born to a mother from an Asian country, being born to a first-time mother, or breast feeding at discharge.

The risk of unrecognised severe hyperbilirubinaemia is also increased if:

- There are gaps in clinical handover between hospital and community-based clinicians
- It is unclear who is responsible for the neonate's healthcare in the first days after discharge
- Parents or caregivers do not know what to look for
- Parents or caregivers do not know when, or how, to access a health care professional for review of their neonate's progress after discharge<sup>11,23</sup>.

It is therefore important that a comprehensive discharge plan is formulated with parents or caregivers of neonates at risk of hyperbilirubinaemia (in line with <u>PD2009\_060</u> <u>Clinical Handover – Standard Key Principles</u>).

### 5.1 Timing of follow-up

Table 11 outlines the recommended maximum timing for post hospital discharge followup by clinicians of neonates with or without risk factors for hyperbilirubinaemia; who are jaundiced or who have received phototherapy, based on their age at discharge.

Table 11: Timing of Follow-up			
Age in hours at discharge	Neonate with risk factors for jaundice or who are jaundiced at discharge or who have received phototherapy	Neonate with no risk factors	
Before 24 hours of age	By 48 hours of age	By 72 hours of age	
Between 24 and 48 hours of age	By 72 hours of age	By 96 hours of age	
Between 49 and 72 hours of age	By 96 hours of age	By 120 hours of age	
Adapted from American Academy of Pediatrics Subcommittee on Hyperbilirubinaemia <sup>1</sup>			



# 5.2 Preparation for discharge and clinical handover

Table 12 outlines the preparation that maternity services should take prior to discharge of a neonate < 48 hours of age with risk factors for hyperbilirubinaemia (see Table 1), or who are jaundiced or who have required phototherapy treatment.

Table 12: Preparation for Discharge and Clinical Handover of Neonates < 48 Hours of Age With Risk Factors for Hyperbilirubinaemia or Who are Jaundice at Discharge or Who Have Received Phototherapy

PHOLO	tnerapy
Step	Action
1	If mother is Rh negative, review the neonate's results for  • Blood group  • DAT
2	A TcB as close to discharge as practical ( <b>if the neonate has not been under phototherapy</b> <sup>9</sup> ) An SBR should be completed <sup>1</sup> for neonates who have risk factors for hyperbilirubinaemia or have been under phototherapy
3	If the TcB measurement is <ul> <li>&lt; 20 micromol/L below the treatment threshold line, measure bilirubin with an SBR</li> </ul> If the SBR measurement is <ul> <li>&lt; 50 micromol/L below phototherapy treatment threshold line consider delay of discharge and repeat the SBR in 12 to 24 hours or ensure the parents are aware of the need to repeat the SBR in line with local processes as outlined in point 4 below</li> <li>&gt; 50 micromol/L below the phototherapy threshold line at discharge then clinical follow-up is still necessary in line with Table 11. Such neonates may be discharged with planned clinical follow-up with consideration given for bilirubin measurement using either TcB or SBR as appropriate (see Section 2.2: Measurement)</li> </ul>
4	<ul> <li>Local processes should be in place for</li> <li>Clinical follow-up at appropriate times in line with Table 11</li> <li>The measurement of bilirubin after discharge (see section 2.2 Measurement)</li> <li>The location where follow-up is to occur e.g. in the home or in a community based setting</li> <li>Clinicians to escalate concerns about the ongoing care of neonates with jaundice</li> </ul>
5	<ul> <li>Consider screening for G6PD deficiency pre-discharge if</li> <li>There is a family history</li> <li>This is a male neonate from a high risk ethnic origin/geographic area; African, Asian, Mediterranean, Middle Eastern<sup>2</sup> descent</li> </ul>
6	Document all results in  The clinical record  The Personal Health Record (Blue Book)  The discharge summary
7	Provide parents or caregivers with  Personal Health Record (Blue Book) Discharge summary Copy of any letters of referral Information sheet on jaundice in preferred language see Section 6 Information for parents and care givers Details of any follow-up appointments



# 5.3 Preparation for discharge of neonates who are jaundiced or had phototherapy

All neonates who are jaundiced at discharge or who have received phototherapy should have a bilirubin measurement with either a TcB or an SBR prior to discharge as appropriate (see Section 2.2: <a href="Measurement">Measurement</a>).

This is particularly important to plan for discharge and clinical handover of neonates discharged < 48hours who have risk factors of hyperbilirubinaemia or who have received phototherapy as they require ongoing surveillance, planned and timely follow-up by a clinician<sup>1</sup> as outlined in Table 11 and Table 12.

# 5.4 Jaundice that develops after discharge from hospital

Neonates who develop jaundice after discharge from hospital should be referred for an urgent SBR by the clinician identifying the condition. If the result is above the phototherapy treatment threshold line on the Jaundice Treatment Threshold Graph for gestational age<sup>1, 2</sup> (see <u>Attachments 1-7</u>) the neonate requires <u>urgent medical review</u>, including investigation and readmission without delay.

Rapid readmission will follow local admission protocols developed in line with <u>PD2011\_038 Children and infants - Recognition of a Sick Baby or Child in the Emergency Department</u> and <u>PD2009\_055 Emergency Department - Direct Admission to Inpatient Wards.</u>

### 6. INFORMATION FOR PARENTS AND CAREGIVERS

# 6.1 Neonatal jaundice

Parents and caregivers play an important role in the detection of jaundice and the support of neonates who are jaundiced. Parents and caregivers should be involved in decisions regarding investigations, treatment and care<sup>17</sup> and should receive verbal and written information on jaundice irrespective of whether or not the neonate appears to be jaundiced. A fact sheet has been developed by the Sydney Children's Hospital Network and Kaleidoscope Fact Sheet: Jaundice in newborn babies May 2015 to inform parents and caregivers. This factsheet is available in the following languages:

- English
- Arabic
- Bengali
- Chinese Simplified
- Chinese Traditional
- Dari
- Dinka

- Farsi
- Hindi
- Japanese
- Khmer
- Korean
- Nepali
- Punjabi

- Somali
- Swahili
- Tamil
- Thai
- Turkish
- Urdu
- Vietnamese.



# 6.2 Glucose-6-phosphate dehydrogenase (G6PD) deficiency

On discharge, written information on G6PD deficiency should be given to all parents whose neonate has been diagnosed with this enzyme deficiency, or whose neonate may be at risk of G6PD if:

- There is family history
- This is a male neonate from a high risk ethnic origin/geographic area (African, Asian, Mediterranean and Middle Eastern)<sup>2</sup>.

In Australia, approximately 5% of people from African, Asian, Mediterranean or Middle Eastern descent have G6PD deficiency. Affected neonates can develop massive haemolysis at virtually any time within hours of exposure to triggers such as:

- Clothes stored with moth balls containing naphthalene
- Fava beans also called broad beans
- Sepsis
- Particular medication including some antibiotics.

Mothers who are breast feeding their neonate diagnosed with G6PD may need to avoid the substances and medications that can trigger haemolysis under the guidance of the medical officer caring for the neonate.

Exposure to these triggers most commonly occurs after discharge. A G6PD deficiency Fact Sheet in English is available from the Royal Children's Hospital Melbourne website. A multi-lingual NSW Health information sheet Naphthalene in Moth Balls and Toilet Deodorant Cakes is available from the NSW Multicultural Health Communication Service website.

Further advice on the health risks of naphthalene can be obtained 24 hours a day, 7 days a week Australia-wide from the <u>NSW Poisons Information Centre on 13 11 26</u>, or from local Public Health Units. Contact information for all NSW Public Health Units is available from NSW Health by telephone on 1300 066 055 or via the website and search facility at <a href="http://www.health.nsw.gov.au/Infectious/Pages/phus.aspx">http://www.health.nsw.gov.au/Infectious/Pages/phus.aspx</a>.



