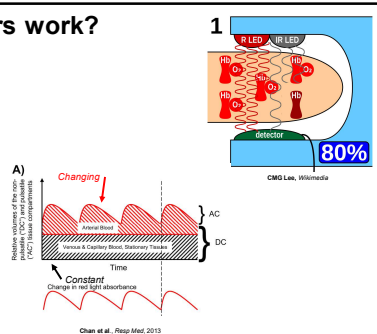


How do pulse oximeters work?

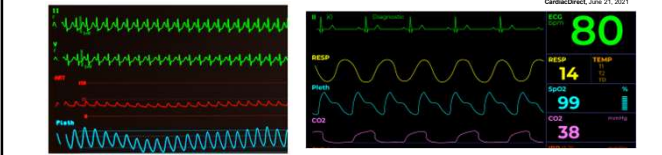
- Pulse oximeters transmit **red** and **infrared (IR)** light through tissue – emitter to detector
- Absorption = emitted light – returned light
- Two components of absorption
 - Non-pulsatile: Skin, bone, fat, muscle, and non-pulsatile blood
 - Pulsatile: arterial blood (mix of oxyhemoglobin (HbO) and deoxyhemoglobin (HbD))
- Ratio of absorbencies compared against calibration table from healthy volunteers to generate SpO₂



7

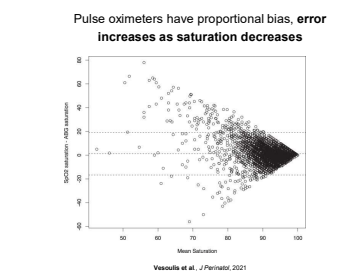
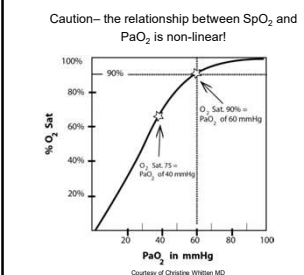
How is pulse ox displayed?

- Absorption of IR light is displayed on patient monitors as the **Pleth waveform**
- Depending on perfusion, may resemble arterial waveform with diastolic notch or may look more like sine wave
- Posted SpO₂ number represents an average (**8-10 seconds**) -- large swings will take time to be reflected



8

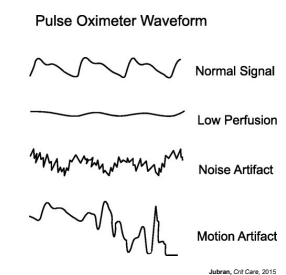
Pulse ox error



9

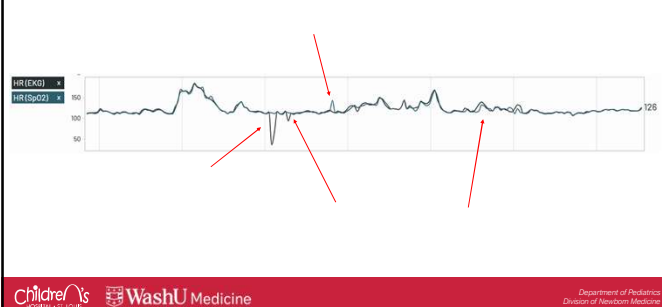
Pleth waveforms – identifying problems

- The pleth waveform is valuable tool for identifying SpO₂ problems
- The appearance of the waveform provides insight about perfusion, motion, noise
 - If SpO₂ number is not posted, pleth waveform can help to diagnose reason
 - If SpO₂ number is posted, pleth waveform can provide information on level of confidence
- Compare ECG and SpO₂ HR – significant discordance suggests unreliable signal



10

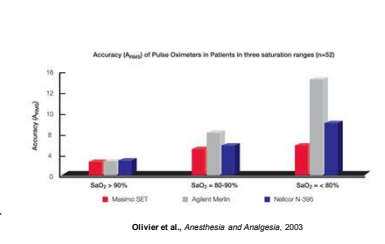
Heart rate – Pulse rate concordance



11

Pulse ox reliability danger zones


- Poor perfusion (e.g., 22-week infant) – pleth wave amplitude too small
 - Brighter light? ↑heat = burns
- Motion
 - Value "hold" during motion
- Lower saturations
 - Exponential increase in error as saturations decrease



12

Role of a pulse oximeter in the NICU

- Non-invasive measure of arterial oxygen saturation (avoids frequent arterial blood gas)
- Can be obtained continuously throughout NICU stay
- Superior to observation alone, cyanosis visually apparent only when SaO₂ < 80%
- Alternative source of heart rate measurement
- Soft and flexible sensor can be positioned on hands, feet, fingers, toes



Reich et al., Case Rep Pediatr, 2013

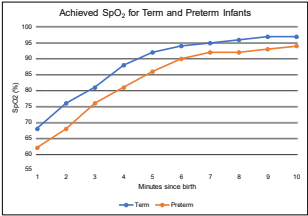
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SpO₂ Use Case – Delivery Room Oxygen Titration

