

**Topic Expert Group:** Medical care and clinical practice

**Neurological monitoring in the high-risk infant: Near-infrared spectroscopy (NIRS)**


**Target group**
- Term and preterm infants at risk for brain injury:
  - Infants with hypoxic-ischaemic encephalopathy (HIE)
  - Infants with encephalopathy for other causes (e.g. metabolic)
  - Infants with suspected or verified seizures
  - Infants requiring intensive care and/or surgery
  - Infants with suspected/confirmed congenital central nervous system (CNS) anomalies
- Parents

**User group**
Healthcare professionals, neonatal units, hospitals, and health services

**Statement of standard**
In order to improve evaluation and outcomes of newborn infants at risk of brain injury, management includes neurological monitoring using a structured, age-appropriate neurological assessment and a range of devices to evaluate brain haemodynamics, oxygen transport, brain function, and imaging, as required.

**Rationale**
Infants requiring neonatal intensive care constitute a high-risk population for developing brain injury, particularly full term and preterm infants exposed to hypoxia-ischaemia, CNS infections, or with congenital anomalies. In the first hours after birth, there is imbalance between blood flow and oxygen supply to the brain due to haemodynamic adaptation during transitional circulation, particularly in the very preterm infant. (1) Low and fluctuating cerebral blood flow are associated with adverse outcomes. (2,3) Experimental models and observational studies confirm that both hyper- and hypoxaemia may cause irreversible brain injury. (4–6) The vulnerability of this population, the severity of underlying clinical conditions, and the complexity of care make continuous, cot-side, and non-invasive monitoring tools valuable. Near-infrared spectroscopy (NIRS) derived regional tissue oxygen saturation of haemoglobin (rStO₂) is an absolute value, which corresponds to mixed blood saturation, used in the clinical setting as a surrogate measure for venous oxygen saturation (SvO₂). (7) Indirect assessment of cerebral blood flow has been shown to correlate with rStO₂. (8) This non-invasive, continuous monitoring system may help to adjust interventions that have effects on blood and oxygen supply to the brain. (9) Bilateral brain monitoring may detect differential perfusion between hemispheres.
Benefits

Short-term benefits
- Reduced burden of cerebral hypo- and hyperoxia in preterm infants in the first 72 h after birth (10,11)
- Improved neuroprotection after asphyxia using combined NIRS and MRI measurements of brain perfusion (12)
- Improved maintenance of theoretically safe cerebral oxygenation levels in infants with congenital heart defects (13)

Long-term benefits
- Reduced all-cause mortality in extremely preterm infants (10)
- Improved long-term outcomes in extremely preterm infants (14)

Components of the standard

<table>
<thead>
<tr>
<th>Component</th>
<th>Grading of evidence</th>
<th>Indicator of meeting the standard</th>
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<tbody>
<tr>
<td>For parents and family</td>
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<tr>
<td>1. Parents are informed by healthcare professionals about the role of near-infrared spectroscopy (NIRS) monitoring.</td>
<td>B (High quality)</td>
<td>Patient information sheet</td>
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<td>For healthcare professionals</td>
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<td>2. A unit guideline on neurological monitoring including NIRS is adhered to by all healthcare professionals, to include</td>
<td>A (High quality)</td>
<td>Guideline</td>
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<td>Newborn infants during resuscitation at birth (≤15 min) (11,15,16)</td>
<td>B (High quality)</td>
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<td>Extremely preterm infants in the first 72 h after birth (9,10,17)</td>
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<td>Asphyxiated newborn infants undergoing therapeutic hypothermia (18,19)</td>
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<td>Infants undergoing surgery with cardio-pulmonary bypass (13,20–22)</td>
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<td>3. Training on NIRS monitoring is attended by all responsible healthcare professionals. (9,17,20,21,23)</td>
<td>A (High quality)</td>
<td>Training documentation</td>
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<td>4. Teams with a focus of interest on neuro-critical care, including neonatologists, neurologists, neurophysiologists, nurses, radiologists, radiographers, and physicists are</td>
<td>B (High quality)</td>
<td>Guideline</td>
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established.

For neonatal unit
5. A unit guideline on neurological monitoring including NIRS is available and regularly updated including standardised operational procedures. (7,9,17,23)
   A (High quality) Guideline
   B (High quality)

For hospital
6. Training on NIRS monitoring is ensured. (7,17,20,21,23)
   A (High quality) Training documentation
   B (High quality)

7. Facilities for NIRS monitoring are provided.
   B (High quality) Audit report

8. An interdisciplinary team for neuro-critical care of high-risk infants in the NICU is supported.
   B (Moderate quality) Audit report

For health service
9. High-risk infants are transferred to NICUs with appropriate neuro-monitoring systems and expertise. (24–26)
   A (High quality) Audit report, guideline

Where to go – further development of care

Further development
Grading of evidence
For parents and family
N/A
For healthcare professionals
- Monitor perioperative NIRS in infants with non-cardiac complex neonatal surgery. (27,28)
   A (Low quality)
For neonatal unit
N/A
For hospital
N/A
For health service
N/A

Getting started

Initial steps
For parents and family
- Parents are verbally informed by healthcare professionals about the role of NIRS.
For healthcare professionals
- Attend training on NIRS monitoring.
- Identify leading healthcare professionals with a focus of interest on neonatal neurological monitoring.