### 7. REFERENCES

- 1. The American Academy of Pediatrics Subcommittee Hyperbilirubinaemia. Management of hyperbilirubinaemia in the newborn infant 35 or more weeks of gestation, clinical practice guideline. *Pediatrics*. 2004;114(1):297-316.
- National Institute for Health and Clincal Care Excellence (NICE). Jaundice in newborn babies under 28 days, Clinical Practice Guideline 98 London; 2010 Updated Oct 2016.
- United States Centers for Disease Control (CDC). Kernicterus in full-term infants
   United States 1994-1998. MMWR Weekly. 2001;50(23):491-494.
- 4. Kumar P., Murki S., Malik G.K., Chawla D., Deorari A.K., Karthi N., Subramanian S., Sravanthi J., Goddam P. and Singh S. Light emitting diodes versus compact flourescent tubes for phototherapy in neonatal jaundice: A multi centre randomised controlled trial. *Indian Pediatrics*. 17 February 2010;47 (February):131 137.
- 5. McDonald S.J., Middleton P., Rowsell T., Morris P.S. Effect of timing of umbilical cord clamping of term infants on maternal and neonatal outcomes. *Cochrane Library*. July DOI: 10.1002/14651858.CD004074pub3 2013.
- 6. Dai J., Parry D.M., Krahn J. Transcutaneous bilirubinometry: Its role in the assessment of neonatal jaundice. *Clinical Biochemistry*. February 1997;30(1):1-9.
- 7. Grohmann K., Roser M., Rolinski B., Kadow I., Muller C., Goerlach-Graw A., Nauck M., Kuster H. Bilirubin measurement for neonates: Comparison of 9 frequently used methods. *Pediatrics*. 2006;117:1174-1183.
- 8. Karen T., Bucher H.U., Fauchere J.C. Comparison of a new transcutaneous bilirubinometer (Bilimed\*) with serum bilirubin measurements in preterm and full-term infants. *BMC Pediatrics*. 2009 9(70).
- 9. Nagar G., Vandermeer B., Campbell S., Kumar M. Effect of phototherapy on the reliability of transcutaneous bilirubin devices in term and near-term infants: A systematic review and meta-Analysis. *Neonatology*. 2016;109:203 212.
- 10. Lease M., Whalen B. Assessing jaundice in infants of 35 week gestation and greater. *Current Opinion in Pediatrics*. June 2010;22:352-365.
- 11. The Health Care Complaints Commission. Confidential report to director-general of health in: NSW Health. Unpublished; 2011.
- 12. Maisels M.J., McDonagh A.F. Phototherapy for neonatal jaundice. *N Engl J Med.* February 2008;358(9):920-928.
- 13. Bhutani V.K. and The Committee on Fetus and Newborn. Technical Report: Phototherapy to prevent severe neonatal hyperbilirubinemia in the newborn. *American Academy of Pediatrics*. 2011;128:e1046-1052.
- 14. Wentworth S. Neonatal phototherapy today's lights, lamps and devices. *Infant.* 2005;1:14-19.
- 15. Malwade U., Jardine L. Home versus hospital-based phototherapy for the treatment of non-haemolytic jaundice in infants at more than 37 weeks gestation. *Cochrane Database of Systematic Reviews.* June DOI: 10.1002/14651858.CD010212.pub2 2014(6).
- 16. National Institute for Health and Clincal Care Excellence (NICE). Clinical Guidleine 98 Neonatal Jaundice, Surveillance proposal Guidance Executive document. August 2014.



- 17. National Institute for Health and Clincal Care Excellence (NICE). Jaundice in newborn babies under 28 days: Quality standard 57. 6 March 2014.
- 18. Alcock G., Liley H. Immunoglobulin infusion for isoimmune haemolytic jaundice in neonates. *Cochrane Database of Systematic Reviews* 2002 (3).
- 19. Onishi S., Itoh S., Isobe K., Ochi M., Kunikata T., Imai T. Effect of the binding of bilirubin to either the first class or the second class of binding sites of the human serum albumin molecule on its photochemical reaction. *Biochem J* 1989;257(3):711-714.
- 20. Kulkarni M., Elsner C., Ovellet D. Pediatric vascular access and blood sampling techniques. *Roberts and Hedges' Clinical Procedures in Emergency Medicine*. 6 ed. Philadelphia: Elsevier; 2014.
- 21. Young P., Korgenski K., Buchi K. Early readmission of newborns in a large health care system. *Pediatrics* 2013(131):e-1538-e1544.
- 22. Lain S., Roberts C., Bowen J., Nassar N. Early discharge of infants and risk of readmission for jaundice. *Pediatrics*. 2015;135(2):314-321.
- 23. Salem-Schatz S., Peterson L., Palmer H., Clanton M., Ezhuthachan S., Luttrell R., Newman C., Westbury R. Barriers to first-week follow-up of newborns: findings from parent and focus groups *Joint Commission Journal on Quality and Patient Safety.* 2004;30(11):593-601.
- 24. MacDonald M. Hidden risks: Early discharge and bilirubin toxicity due to glucose 6-phosphate dehydrogenase deficiency. *Pediatrics* 1995;96(4):734-738.
- 25. Vandborg P.K, Hansen, B.M., Greisen, G., Ebbesen, F. Dose-response relationship of phototherapy for hyperbilirubinaemia. *Pediatrics*. 2012:e352 to e357



# 8. APPENDIX A: ABBREVIATIONS

Appendix A:	Abbreviations
CERS	Clinical Emergency Response System
DAT	Direct antiglobulin test (also known as the Coombs test)
FBC	Full blood count
G6PD	Glucose-6-phosphate dehydrogenase
LHD	Local Health District
Micromol/L	Micromol per litre
NETS	Newborn and paediatric Emergency Transport Service (NETS NSW)
NICU	Neonatal Intensive Care Unit
PSN	Pregnancy and newborn Services Network
Rh	Rhesus antigen on red blood cells
SBR	Serum bilirubin
SCN	Special care nursery
SNOC	Standard neonatal observation chart
TcB	Transcutaneous bilirubin
UVC	Umbilical venous catheter
μW.cm <sup>-2</sup> nm <sup>-1</sup>	Light irradiance
	cm <sup>2</sup> - body surface area
	nm - light source



### 9. APPENDIX B: MAXIMISING PHOTOTHERAPY EFFICACY

### **Appendix B: Maximising Phototherapy Efficacy**

#### Phototherapy efficacy depends on three criteria

- Effectiveness of the light source
- Dose (light intensity or irradiance) of phototherapy administered