- Oxygen saturations naturally rise following birth as PVR drops and cardiac output increases
- Dawson et al. studied 468 infants and 61,650 SpO₂ points to define reference SpO₂ values by time
 - Helpful in guiding resuscitation!
- Failure to achieve 80% by 5 minutes associated with increased risk of IVH in VLBW (Oei et al., Arch Dis Child, 2016)
- Term infants with severe birth asphyxia (Apgar <4 at 1 minute) have lower SpO₂ than infants with mild asphyxia (Saugstad et al., Acta Paediatr, 2005)



Adapted from Dawson et al., Pediatrics, 2010

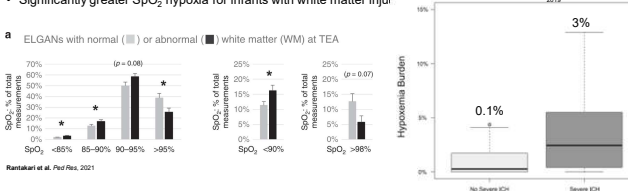
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Hypoxia outcomes – Brain Injury

- Preterm infants are at risk for intraventricular hemorrhage (IVH), cerebellar hemorrhage (CH), and white matter injury (WMI)
- Acute and chronic hypoxia are driving factors behind preterm brain injury
- 30x increase in SpO₂ hypoxia for infants with severe IVH in one study
- Significantly greater SpO₂ hypoxia for infants with white matter injury



Wessells et al., J Perinatol, 2016

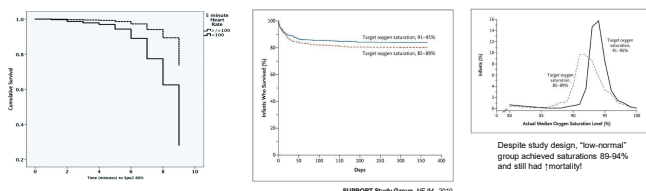
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Hypoxia outcomes – NICU Mortality

- Excessive supplemental oxygen causes ROP, but too little increases risk of death
- Target limits have been intensely debated; SUPPORT trial had surprising finding of increased mortality in "low normal" target group (19.9% vs 16.2%)



Oei et al., Arch Dis Child, 2016


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How are pulse oximeters validated?

- Regulatory clearance provided by through FDA 510(k) clearance pathway, based on ISO 80601-2-61 standard (last updated in 2017)
 - Invasive laboratory testing on healthy adult volunteers
 - Controlled hypoxia between 70-100% SpO₂ using a controlled air-nitrogen-carbon dioxide rebreather circuit with nose pinched shut
 - 10 individuals, at least 2 of which are "darkly pigmented"
 - At least 200 paired SaO₂-SpO₂ samples
- Must achieve A_{95%} ≤ 3 % for approval
 - A_{95%} = $\sqrt{\frac{1}{25} \left(\frac{1}{SpO_2} - \frac{1}{SaO_2} \right)^2}$
 - Combines measures of bias and precision (consistently close to real value)
- Big gaps
 - Most test subjects are repeat volunteers at UCSF, are they even "normal"?
 - Skin tone not assessed in objective way
 - Sick patients are never tested
 - Children (much less neonates) generally not tested
 - "Adult subjects are acceptable in this case due to the uncertainty of determining the accuracy of sensors intended for use in neonates"
 - We found A_{95%} = 8.9% for White neonates, 9.5% for Black neonates
 - Validation data rarely available to the public



Volunteer undergoing testing at UCSF Hypoxia lab


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The current state of neuromonitoring in the NICU, 2025

- Neonates are at risk for stroke, hemorrhage, hypoxic-ischemia, elevated intracranial pressure, and seizures
- Neonates have the highest risk of seizures across the lifespan
- Standard ICU monitoring is comprehensive, EXCEPT FOR THE BRAIN



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
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What is the ideal monitor?

- Continuous, real-time
- Non-invasive
- Organ (brain) specific
- Includes elements of oxygen delivery and consumption
- Regulatory approval for neonates

Cerebral NIRS!



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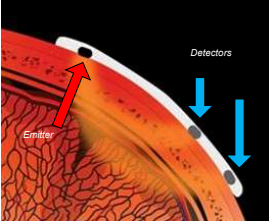
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
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What is NIRS and how does it work?