For neonatal unit
- Develop and implement a unit guideline on neurological monitoring including NIRS.
- Develop parental information material about NIRS monitoring also including parent perspectives.
- Provide resources for specific training on NIRS monitoring.

For hospital
- Support healthcare professionals to participate in training on NIRS monitoring.

For health service
- Create systems to effectively transfer high-risk infants to NICUs with appropriate neuro-monitoring systems and expertise.

Description

The NIRS sensor is placed at the forehead avoiding cavities, superior sagittal sinus, intra or extra-cranial huge blood collections, or vascular malformations, if known. Scalp oedema will also influence the quality of the NIRS signal. In the smallest newborn infants and those with poor perfusion states sensor position is rotated to avoid tissue injury related to compression or heat. (7,17)

Commercial NIRS devices incorporate similar technology but different wavelengths and computational algorithms translating changes in light absorption into rStO2 absolute values. (7) Systematic approach has evidenced huge differences in rStO2 according to device or probe (23,29), so that device-specific reference ranges or limits have to be used.

Neonatal resuscitation after birth: Clinical assessment of the newborn infant carries high inter-observer variability particularly when scoring preterm or term infants in need of resuscitation. (30) Oxygen saturation targeting and the use of supplemental oxygen during transition remain controversial topics. (31) The use of pulse oximetry or heart rate monitoring during resuscitation has not led to improvements on the short or long-term outcomes. (32) rStO2 and fractional oxygen extraction reference ranges and percentile charts for the interpretation of cerebral oxygenation during immediate transition to avoid hypo- and hyperoxia of the brain during resuscitation appears promising. (11,15,16) Yet, routine interventions based on rStO2 during resuscitation need development and evaluation.

Extremely low gestational age newborn infants: Recent studies have shown an association of cerebral rStO2 levels and clinical outcomes. (33) Low rStO2 on the first day of life associates surrogated measures of compromised systemic blood flow and risk of intraventricular haemorrhage. (34) Impaired cerebral blood flow autoregulation assessed by NIRS and arterial blood pressure monitoring associates abnormal systemic (and cerebral) blood flow distribution, death and severe brain injury. (35,36) Cerebral oxygenation can be stabilised in the preterm infant during the first 72 hours from birth by the combined use of rStO2-NIRS monitoring and a pathophysiological, brain oriented treatment guideline with no record of severe adverse events. (9,10) The quality of evidence supporting some of the listed statements in the intervention
algorithm is generally low, however, are all routinely used in clinical care of these patients. (9) Although important early surrogate outcomes, such as aEEG at day 3 of postnatal life or neuroimaging, did not significantly differ between the study groups (37,38), post hoc analyses showed that early burden of cerebral hypoxia was significantly associated with low brain electrical activity and severity of intracranial haemorrhage. (14) So far, definitive evidence of benefit for improvement of long-term clinical outcomes is needed as the technology is not cheap, requires manipulation and additional staff time, and may have unwanted effects. (10)

**HIE:** Cerebral hypoperfusion during the first hours after birth is followed by hyperperfusion, even during treatment with moderate hypothermia. Potential differences according to the severity of brain injury (moderate vs severe) have been identified. (12,18) NIRS measurements of oxygenation and MRI measurements of brain perfusion show good correlation. (12) However, the predictive capacity of NIRS changes lacks consistency. (18,19) As yet, widespread recommendation of NIRS monitoring to guide important clinical decisions in asphyxiated newborn infants cannot be made.

**Congenital heart disease (CHD):** NIRS may be a useful adjunct particularly during cardiopulmonary bypass to optimise perfusion. NIRS-derived measures of systemic oxygen balance correlate with global circulatory measures and biochemical indicators of shock. (20) Algorithms have been developed to guide interventions based on rStO₂ values during the perioperative period. (21) However, the current literature on the use of NIRS alone does not demonstrate improvement in neurologic outcome. (22) Prospective data evaluating NIRS findings and relevant outcomes in this population difficult to compare, because of the variable disease physiology, variable baseline values, and small sample sizes. These issues prevent extrapolation to wider CHD population.

Other complex surgical procedures conducted during the neonatal period, such as congenital diaphragmatic hernia or esophageal atresia (27,28), might be additional scenarios where NIRS may play a role to guide surgeons and anesthetists during the intervention procedures.

**Source**


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5 years/next revision: 2023

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