#### The skin surface area effectively illuminated by the phototherapy light Lights in the blue and blue-green spectrum on conventional devices have both been found to be effective <sup>13,14</sup>. Wavelengths in the blue-green spectrum (~460-Light source effectiveness 490 nm) are effective with special blue being the most effective (~460 nm)<sup>1</sup> Do not use white lights painted blue or covered with blue plastic sheaths<sup>13</sup> Position the light as close to the neonate as manufacturer's instructions allow Position the light rays perpendicular to the surface of the humidicrib to minimise reflectance and loss of efficacy<sup>13</sup> Fibre optic phototherapy devices use a standard light source, usually a quartz halogen bulb. Filtered light passes through a fibre optic bundle into a pad of woven optic fibres that can be placed next to the neonate's skin<sup>14</sup>. The effect of fibre optic devices may be limited by the size of the device and the surface area of skin exposed<sup>2</sup> particularly when used for larger neonates Dose of • Light intensity output (or irradiance) varies widely between devices and depends phototherapy on factors such as the number and quality of bulbs, tubes or light sources<sup>1</sup> administered • Light intensity output is displayed on each device and is usually measured in microwatts per cm<sup>2</sup> of exposed skin (μW/sq cm<sup>2</sup>), confirmed using the irradiance meter recommended by the device manufacturer, calibrated over the appropriate wavelength range<sup>13</sup> Evidence suggests phototherapy increases effectiveness in a linear relationship from 20 to 55 μW/cm<sup>2</sup>/nm<sup>-1</sup> and demonstrates a decrease in TcB after 24hrs of therapy. No evidence of saturation point was demonstrated <sup>25</sup> Check phototherapy devices regularly as per local protocols in accordance with the manufacturer's instructions<sup>13</sup>. With use, the irradiance of all lamps decreases, so do not utilise beyond the manufacturer's useful-lifetime estimates 13 NOTE: Heat generation from halogen or tungsten lights can cause a burn so manufacturer's instructions should always be followed for the minimum distance from the light to the neonate as this can vary from 25cm to 50cm<sup>13</sup> The skin Ensure the light is not obstructed by equipment or objects that decreases the surface area exposed skin surface area, such as effectively Radiant warmers exposed to Head covers phototherapy Large nappies treatment Large eye masks that cover large areas of the scalp Electrode patches Insulating plastic covers<sup>13</sup>



# 10. APPENDIX C: RELEVANT DOCUMENTS

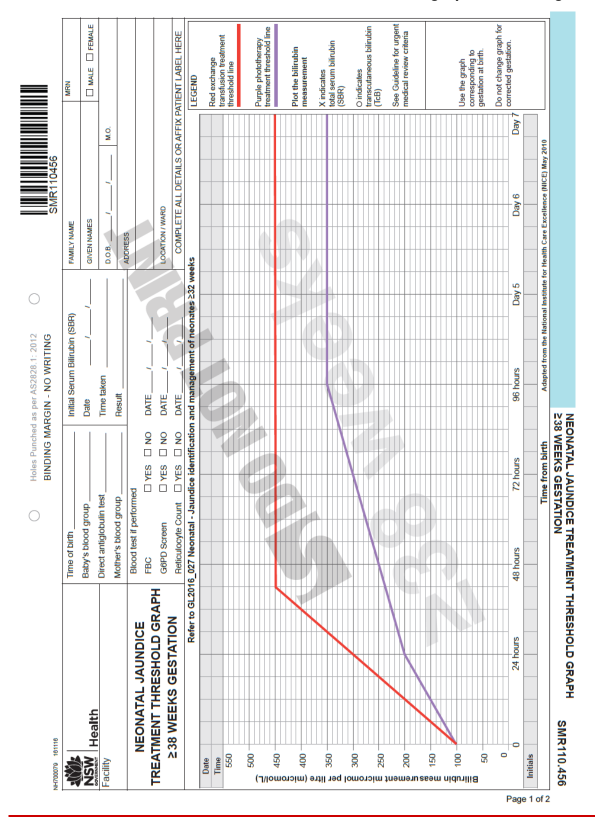
Appendix D	: Relevant Documents	
Туре	Publisher	Title
Guideline	NSW Health	GL2016_018 Maternity and Neonatal Service Capability Framework
Policy Directive	NSW Health	PD2010 69 Critical Care Tertiary Referral Networks (Perinatal)
Policy Directive	NSW Health	PD2010 030 Critical Care Tertiary Referral Networks (Paediatrics)
Policy Directive	NSW Health	PD2011 015 Care Coordination: Planning from Admission to Transfer of Care in NSW Public Hospitals
Policy Directive	NSW Health	PD2009_060 Clinical Handover - Standard Key Principles
Policy Directive	NSW Health	PD2010_022 National Midwifery Guidelines for Consultation and Referral
Policy Directive	NSW Health	PD2011 038 Children and infants - Recognition of a Sick Baby or Child in the Emergency Department
Policy Directive	NSW Health	PD2009 055 Emergency Department - Direct Admission to Inpatient Wards
Policy Directive	NSW Health	PD2013 049 Recognition and Management of Patients Who are Clinically Deteriorating
Guideline	NSW Health	GL2008 015 Term Changeover - ensuring an effective handover of patient care
Resource	Pregnancy and newborn Services Network	Neonatal Exchange Transfusion in a Non-Tertiary Hospitals - How to guide
Resource	NSW Health	My Personal Health Record (Blue Book)
Resource	The Sydney Children's Hospitals Network and Kaleidoscope Hunter Children's Health Network.	Fact Sheet: Jaundice in newborn babies. Available in 20 languages
Resource	NSW Health	NSW Health: Naphthalene in moth balls and toilet deodorant cakes
Resource	Royal Children's Hospital Melbourne	G6PD Deficiency Fact Sheet 2011