- Near-infrared light can penetrate through skin, bone, connective tissue
 - Depth depends on spacing between emitter/detector
- NIR light is scattered, reflected, absorbed – but some returns to the sensor
 - Scatter and reflection remain constant
 - HbD absorbs more red, less NIR light
 - HbO absorbs more NIR, less red light
- NIRS measures oxygen saturation in all tissue spaces
 - Most blood is in venous space (70% venous, 25% arterial, 5% capillary) so NIRS is a **venous weighted measure**
- Second shallow detector removes superficial blood flow signal



Santi Barritt, Harvard Medical School



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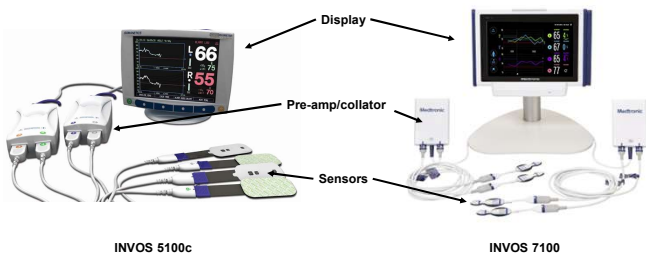
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
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NIRS hardware anatomy



INVOS 5100c INVOS 7100



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NIRS sensor placement

- Cerebral (temporal placement to right or left of midline)
- Splanchnic (right or left lower quadrant)
- Renal (right or left flank)
- Most monitors support up to 4 simultaneous channels

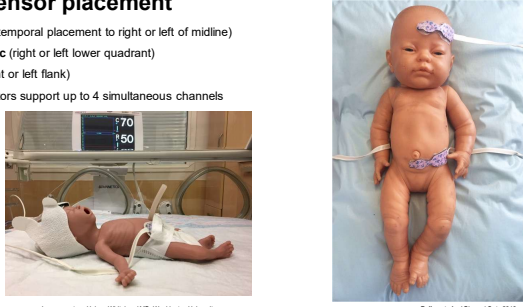



Image courtesy Helene Whitehead MD, Washington University *Bailey et al., J Biomed Opt. 2016*



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
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NIRS is not just a pulse oximeter

Pulse oximeter	NIRS
Uses red and NIR light to measure oxygenation	Uses red and NIR light to measure oxygenation
Non-invasive continuous monitor	Non-invasive continuous monitor
Measures of hypoxia are associated with bad outcomes	Measures of hypoxia are associated with bad outcomes
Transmission-based measure, limited to thin body parts	Reflectance-based measure, can be applied to any tissue
Arterial saturation: measure of oxygen supply	Mixed-venous saturation: measure of oxygen delivery and consumption
Detects oxygenation problems in whole body	Detects oxygenation problems in specific tissue

↑ SpO₂ = too much supplemental oxygen
↓ SpO₂ = not enough supplemental oxygen

↑ rSO₂ = too much supplemental oxygen or decreased extraction by tissue
↓ rSO₂ = not enough supplemental oxygen or increased extraction by tissue



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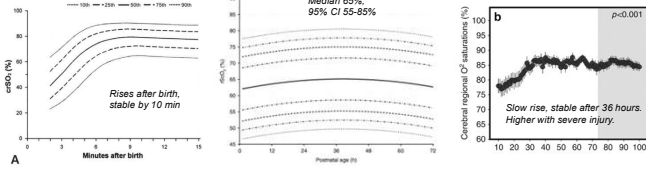
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
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What is a "normal" cerebral NIRS value?

- Large reference studies have been done in delivery room, VLBW, HIE populations
- rSO₂ 55-85% (median 65%) widely used in preterm population as "normal" bounds
- These values were established using INVOS 5100c and small adult sensor. **INVOS neonatal sensor equivalent is 63-93% (other device-sensor combinations will vary)**



Prober et al., J Pediatrics, 2013 *Alberici et al., Pediatr Res, 2015* *Ferman et al., Pediatr Res, 2017*



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Fractional tissue oxygen extraction (FTOE)

- The fractional tissue oxygen extraction (FTOE) is the percentage of delivered oxygen that is consumed
- It can be calculated from the NIRS signal (rSO₂) and the pulse oximeter (SpO₂)
- Should be about 20% in healthy humans
- Influenced by functional metabolism, inflammation, injury

$$FTOE = \frac{SpO_2 - rSO_2}{SpO_2}$$

FTOE = 0.305

Infants with chorioamnionitis (shaded bars) have lower rSO₂ and higher FTOE.

Roescher et al., *Pediatr Res*, 2015; Chen et al.

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Between-device differences in NIRS

- Conceptually NIRS monitors all work the same way
- BUT each device has subtle differences in hardware and software which result in **different** measurements on the **same** patient
- Kleiser et al. have evaluated a broad range of devices using a blood lipid phantom and have published transforms to convert measures

Blood-lipid phantom

SpO₂ (%) (INVOS)

rSO₂ (%) (INVOS)

NIRS Device	Hypoxic threshold %
FORESIGHT small	66
FORESIGHT non-adhesive small	67
NIRO small	61
NIRO small re-usable	63
NIRO large	62
NIRO large re-usable	62
INVOS neo	63
ScenSmart neo 8004CB-NA	66

- Kleiser et al., *Biomed Opt Express*, 2017
- Kleiser et al., *Adv Exp Med Biol*, 2016
- Hansen et al., *NEJM*, 2023

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Head size may make a difference

a

GA (weeks)	HC (cm)	Diameter (cm)	Radius (cm)	NIRS probe location (2.5 cm measure - red arch)
22	19.5	6.2	3.1	Ventricle
24	21.4	6.8	3.4	Ventricle
26	24.5	7.8	3.9	Ventricle
28	25.3	8.1	4.0	Ventricle
30	27.0	8.6	4.3	Ventricle
31	28.4	9.0	4.5	Corpus callosum/ventricle
32	29.5	9.4	4.7	Corpus callosum/ventricle
33	29.8	9.5	4.7	Corpus callosum/ventricle
34	30.4	9.7	4.8	Anterior limb/white matter tract
35	31.7	10.1	5.0	Anterior limb/white matter tract
36	32.0	10.2	5.1	White matter tract
37	32.0	10.2	5.1	White matter tract
38	32.3	10.3	5.1	White matter tract

b

Kabak et al., *Pediatr Res*, 2023

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What causes cerebral hypoxia?