### 11. ATTACHMENTS

### **NEONATAL JAUNDICE TREATMENT THRESHOLD GRAPHS**

### Attachment 1: Neonatal Jaundice treatment threshold graph 38 weeks gestation





	Use single light phototherapy if	FAMILY NAME MICN	
NSW Leafth	<ul> <li>SBR is at or above the phototherapy treatment threshold line</li> </ul>	GIVEN NAMES	FEMALE
-1	Use multiple light phototherapy if		
-acmiy	<ul> <li>SBR is rising rapidly (&gt;8.5 micromol/L per hour)</li> </ul>	D.O.B// M.O.	
	<ul> <li>SBR is &lt;50 micromol/L below the RED exchange transfusion line</li> </ul>	ADDRESS	
NEONATAL JAUNDICE	<ul> <li>SBR fails to respond to single light phototherapy</li> </ul>		
LONG O ICHOSTON TABLES	If the SBR is rapidly rising or approaching the RED exchange		
Vac WITTE OFFITTION	transfusion treatment threshold line an urgent medical review	LOCATION / WARD	
230 WEERS GESTATION		COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE	L HERE
Neonate with jaundice <24 hours of age or greater than 14 days of age	ys of age Table A Risk Factors and Causes of Neonatal Jaundice	dice	
should have urgent medical review and	Jaundice <24 hours of age - Suspect haemolysis until proven otherwise	ntil proven otherwise	
Measure the SDR and plot on the jaundre treatment unestrong graph.  Urgent medical review will determine when to start phototherapy.	Jaundice due to haemolysis	Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E. Non-immune - e.g. G6PD	
Consider staining photomerapy at a lower ober if the neonate has risk factors for neonatal jaundice (see Table A) or is unwell	Individual neonatal risk factors • Prematurity • Asphyxia		
Measure the SBK every o 6 hours until the SBR is both below the phototherapy treatment threshold line and stable or falling then		Apgar <7 at 5 minutes or acidosis pH <7 or base excess <12 mEq/L Low serum albumin <30 grams per litre Sersis or connential infections	
o 12-24 hourly for the duration of treatment	Material diabeter mocaons     Material diabeter     Carbalohamatoma / bruising	/ bruising	
Consider o Additional investigations (see <i>Table B</i> )	History of subling with family     G6PD nisk with family	oppragnos assuments as a supersection of the s	
o Transfer to a higher level facility if appropriate	Table B Additional Investigations		
Neonate with jaundice ≥24 hours of age	Clinical Feature Investigation		
Do a transcutaneous bilirubin (1cB) if well and ≥35 weeks or	Neonate of Rhesus negative Blood Group		
o Unwell or <35 weeks	mother Direct Antiglobulin Test (DAT)	(DAT)	
o The TcB is ≥250 micromol/L or		An immediate 55K is required if the DAT is positive and the 55K is unknown	
o The TcB is <20 micromol/L below the treatment threshold line	Neonate with jaundice within the first 24 hours of age	Full blood count (FBC) and film with reticulocyte Blood group	
Medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk factors for neonatal	OR Neonate with a ranidly rising total	DAT Santic craan including blood and uning cultura & cancitivity if cancic cusparted	
jaundice (see Table A) or is unwell	SBR (>8.5 micromol/L per hour)	מספת מוות תוווני בתונתו כית ספו שותאול זו ספקסים סתסקיפינים	
If SBR <50 micromol/L below the phototherapy treatment threshold line repeat the SBR in 12-24 hours	OR Neonate with a total SBR above the	I here is a family history This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian	n, Asian
If SBR >50 micromol/L below the phototherapy treatment threshold line	phototherapy threshold	Mediterranean and Middle Eastern descent	
continue regular visual assessments If phototherapy is commenced measure SBR	Identification of maternal blood group should also be considered with the above investigations	sidered with the above investigations	
o After 6 hours to ensure SBR is stable or falling, then	Neonate with a total SBR approaching exchange transfusion thresholds	Serum albumin level     Liver function tests	
o Every 12-24 hours for the duration of treatment Consider		Conjugated bilirubin	
o Additional investigations (see Table B)	A neonate of any gestation with a conjugated bilirubin >20 mic	A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior	nost senior
o Transfer to a higher level facility if appropriate	medical office (same day before discharge norm nospiral)		
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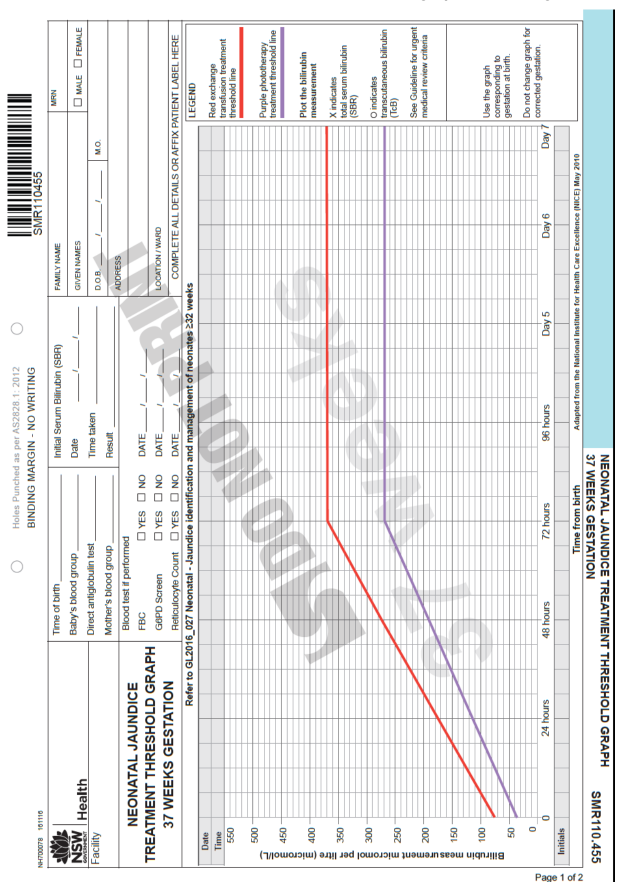
 Every 12-24 hours for the duration of tr o Additional investigations (see Table B) o Transfer to a higher level facility if appri Consider

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# Attachment 2: Neonatal Jaundice treatment threshold graph 37 weeks gestation





	Use single light phototherapy if	FAMILY NAME	
WSW Localth	<ul> <li>SBR is at or above the phototherapy treatment threshold line</li> </ul>	GIVEN NAMES	FEMALE
	Use multiple light phototherapy if		
Facility	<ul> <li>SBR is rising rapidly (&gt;8.5 micromol/L per hour)</li> </ul>	D.O.B. / M.O.	
	<ul> <li>SBR is &lt;50 micromol/L below the RED exchange transfusion line</li> </ul>	ADDRESS	
NEONATAL JAUNDICE	<ul> <li>SBR fails to respond to single light phototherapy</li> </ul>		
TREATMENT THRESHOLD GRAPH	If the SBR is rapidly rising or approaching the RED exchange transfusion treatment threshold line an urgent medical review	1 OCATION (MARDI	
37 WEEKS GESTATION	should occur	COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE	EL HERE
Neonate with jaundice <24 hours of age or greater than 14 days of age	s of age Table A Risk Factors and Causes of Neonatal Jaundice	90	
should have urgent medical review and	Jaundice <24 hours of age - Suspect haemolysis until proven otherwise	I proven otherwise	
Measure the SBK and plot on the jaundice treatment threshold graph     Ugent medical review will determine when to start phototherapy.	Jaundice due to haemolysis	Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Non-immune - e.g. G6PD	
Consider starting photomerapy at a lower SDR if the neonate has risk factors for neonatal jaundice (see Table A) or is unwell	Individual neonatal risk factors		
<ul> <li>Measure the SBR every</li> <li>6 hours until the SBR is both below the phototherapy treatment</li> </ul>		Asphyxia Apgar ≺7 at 5 minutes or acidosis pH <7 or base excess ≤12 mEq/L Low serum albumin <30 grams per lftre	
threshold line and stable or falling, then	Sepsis or congenital infections     Maternal diabetes	fections	
Consider	Cephalohaematoma / bruising     Director of cilding upon user in uncertainty	Cephalohaematoma / bruising	
o Additional investigations (see Table B)	G6PD risk with family b	nistory of starting who was jaturianced as a recritate G6PD risk with family history or with exposure to trigger	
o Transfer to a higher level facility if appropriate	Table B Additional Investigations		
Neonate with jaundice ≥24 hours of age	Clinical Feature Investigation		
<ul> <li>Do a transcutaneous bilirubin (IcB) if well and ≥35 weeks or</li> </ul>	Neonate of Rhesus negative Blood Group		
o Unwell or <35 weeks	mother Direct Antiglobulin Test (DAT)	AT)	
o The TcB is ≥250 micromol/L or	1	An immediate SBK is required if the DALL is positive and the SBK is unknown	
o The TcB is <20 micromol/L below the treatment threshold line  Medical review will determine when to start nhototherary. Consider starting	Neonate with jaundice within the Full blood count (FBC) and film with reticulocyte floor sider starting to be sider starting.	d film with reticulocyte	
phototherapy at a lower SBR if the neonate has risk factors for neonatal jaundice (see <i>Table A</i> ) or is unwell	Neonate with a rapidly rising total SBR (>8.5 micromol/L per hour)	Septic screen including blood and urine culture & sensitivity if sepsis suspected A G6PD screen if	
• If SBR <50 micromol/L below the phototherapy treatment threshold line repeat the SBR in 12-24 hours	OR Neonate with a total SBR above the	There is a family history This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian	an, Asian
If SBR >50 micromol/L below the phototherapy treatment threshold line	shold line phototherapy threshold Mediterranean and Middle Eastern descent	idle Eastern descent	
United Education Vision assessments If phototherapy is commenced measure SBR	Identification of maternal blood group should also be considered with the above investigations	lered with the above investigations	
o After 6 hours to ensure SBR is stable or falling, then	Neonate with a total SBR approaching exchange transfusion thresholds	Serum albumin level Liver function tests	
o Every 12-24 hours for the duration of treatment		Conjugated bilirubin	
o Additional investigations (see Table B)	A neonate of any gestation with a conjugated bilirubin >20 micro	A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior	most senior
I a a serial management of the serial	medical officer (same day before discharge from hospital)		

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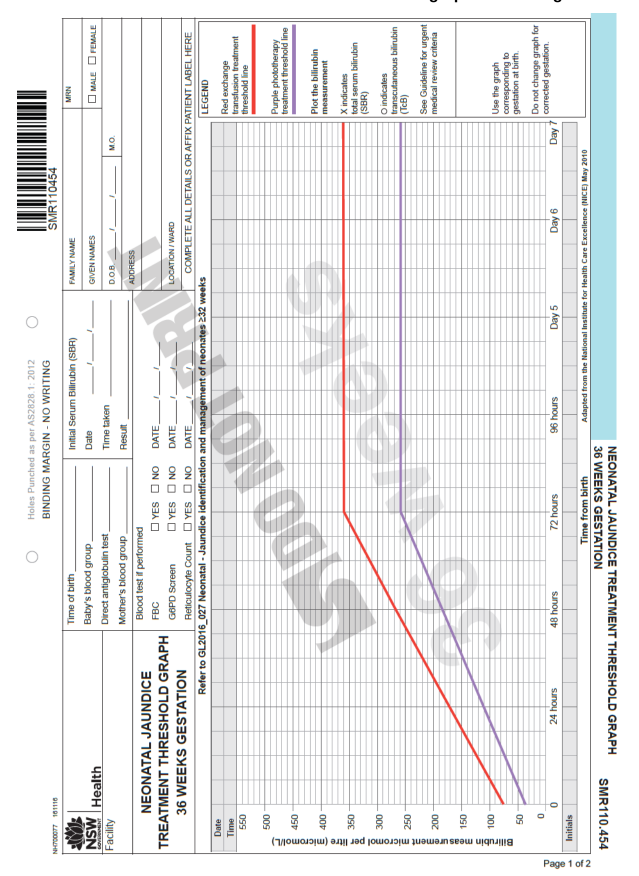
o Transfer to a higher level facility if appropriate

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### Attachment 3: Neonatal Jaundice treatment threshold graph 36 weeks gestation





FEMALE A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE There is a family history This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian MALE Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Septic screen including blood and urine culture & sensitivity if sepsis suspected A G6PD screen if An immediate SBR is required if the DAT is positive and the SBR is unknown Apgar <7 at 5 minutes or acidosis pH <7 or base excess ≤12 mEq/L Identification of maternal blood group should also be considered with the above investigations History of sibling who was jaundiced as a neonate G6PD risk with family history or with exposure to trigger LOCATION / WARD **GIVEN NAMES** FAMILY NAME Jaundice <24 hours of age - Suspect haemolysis until proven otherwise Full blood count (FBC) and film with reticulocyte Blood group DAT Serum albumin level
 Liver function tests Mediterranean and Middle Eastern descen ADDRESS Low serum albumin <30 grams per litre D.O.B. Sepsis or congenital infections Cephalohaematoma / bruising Direct Antiglobulin Test (DAT) Table A Risk Factors and Causes of Neonatal Jaundice Non-immune - e.g. G6PD SBR is <50 micromol/L below the RED exchange transfusion line</li> transfusion treatment threshold line an urgent medical review If the SBR is rapidly rising or approaching the RED exchange Maternal diabetes medical officer (same day before discharge from hospital) SBR is at or above the phototherapy treatment threshold line Investigation Prematurity Blood Group Asphyxia Neonate with a total SBR approaching exchange transfusion thresholds SBR fails to respond to single light phototherapy SBR is rising rapidly (>8.5 micromol/L per hour) first 24 hours of age
OR
Neonate with a rapidly rising total
SBR (>8.5 micromol/L per hour)
OR
Neonate with a total SBR above the Neonate with jaundice within the Individual neonatal risk factors Neonate of Rhesus negative Jaundice due to haemolysis Use multiple light phototherapy if Jse single light phototherapy if phototherapy threshold Table B Additional Clinical Feature mother should occur Medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk factors for neonatal Neonate with jaundice <24 hours of age or greater than 14 days of age If SBR <50 micromol/L below the phototherapy treatment threshold line If SBR >50 micromol/L below the phototherapy treatment threshold line Urgent medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk Measure the SBR and plot on the jaundice treatment threshold graph o 6 hours until the SBR is both below the phototherapy treatment o The TcB is <20 micromol/L below the treatment threshold line Do a transcutaneous bilirubin (TcB) if well and ≥35 weeks or **IREATMENT THRESHOLD GRAPH** factors for neonatal jaundice (see Table A) or is unwell o After 6 hours to ensure SBR is stable or falling, then **NEONATAL JAUNDICE** 36 WEEKS GESTATION Every 12-24 hours for the duration of treatment Transfer to a higher level facility if appropriate Transfer to a higher level facility if appropriate If phototherapy is commenced measure SBR o 12-24 hourly for the duration of treatment threshold line and stable or falling, then o Additional investigations (see Table B) o Additional investigations (see Table B) Neonate with jaundice ≥24 hours of age should have urgent medical review and continue regular visual assessments jaundice (see Table A) or is unwell o The TcB is ≥250 micromol/L or

Consider

repeat the SBR in 12-24 hours

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Consider

o Unwell or <35 weeks

Do an SBR if

Measure the SBR every

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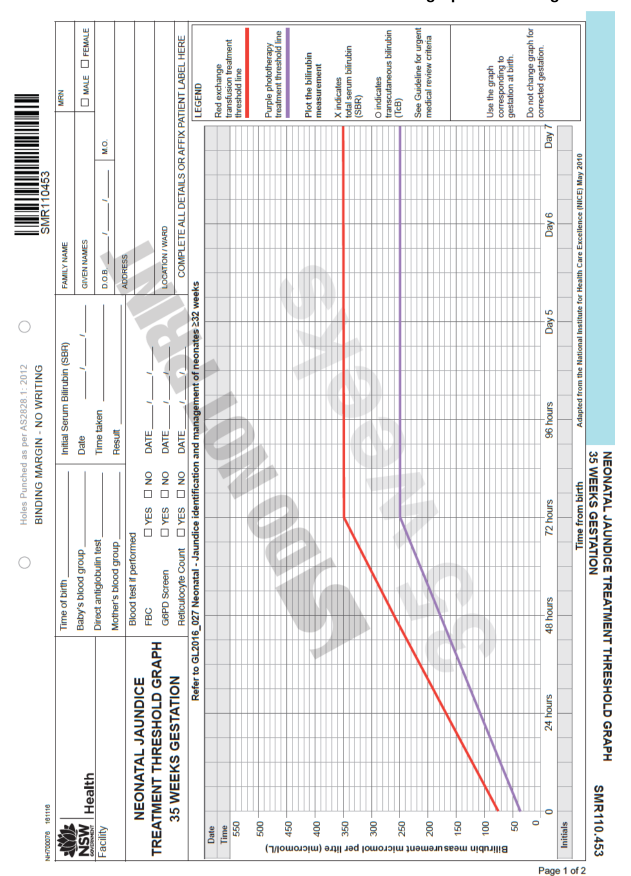
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### Attachment 4: Neonatal Jaundice treatment threshold graph 35 weeks gestation