Oxygen carrying capacity
Systemic oxygenation
Cardiac output

Supply = Demand
Balanced

Brain activity
Growth

Supply < Demand
Hypoxia

Anemia
Respiratory Disease
Shock

Brain activity
Growth
Seizures
Inflammation From injury

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NIRS and anemia

- We studied 39 healthy VLBW infants (no IVH, no congenital anomalies) with paired cerebral NIRS recording and hemoglobin measurement
- Cerebral saturation drops, FTOE increases as hemoglobin decreases
- Potential implications for pRBC transfusion threshold- what FTOE threshold risks injury?

Mean of FTOE

Hemoglobin (g/dL)

Change in FTOE

Hemoglobin (g/dL)

63% threshold, -2 SD

Halana Whitehead

WashU Medicine

Whitehead et al., *J Perinatol*, 2018

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NIRS and IVH

- Cerebral oxygenation patterns in IVH have two phases
 - Hyperoxia** (impaired autoregulation, temporary decrease in O₂ demand by injured tissue)
 - Hypoxia** (increased O₂ demand from inflammation, seizures, ↑metabolism)

Acute ↑ rSO₂

Hyperoxia

Hypoxia

↓ rSO₂ after bleed

Beauchamp et al., *Sci Rep*, 2018

Altshuler et al., *J Pediatr*, 2012

Verheijen et al., *Stroke*, 2010

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Cerebral hypoxia over time and injury

- We made 1287 NIRS recordings from 187 VLBW infants, each 6-8 hours long
- Cerebral saturations drop over time (with anemia)
- Infants with IVH have lower saturations, remain lower for at least 60 days
- High grade IVH has more cerebral desaturation than mild IVH or no IVH

Source: Vesoulis et al., Pediatric Res, 2021

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Silent cerebral hypoxia

- Apnea-bradycardia-desaturation (ABD) events occur in 100% of infants < 32 weeks and may cause impaired cerebral autoregulation
- Recent evidence suggests presence of **significant silent cerebral hypoxia** (low rScO₂ with normal SpO₂)
- We made 1616 recordings from 209 infants with mean length 5.1 hours
 - Daily intermittent cerebral hypoxia (ICH) events increased from 32/day at birth to 49/day at 21 days of life
 - Most events (48%) were silent, increased proportion in infants with IVH (67%)

Source: Fairchild et al., Pediatric Res, 2018

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Silent cerebral hypoxia and respiratory support

- Many NICUs are focusing on less invasive respiratory support, more rapid weaning guided by pulse oximetry and blood gases
- We made 1013 recording from 174 infants over the entire NICU course, mean rScO₂ and hypoxia burden were evaluated
- Infants on lowest respiratory support (room air, low flow nasal cannula) had highest cerebral hypoxia burden, no difference in SpO₂ values

Source: Monro Norozi-Cleaver, J Perin, 2023

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Cerebral autoregulation

Source: Adapted from Meng et al., Anesthesiology, 2015

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Balancing factors in cerebral autoregulation

- The goal of cerebral blood flow is **sufficient oxygen delivery for maintenance of cerebral metabolism**
- Optimally, changes in CBF are driven **ONLY** by changing metabolic needs
- In reality, CBF can be influenced by many factors
 - pCO₂
 - Medications (especially sedation)
 - Systemic blood pressure/cardiac output
 - Position
- Theoretical control mechanisms
 - Myogenic (stretch)
 - Neurogenic (autonomic)
 - Metabolic

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What is “time domain” ?

- Examines the correlation **over time** between two variables **without** mathematical transformation
- Cerebral blood flow and cardiac output should be independent (zero correlation)
- Relationship can be examined in blocks, over time

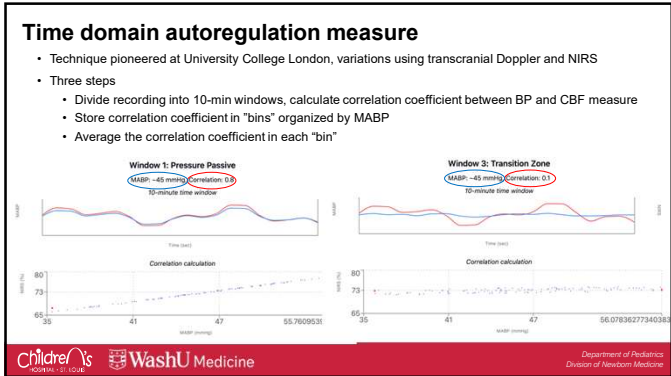
Source: Adapted from Meng et al., Anesthesiology, 2015

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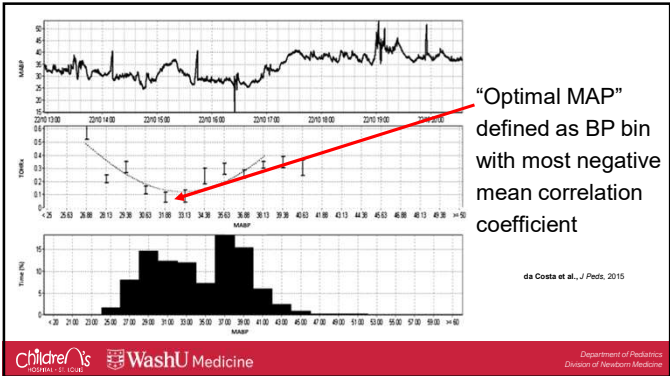
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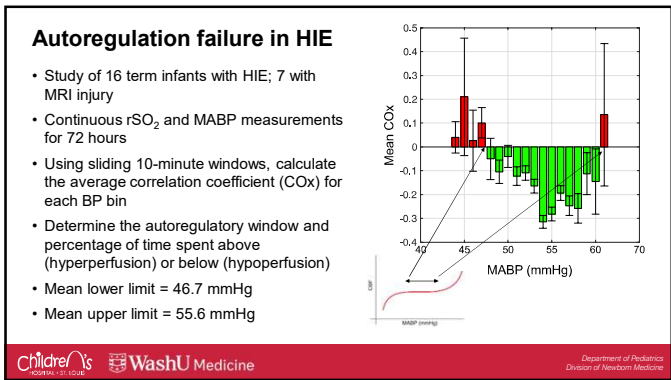
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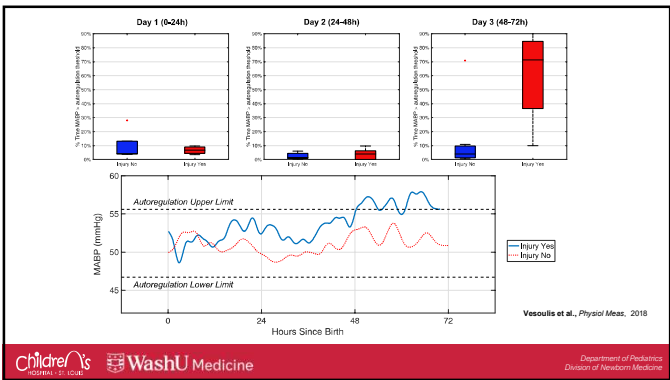
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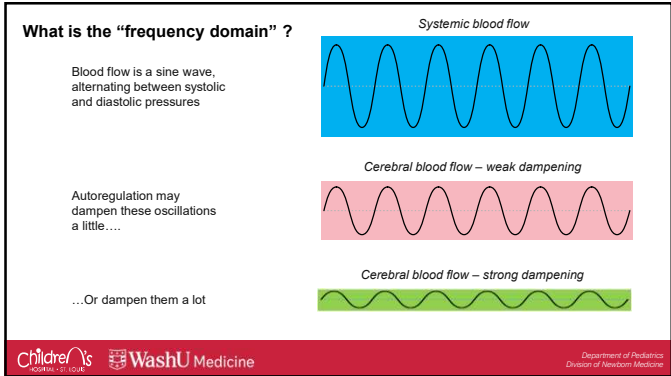
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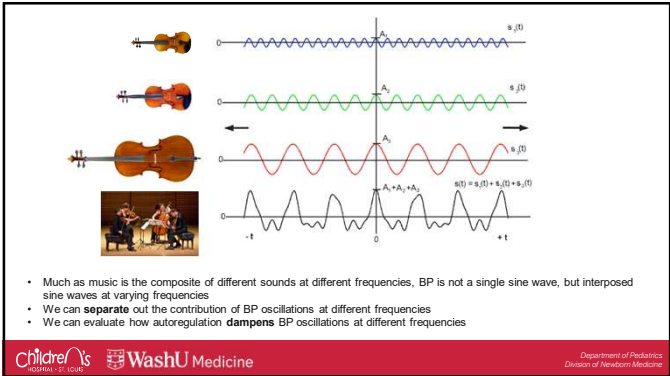
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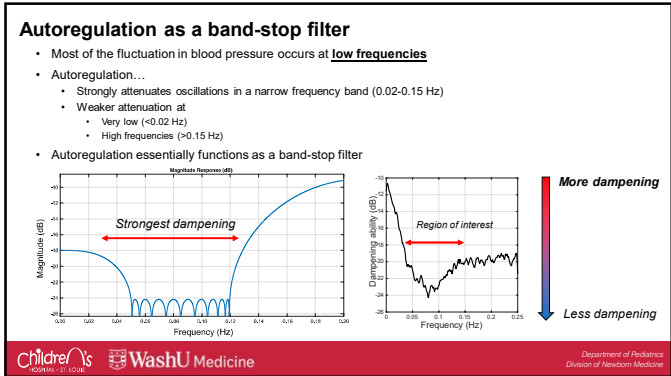