☐ FEMALE A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior medical officer (same day before discharge from hospital) COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian MALE Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Septic screen including blood and urine culture & sensitivity if sepsis suspected An immediate SBR is required if the DAT is positive and the SBR is unknown Apgar <7 at 5 minutes or acidosis pH <7 or base excess ≤12 mEq/L identification of maternal blood group should also be considered with the above investigations Cephalohaematoma / bruising History of sibling who was jaundiced as a neonate G6PD risk with family history or with exposure to trigger LOCATION / WARD GIVEN NAMES Jaundice <24 hours of age - Suspect haemolysis until proven otherwise FAMILY NAME Full blood count (FBC) and film with reticulocyte Serum albumin level Liver function tests ADDRESS Mediterranean and Middle Eastern descen D.O.B. Low serum albumin <30 grams per litre Sepsis or congenital infections Direct Antiglobulin Test (DAT) Table A Risk Factors and Causes of Neonatal Jaundice There is a family history Non-immune - e.g. G6PD . . SBR is <50 micromol/L below the RED exchange transfusion line</li> If the SBR is rapidly rising or approaching the RED exchange transfusion treatment threshold line an urgent medical review Maternal diabetes SBR is at or above the phototherapy treatment threshold line A G6PD screen if Investigation BINDING MARGIN - NO WRITING Prematurity Blood Group Blood group Neonate with a total SBR approaching exchange SBR fails to respond to single light phototherapy SBR is rising rapidly (>8.5 micromol/L per hour) first 24 hours of age
OR
Neonate with a rapidly rising total
SBR (>8.5 micromol/L per hour)
OR
Neonate with a total SBR above the Table B Additional Investiga Veonate with jaundice within the Individual neonatal risk factors Neonate of Rhesus negative Jaundice due to haemolysis Use multiple light phototherapy if Jse single light phototherapy if phototherapy threshold ransfusion thresholds Clinical Feature mother should occur Medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk factors for neonatal If SBR <50 micromol/L below the phototherapy treatment threshold line If SBR >50 micromol/L below the phototherapy treatment threshold line Neonate with jaundice <24 hours of age or greater than 14 days of age Urgent medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk Measure the SBR and plot on the jaundice treatment threshold graph o 6 hours until the SBR is both below the phototherapy treatment o The TcB is <20 micromol/L below the treatment threshold line Do a transcutaneous bilirubin (TcB) if well and ≥35 weeks or TREATMENT THRESHOLD GRAPH factors for neonatal jaundice (see Table A) or is unwell After 6 hours to ensure SBR is stable or falling, then 35 WEEKS GESTATION **NEONATAL JAUNDICE**  Every 12-24 hours for the duration of treatment o Transfer to a higher level facility if appropriate Transfer to a higher level facility if appropriate If phototherapy is commenced measure SBR 12-24 hourly for the duration of treatment o Additional investigations (see Table B) threshold line and stable or falling, then Additional investigations (see Table B) Neonate with jaundice ≥24 hours of age should have urgent medical review and continue regular visual assessments jaundice (see Table A) or is unwell The TcB is ≥250 micromol/L or

Page 2 of 2

Consider

repeat the SBR in 12-24 hours

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Consider

o Unwell or <35 weeks

Do an SBR if

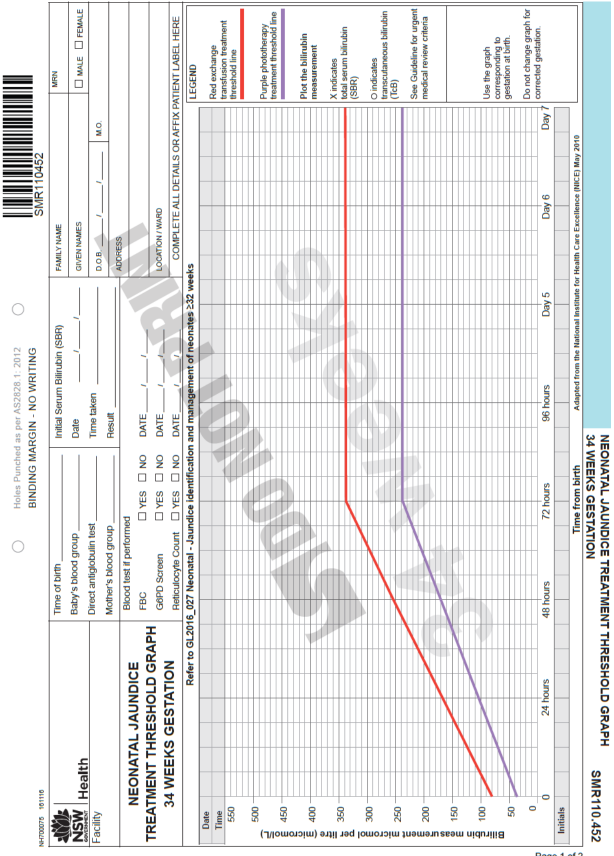
Measure the SBR every

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# Attachment 5: Neonatal Jaundice treatment threshold graph 34 weeks gestation