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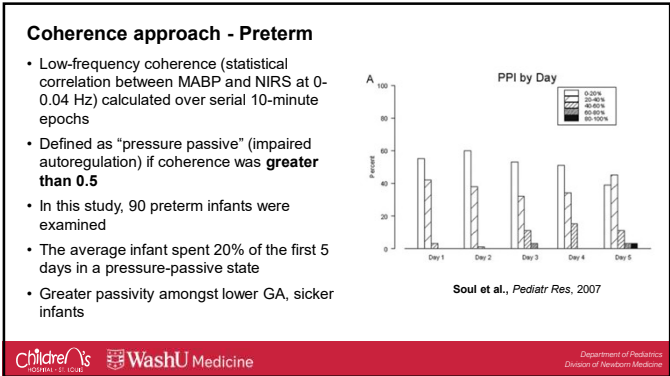
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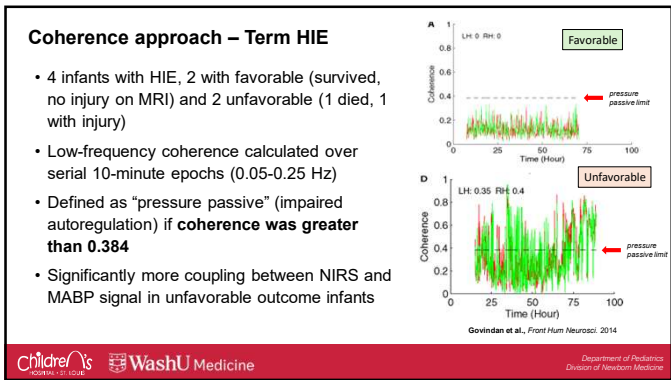
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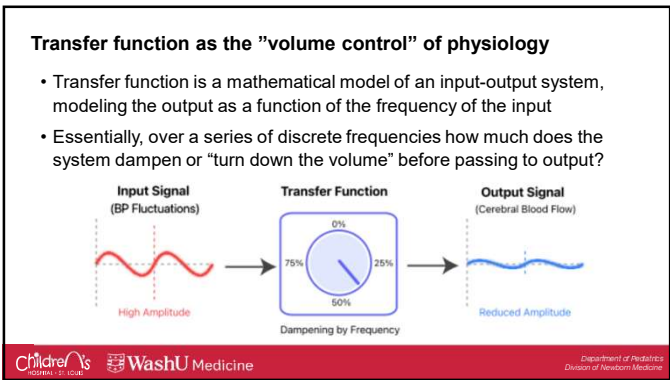
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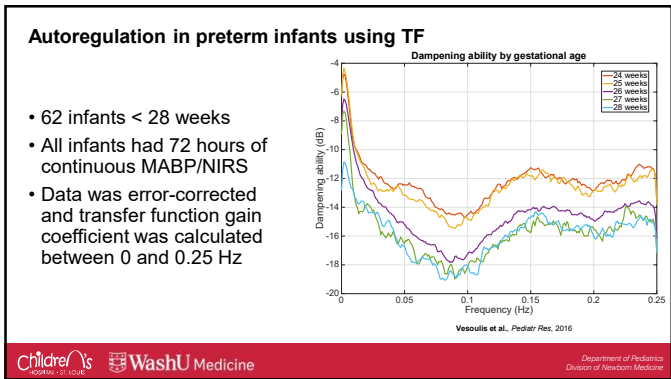
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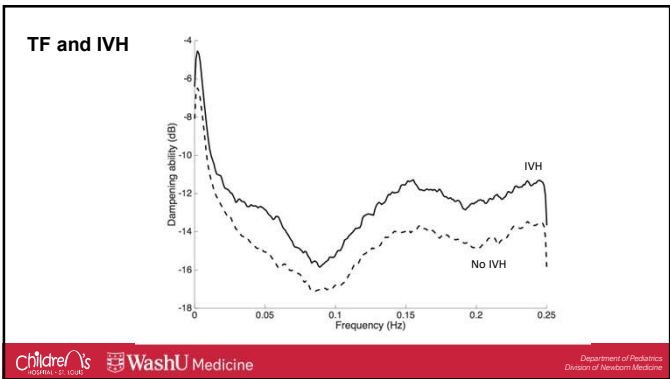
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NIRS and NEC

Bella Anadiak

van der Heide et al., Pediatr Res, 2021

- NIRS applied at first suspicion of NEC in 75 babies
- Infants with NEC had lower saturations and less variability
- NEC ruled out (100% specificity) if rSO2 > 87.8%
- NEC ruled in (100% specificity) if rSO2 < 51.3%

No normative values in preterm infants!

- N=194 infants <32 weeks, weekly monitoring 1976 recordings
- Overall pattern resembles Hb in anemia of prematurity
- Infants who later develop NEC have early drop in intestinal oxygenation

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Renal NIRS and AKI

Persistent rSO2 after birth related AKI

Matt Harer

Harer et al., Pediatr Nephrol, 2021

Credit et al., Preprint

- NIRS is also useful for identifying "silent" kidney injury
- Laboratory markers lag AKI, NIRS provides early recognition
- As with brain injury, renal injury induces persistent tissue hypoxia
- Recent study shows strong correlation between decreasing renal rSO2 and increasing risk of AKI

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Renal NIRS and PDA

- Useful for detection of hemodynamically significant PDA
- Two studies
 - Renal rSO2 < 43% is 85% sensitive, 83% specific (Underwood et al., Neonatology, 2007)
 - Renal rSO2 < 66% is 81% sensitive, 77% specific (Chock et al., Pediatr Res, 2016)
- rSO2 values are restored to normal values after treatment

Wolf et al., JNIRS, 2012

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If NIRS monitoring is so great, why isn't it more widely used?

-RCT evidence not compelling
-Implementation challenges

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COSGOD-II

- 2-center RCT of 60 infants < 32 weeks GA
- Randomized to NIRS visible/not visible in delivery room
- Cerebral hypoxia burden reduced by 55% in NIRS visible group

Pechter et al., J Pediatr, 2016

COSGOD-III

- 11-center RCT of 607 infants < 32 weeks GA
- Randomized to NIRS visible vs. standard of care for first 15 minutes after birth
- Improved survival without brain injury for NIRS group (82.9 vs 78.5%) but not statistically significant

Pechter et al., BMJ, 2023

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SafeBoosC-II

- 8-center RCT of 166 infants < 28 weeks GA
- Randomized to NIRS visible/not visible for 72h after birth
- Standardized interventions for abnormal rSO2
- Combined hyperoxia/hypoxia burden reduced by 58% in NIRS group
- Reduction in mortality or severe brain injury (25% vs 14%, not statistically significant)

Hyatt-Sorenson et al., BMJ, 2015

SafeBoosC-III

- 70-center RCT of 1601 infants < 28 weeks GA
- Randomized to NIRS visible vs. standard of care for 72h after birth
- No difference in mortality or severe brain injury on ultrasound (35.2 vs 34%)
- Small non-significant improvement in MRI score (Kidokoro score 2 vs. 3, p=0.11)

Hansen et al., NEJM, 2023

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Three challenges ahead

- **Devices are not validated against outcomes**
 - FDA requires monitoring devices to accurately measure **current physiology** against gold standards (heart rate, SaO₂, SvO₂, etc.)
- **RCTs study protocols, not devices**
 - Represent decisions about population, timing, threshold, and choice of interventions
- **Provider trust is essential, an ignored “box in the corner” does not help the patient!**

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Didn't trust it – an example from the literature

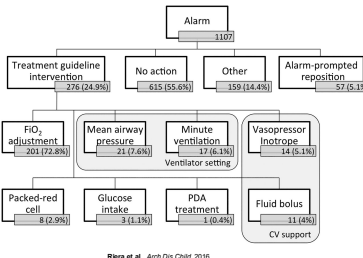
- 74 infants randomized
- Hgb-based threshold
 - 14 g/dL for FiO₂ > 0.35
 - 12 g/dL for FiO₂ < 0.35
 - Any time for VS instability
- NIRS-based threshold
 - Fractional tissue oxygen extraction (FTOE) > 47%
- FTOE improved significantly after transfusion, regardless of arm
- No difference in number of transfusions, death, LOS, ROP
- Failure of randomization, nearly 70% of transfusions in NIRS arm were given on “clinical basis” before hitting NIRS threshold



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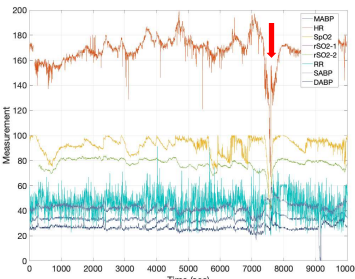
Didn't use it or used it wrong – an example from the literature

- SafeBoosC-II, 8-center phase II RCT of NIRS visible vs. not visible
- Treatment algorithm with 10 different options
- Number and type of interventions per site were tracked
- More than half of alarms were ignored
- 73% of interventions were **↑FiO₂**



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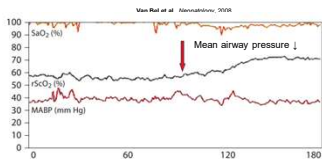
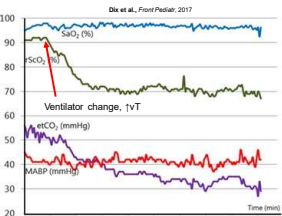
Case example 1



Bradycardia event (arrow) has broad impact on BP, SpO₂, rSO₂. NIRS doesn't provide much additional information here.