Page 1 of 2



	Use single light phototherapy if		FAMILY NAME	MRN
NSW Health	• SBR is at or above the phototherapy treatment threshold line	reshold line	GIVEN NAMES	□ MALE □ FEMALE
Government Incoluit Facility	Use multiple light phototherapy if  SBR is rising rapidly (>8.5 micromol/L per hour)		D.O.B// M.O.	
	SBR is <50 micromol/L below the RED exchange transfusion line	transfusion line	ADDRESS	
NEONATAL JAUNDICE	SBR fails to respond to single light phototherapy     He the EBD is president in the case of the CBD is president in the case of the			
TREATMENT THRESHOLD GRAPH	in the SDK is raptury fishing or approaching the KED exchange transfusion treatment threshold line an urgent medical review	medical review	LOCATION / WARD	
34 WEEKS GESTATION	should occur		COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE	TENT LABEL HERE
Neonate with jaundice <24 hours of age or greater than 14 days of age	/s of age Table A Risk Factors and Causes of Neonatal Jaundice	of Neonatal Jaundice		
should have urgent medical review and	Jaundice <24 hours of age - Suspect haemolysis until proven otherwise	ct haemolysis until proven	otherwise	
<ul> <li>Measure the SBK and plot on the jaundice treatment threshold graph.</li> <li>Urgert medical review will determine when to start phototherapy.</li> </ul>	Jaundice due to haemolysis	Immune - e.g. ABO blood group inco Non-immune - e.g. G6PD	Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Non-immune - e.g. G6PD	
Consider starting prodouteraby at a lower SDR if the neonate has risk factors for neonatal jaundice (see <i>Table A</i> ) or is unwell	Individual neonatal risk factors	Prematurity Asphyxia		
<ul> <li>Measure the SbK every of hours until the SBR is both below the phototherapy treatment threshold line and stable or falling then</li> </ul>		Apgar <7 at 5 minutes or acidosis pH	Apgar <7 at 5 minutes or acidosis pH <7 or base excess ≤12 mEq/L Low secum albumist <30 grams per litre Convesion concaminal infections	
o 12-24 hourly for the duration of treatment		Maternal diabetes		
<ul> <li>Consider</li> <li>Additional investigations (see Table B)</li> </ul>		Oephaonaemaoma Fousning History of sibling who was jaundiced as a neonate GEDD risk with family history or with exposure to tringer	ced as a neonate	
o Transfer to a higher level facility if appropriate	Table B Additional Investigations			
Neonate with jaundice ≥24 hours of age	Clinical Feature Inve	Investigation		
· Do a transcutaneous bilirubin (TcB) if well and ≥35 weeks or	Neonate of Rhesus negative Blood	Blood Group		
- Do an SBK III	mother Direc	Direct Antiglobulin Test (DAT)		
o The TcB is >250 micromol/l or	An in	mmediate SBR is required if the	An immediate SBR is required if the DAT is positive and the SBR is unknown	
o The TcB is <20 micromol/L below the treatment threshold line	Neonate with jaundice within the first 24 hours of age	Full blood count (FBC) and film with reticulocyte Blood group	reticulocyte	
<ul> <li>Medical review will determine when to start photomerapy. Consider starting phototherapy at a lower SBR if the neonate has risk factors for neonatal</li> </ul>	OR Noonate with a ranidly rising total	fic screan including blood and	DAT Santin ceraan including blood and uring pullura & cancitivity if cancic cuspartad	
jaundice (see Table A) or is unwell	SBR (>8.5 micromol/L per hour)	A G6PD screen if	ille cuitile & selisium   1 sepsis suspecieu	
<ul> <li>If SBR &lt;50 micromol/L below the phototherapy treatment threshold line repeat the SBR in 12-24 hours</li> </ul>	OR Neonate with a total SBR above the	There is a family history This is a male neonate with dark	l here is a family history This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian	ea e.g. African, Asian
If SBR >50 micromol/L below the phototherapy treatment threshold line	phototherapy threshold	Mediterranean and Middle Eastern descent	m descent	
If phototherapy is commenced measure SBR	Identification of maternal blood group should also be considered with the above investigations	hould also be considered with	n the above investigations	
o After 6 hours to ensure SBR is stable or falling, then be Every 12-24 hours for the duration of treatment	Neonate with a total SBR approaching exchange transfusion thresholds		Serum albumin level Liver function tests Coniugated bilirubin	
Consider		an Balino		
o Additional investigations (see <i>Table B</i> ) o Transfer to a higher level facility if appropriate	A neonate of any gestation with a conjugated bilirubin >20 medical officer (same day before discharge from hospital)	ed bilirubin >20 micromol/L or > e from hospital)	A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior medical officer (same day before discharge from hospital)	riew by the most senior
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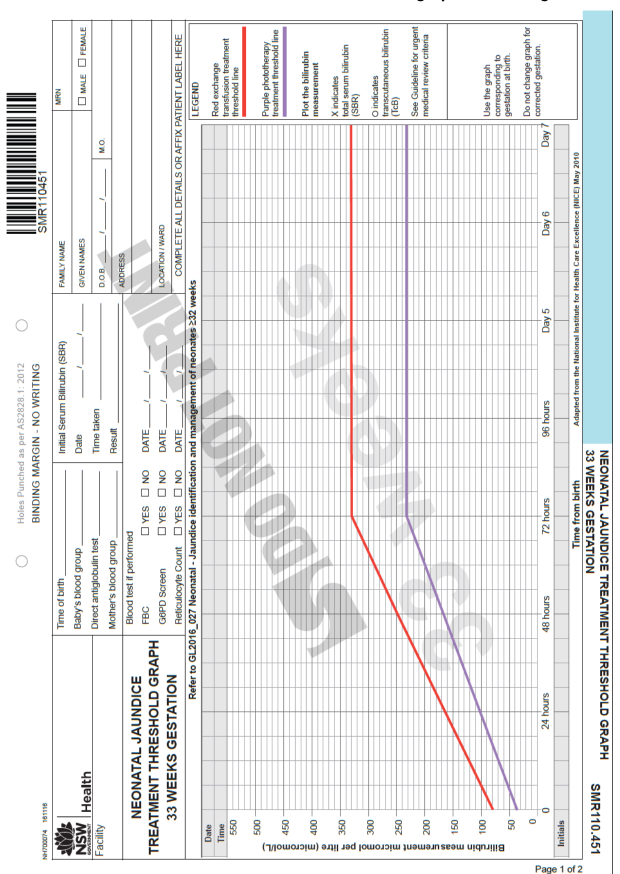
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### Attachment 6: Neonatal Jaundice treatment threshold graph 33 weeks gestation





	Use single light phototherapy if	FAMILY NAME	MKN
NSN III	<ul> <li>SBR is at or above the phototherapy treatment threshold line</li> </ul>	GIVEN NAMES	MAIF FEMAIE
GOVERNMENT FREAITH	Use multiple light phototherapy if		
Facility	<ul> <li>SBR is rising rapidly (&gt;8.5 micromol/L per hour)</li> </ul>	D.O.B// M.O.	
	<ul> <li>SBR is &lt;50 micromol/L below the RED exchange transfusion line</li> </ul>	ADDRESS	
NEONATAL JAUNDICE	<ul> <li>SBR fails to respond to single light phototherapy</li> </ul>		
TREATMENT THRESHOLD GRAPH	If the SBR is rapidly rising or approaching the RED exchange transfusion treatment threshold line an urgent medical review	OCATION AWARD	
33 WEEKS GESTATION	should occur	COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE	ENT LABEL HERE
Neonate with jaundice <24 hours of age or greater than 14 days of age	of age Table A Risk Factors and Causes of Neonatal Jaundice		
should have urgent medical review and	Jaundice <24 hours of age - Suspect haemolysis until proven otherwise	ren otherwise	
Measure the SbK and plot on the jaundlee treatment threshold graph     Urgent medical review will determine when to start phototherapy.	apn Jaundice due to haemolysis	Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Non-immune - e.g. G6PD	
Consider stating protouneraby at a lower SDR in the neonate has risk factors for neonatal jaundice (see <i>Table A</i> ) or is unwell	Individual neonatal risk factors • Prematurity		
<ul> <li>Measure the SBR every</li> <li>6 hours until the SBR is both below the phototherapy treatment</li> </ul>		Appar <7 at 5 minutes or acidosis pH <7 or base excess ≤12 mEq/L Low serum albumin <30 grams per litre	
threshold line and stable or falling, then		· SI	
Consider	Cephalohaematoma / bruising	g of	
o Additional investigations (see Table B)	G6PD risk with family history or with exposure to trigger	or with exposure to trigger	
o Transfer to a higher level facility if appropriate	Table B Additional Investigations		
Neonate with jaundice ≥24 hours of age	Clinical Feature Investigation		
• Do a transcutaneous bilirubin (1cb) if well and ≥35 weeks or	Neonate of Rhesus negative Blood Group		
• Do an SDR II	mother Direct Antiglobulin Test (DAT)		
o The TcB is ≥250 micromol/L or		An immediate SBR is required if the DAT is positive and the SBR is unknown	
o The TcB is <20 micromol/L below the treatment threshold line	Neonate with jaundice within the Full blood count (FBC) and film with reticulocyte first 24 hours of age	with reticulocyte	
<ul> <li>Medical review will determine when to start phototherapy. Consider starting phototherapy at a lower SBR if the neonate has risk factors for neonatal</li> </ul>	OR Neonate with a rapidly rising total	DAT Sentic screen including blood and unite culture & sensitivity if sensis suspected	
jaundice (see Table A) or is unwell	SBR (>8.5 micromol/L per hour)		
<ul> <li>If SBK &lt;50 micromol/L below the phototherapy treatment threshold line repeat the SBR in 12-24 hours</li> </ul>	OR Neonate with a total SBR above the	nere is a family history This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian	a e.g. African, Asian
<ul> <li>If SBR &gt;50 micromol/L below the phototherapy treatment threshold line continue regular visual assessments.</li> </ul>	old line phototherapy threshold Mediterranean and Middle Eastern descent	astern descent	
If phototherapy is commenced measure SBR	Identification of maternal blood group should also be considered with the above investigations	with the above investigations	
o After 6 hours to ensure SBR is stable or falling, then	Neonate with a total SBR approaching exchange	Serum albumin level	
o Every 12-24 hours for the duration of treatment	•	Conjugated bilirubin	
• Consider	Management of a side will be about in a continuous of the continuo	in leading a supply bloods of the control of the co	1
o Additional investigations (see Table B)	A neonate of any gestation with a conjugated bilirtibin >20 micromovil of >20% of the total SDK, should have a medical review by the most senior medical officer (same day before discharge from hospital)	of >20% of the total SBK, should have a medical revi	lew by the most senior
<ul> <li>Transfer to a higher level facility if appropriate</li> </ul>			