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Case example 2: Ventilator management

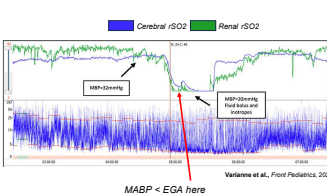


This preterm infant is on high-frequency oscillatory ventilation. rSO₂ monitoring demonstrates saturations at threshold with normal SpO₂ and MABP. Mean airway pressure was decreased (red arrow) and rSO₂ improved into normal range.

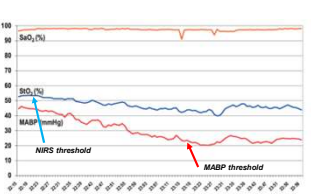
The tidal volume was increased on this child's ventilator and resulted in overventilation. Rapid cerebral desaturation follows pCO₂, which would otherwise be undetectable without NIRS (normal HR, normal BP, normal SpO₂).

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Case example 3: Hypotension



In this infant with **septic shock**, an early decrease in renal rSO₂ is noted, despite normotension. A rapid drop in MABP is accompanied by falling cerebral and renal rSO₂. After fluid bolus and inotropes, a recovery of renal and cerebral rSO₂ is noted.



This 24-week infant gradually becomes hypotensive. Standard treatment would indicate intervention for MABP < 24 mmHg (red arrow). NIRS-guided management would indicate intervention nearly 1 hour earlier (blue arrow). Note SpO₂ remains normal the entire time.

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Case examples 4: Necrotizing enterocolitis

2400g term infant with pulmonary valvulopathy atresia with MAPCAs awaiting surgical treatment when larger size. Diagnosed with NEC (pneumatosis, hematochezia). NIRS monitoring started 48 hours later, showed low-normal cerebral rSO2 and mesenteric rSO2 at lower limits of detection. Gradually improved and feeding restarted once normal.

26-week preterm infant, donor twin of twin-twin transfusion syndrome. Abdominal distention and pneumatosis at 2 weeks. Mesenteric rSO2 drops with onset of pneumatosis, continuing for multiple days, improving only after resection of necrotic bowel and washout.

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Case examples 5: ECMO complications

- Term neonate with meconium aspiration syndrome placed on VA ECMO for hypoxic respiratory failure.
- Cerebral and renal NIRS (green and blue, bottom panel) gradually decline over a 24-hour period.
- Ultrasound revealed large pericardial effusion which was drained with rapid improvement in cerebral and renal rSO2.
- Note that MABP, SpO2 remained normal throughout, subtle tachycardia.

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Case examples 6: More ECMO complications

- Term neonate placed on VA ECMO for congenital diaphragmatic hernia and hypoxic respiratory failure.
- Marked decrease in renal rSO2 (blue bottom panel) with preserved cerebral rSO2 (green)
- pRBC transfusion given (first red arrow) with transient improvement, second pRBC transfusion given (second red arrow) also with transient improvement.
- Abdomen became rigid and discolored, exploratory laparotomy demonstrated 400 mL abdominal hematoma

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NIRS monitoring indications

- Preterm infants
 - First 72 hours for early recognition of acute cerebral hypoxia related brain injury (IVH)
 - Identifying chronic hypoxia over NICU course, especially silent cerebral hypoxia
 - Evaluating response to treatments (hypotension, ventilator, transfusion, PDA)
- Term infants
 - HIE
 - Early detection of acute kidney injury
 - Adequate cerebral/renal perfusion, safety monitor during ECMO
 - Hemodynamic management during anesthesia

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NIRS-directed hemodynamic management

- NIRS provides regional circulation monitoring and the opportunity to provide tissue-specific, directed interventions
- For many neonates, reference ranges (63-93%) can provide helpful target values
- For all patients, a lower threshold of 45% should be strictly avoided
- Baseline-Bottomline approach for critical care
 - Establish rSO2 baseline in awake, stable patients
 - Monitor deviation from baseline, use corroborating evidence to provide directed interventions to maintain baseline values or higher
 - Decrease from baseline of more than 20% (bottom line) associated with adverse outcomes

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Conclusions

- Hypoxia is an important driver of neonatal disease
- Pulse oximeters have a high degree of utility in clinical practice, but we must be cognizant of pitfalls
- NIRS is a distinct tool with a distinct use; it isn't just a fancy pulse oximeter
- There is compelling observational data that cerebral hypoxia is associated with true pathology -- this has not translated to successful NIRS RCTs

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SafeBoosC-III Intervention Algorithm

[illegible]

Footnote: There are not well-established guidelines for normative blood pressure in preterm infants. The most common conversion is the Dubow definition (1) which targets a mean arterial blood pressure = gestational age in weeks (e.g., 34 weeks for infant born at 24 weeks GGA). However, treatment decisions should not be based solely on a number, but rather the combination of cardiac deceleration, hypoxemia, and evidence of impaired perfusion.

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