Consider

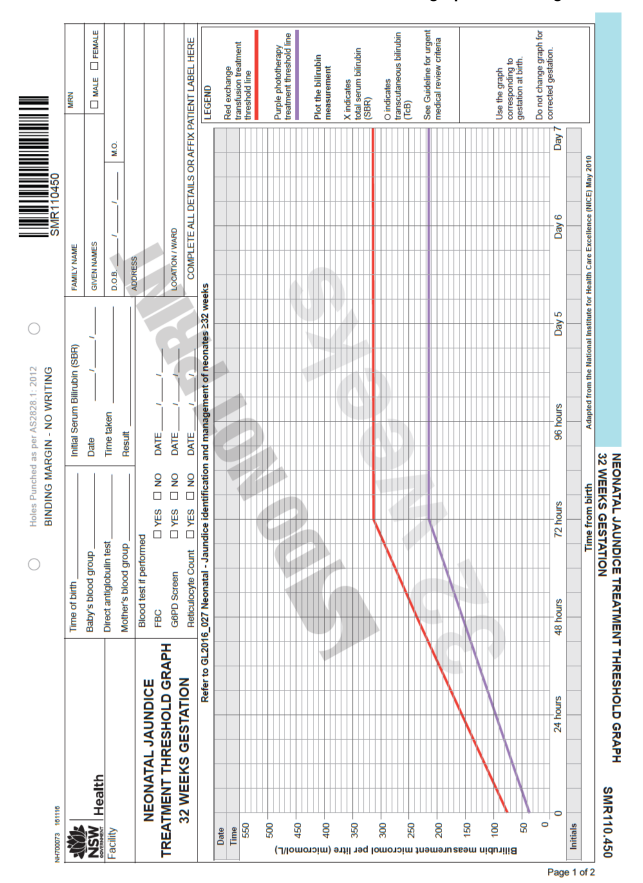
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**BINDING MARGIN - NO WRITING** Holes Punched as per AS2828.1: 2012



### Attachment 7: Neonatal Jaundice treatment threshold graph 32 weeks gestation



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A neonate of any gestation with a conjugated bilirubin >20 micromol/L or >20% of the total SBR, should have a medical review by the most senior medical officer (same day before discharge from hospital)

Conjugated bilirubin

	Use single light phototherapy if		FAMILY NAME	MKN
NSN USA	<ul> <li>SBR is at or above the phototherapy treatment threshold line</li> </ul>	nt threshold line	GIVEN NAMES	☐ MALE ☐ FEMALE
-1	Use multiple light phototherapy if			- 1
raciniy	<ul> <li>SBR is rising rapidly (&gt;8.5 micromol/L per hour)</li> </ul>	ur)	D.O.B// M.O.	
	• SBR is <50 micromol/L below the RED exchange transfusion line	ange transfusion line	ADDRESS	
NEONATAL JAUNDICE	<ul> <li>SBR fails to respond to single light phototherapy</li> </ul>	apy		
TREATMENT THRESHOLD GRAPH	If the SBR is rapidly rising or approaching the RED exchange	the RED exchange		
32 WEEKS GESTATION	transfusion treatment threshold line an urgent medical review	ent medical review	LOCATION / WARD	
32 WEEKS GESTATION	Silouid occur		COMPLETE ALL DETAILS OR AFFIX PATIENT LABEL HERE	ENT LABEL HERE
Neonate with jaundice <24 hours of age or greater than 14 days of	s of age Table A Risk Factors and Causes of Neonatal Jaundice	es of Neonatal Jaundice		
should have urgent medical review and	Jaundice <24 hours of age - Suspect haemolysis until proven otherwise	spect haemolysis until pr	oven otherwise	
<ul> <li>Measure the SBK and plot on the jaundice treatment threshold graph</li> <li>Urgent medical review will determine when to start phototherapy.</li> </ul>	Jaundice due to haemolysis	Immune - e.g. ABO blood grou Non-immune - e.g. G6PD	Immune - e.g. ABO blood group incompatibility Rhesus disease, Kell, Duffy, anti-E Non-immune - e.g. G6PD	
Consider starting photomerapy at a lower SDK if the neonate has risk factors for neonatal jaundice (see <i>Table A</i> ) or is unwell	Individual neonatal risk factors	Prematurity		
Measure the SBR every		<ul> <li>Aspnyxia</li> <li>Apgar &lt;7 at 5 minutes or a</li> </ul>	Applyxia Appar <7 at 5 minutes or acidosis pH <7 or base excess <12 mEa/l.	
o 6 hours until the SBR is both below the phototherapy treatment	ent	Low serum albumin <30 grams per litre	ams per litre	
threshold line and stable of railing, then o 12-24 hourly for the duration of treatment		<ul> <li>Sepsis or congenital infections</li> <li>Maternal diabetes</li> </ul>	ons	
• Consider		Cephalohaematoma / bruising	ing	
o Additional investigations (see Table B)		G6PD risk with family histo	nistory of stolling with was jaurinited as a recriate G6PD risk with family history or with exposure to trigger	
o Transfer to a higher level facility if appropriate	Table B Additional Investigations			
Neonate with jaundice ≥24 hours of age	Clinical Feature	Investigation		
<ul> <li>Do a transcutaneous bilirubin (TcB) if well and ≥35 weeks or</li> </ul>	Neonate of Rhesiis negative	Blood Groun		
• Do an SBR if		Direct Antidobulin Test (DAT)		
o Unwell or <35 weeks		An immediate SBR is required	An immediate SBR is required if the DAT is positive and the SBR is unknown	
o The TcB is ≥250 micromol/L or	A ridhin colomoi din choncol	Cill blood sound (CDC) and file	- discolution disco	
o The TcB is <20 micromol/L below the treatment threshold line	first 24 hours of age	rull blood count (rbc.) and lilim with reticulocyte Blood group	n with reticulocyte	
<ul> <li>Medical review will determine when to start phototherapy. Consider</li> </ul>	starting OR	DAT		
phototherapy at a lower SBR if the neonate has risk factors for neonatal jaundice (see Table A) or is unwell	Neonate with a rapidly rising total SBR (>8 5 micromol/l ner hour)	Septic screen including blood A G6PD screen if	Septic screen including blood and urine culture & sensitivity if sepsis suspected A GADD screen if	
<ul> <li>If SBR &lt;50 micromol/L below the phototherapy treatment threshold</li> </ul>	line OR	<ul> <li>There is a family history</li> </ul>		
repeat the SBR in 12-24 hours		This is a male neonate with	This is a male neonate with dark hair from a high risk ethnic origin/geographic area e.g. African, Asian	e.g. African, Asian
If SBR >50 micromol/L below the phototherapy treatment threshold	hold line phototherapy threshold	Mediterranean and Middle Eastern descent	Eastern descent	
Continue regular visual assessments  If phototherapy is commonsed measure SBD	Identification of maternal blood group should also be considered with the above investigations	up should also be considere	d with the above investigations	
o After 6 hours to ensure SBR is stable or falling, then	Neonate with a total SBR approaching exchange		Serum albumin level	
First 512 34 Level for the direction of the direction	transfusion thresholds	•	Liver function tests	

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o Transfer to a higher level facility if appropriate o Additional investigations (see Table B)

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Consider

o Every 12-24 hours for the duration of treatment