Visual light spectroscopy reflects flow-related changes in brain oxygenation during regional low-flow perfusion and deep hypothermic circulatory arrest

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Objectives: Regional low-flow perfusion has been used to minimize ischemic brain injury during complex heart surgery in children. However, optimal regional low-flow perfusion remains undetermined. Visible light spectroscopy is a reliable method for continuous determination of capillary oxygen saturation (SgvO2). We used visible light spectroscopy to follow deep and superficial brain SgvO2 during cardiopulmonary bypass, regional low-flow perfusion, and deep hypothermic circulatory arrest.

Methods: Visible light spectroscopy probes were inserted into the superficial and deep brain of neonatal (3.9-4.5 kg) piglets, targeting the caudate and thalamic nuclei. The piglets were subjected to cardiopulmonary bypass and cooled to a rectal temperature of 18°C using pH stat. Regional low-flow perfusion was initiated through the innominate artery at 18°C, and pump flows were adjusted to 40, 30, 20, and 10 mL/kg/min for 10-minute intervals followed by 30 minutes of deep hypothermic circulatory arrest. Regional low-flow perfusion was reestablished, and flows were increased in a stepwise manner from 10 to 40 mL/kg/min. SgvO2 was continuously monitored. Carotid flow was measured using a flow probe, and cerebral blood flow (milliliters per kilogram body weight per minute) was calculated.

Results: There were no significant differences between the deep and superficial brain tissue oxygenation during regional low flow brain perfusion before deep hypothermic circulatory arrest. However, after deep hypothermic circulatory arrest, the superficial brain SgvO2 was lower than the deep brain SgvO2 (24 ± 12 vs 55.3 ± 8, P = .05, at flows of 30 mL/kg/min, and 34.2 ± 17 vs 62.5 ± 8, P = .06, at a flow rate of 40 mL/kg/min). During regional low-flow perfusion, SgvO2 was maintained at flows of 30 to 40 mL/kg/min (cerebral blood flows of 15 to 21 mL/kg/min and 19 to 24 mL/kg/min, respectively), but was significantly lower at pump flows of 20 mL/kg/min (cerebral blood flow of 10 to 14 mL/kg/min) and 10 mL/kg/min (cerebral blood flow of 5 to 9 mL/kg/min) compared with the values obtained just before regional low-flow perfusion (pre–deep hypothermic circulatory arrest, 37 ± 6 vs 65.5 ± 4.4, P < .05, and 21.6 ± 3.7 vs 65.5 ± 4.4, P < .01, respectively; and post–deep hypothermic circulatory arrest, 32 ± 4.5 vs 65.5 ± 4.4, P < .05, and 16.6 ± 4.7 vs 65.5 ± 4.4, P < .01, respectively).

Conclusions: Regional low-flow perfusion at pump flows of 30 to 40 mL/kg/min with resulting cerebral blood flows of 14 to 24 mL/kg/min was adequate in maintaining both deep and superficial brain oxygenation. However, lower pump flows of 20 and 10 mL/kg/min, associated with cerebral blood flow of 9 to 14 mL/kg/min, resulted in significantly reduced SgvO2 values.

Deep hypothermic circulatory arrest (DHCA) has facilitated complex cardiac surgery, especially in small patients, enabling the surgeon to attain a bloodless operative field so precise anatomic reconstruction can be achieved. However, the use of DHCA has been associated with both immediate and
late adverse neurodevelopmental outcomes.\textsuperscript{1-6} Although multifactorial in origin, hypoxic-ischemic brain injury is the most likely cause of these outcomes.

Over the decades, cardiopulmonary bypass (CPB) hardware has improved, and safer alternatives to DHCA continue to be explored. More recently, surgical techniques for the repair of complex intracardiac and aortic arch lesions during CPB without the use of DHCA have been developed to minimize the risks of ischemic brain injury.\textsuperscript{7-11} Regional low-flow perfusion (RLFP) is one such alternative to DHCA in which CPB flow is maintained to the brain.\textsuperscript{10} The use of RLFP to maintain continuous cerebral oxygen delivery seems intuitively rational; however, many questions remain relating to the optimal management of the standard set of variables associated with perfusion practice including flow rates, arterial blood gas management, and optimal hematocrit.

Despite the technical challenges involved in the arterial cannulation for RLFP in small neonates and infants, uninterrupted cerebral oxygen delivery is the goal. Newer technologies, such as near-infrared spectroscopy (NIRS), that allow real-time, continuous measurement of cerebral oxygen saturation in the operating room during CPB have shown that cerebral saturation is maintained during RLFP.\textsuperscript{12} However, NIRS is an indirect measure of global cerebral oxygen delivery. Visible light spectroscopy (VLS) is an emerging technology that has recently become available for continuous determination of capillary oxygen saturation (S\textsubscript{vgO2}). Unlike NIRS, VLS tissue oximetry uses shallow-penetrating visible light to measure microvascular hemoglobin oxygen saturation (S\textsubscript{vgO2}) in small, thin-tissue volumes. When VLS technology is compared with the standard NIRS, VLS oximetry measures small, subsurface tissue volumes; in contrast, NIRS measures larger, deeper volumes of tissue.\textsuperscript{13} We used VLS probes in the brain to track S\textsubscript{vgO2} during CPB, DHCA, and variable RLFP flows in a neonatal piglet model.

**Materials and Methods**

Animals received humane care in compliance with the “Principals of Laboratory Animal Care,” formulated by the National Society of Animal Research, and the “Guide for the Care and Use of Laboratory Animal,” prepared by the Institute of Laboratory An-

### Abbreviations and Acronyms

- CPB = cardiopulmonary bypass
- DHCA = deep hypothermic circulatory arrest
- NIRS = near-infrared spectroscopy
- RLFP = regional low-flow perfusion
- S\textsubscript{vgO2} = capillary oxygen saturation
- VLS = visible light spectroscopy

**Visual Light Spectroscopy and Tissue Oximetry**

The VLS oximeter (T-Stat, model 303) used in this study was supplied by Spectros Corporation (Portola Valley, Calif). The probes in this study consisted of a 1-mm needle tip for deep brain measurements or a 3 \times 3-mm disc probe for the superficial brain measurements (Figure 2, A, B). Briefly, the VLS probe emits white light from a probe and collects any light returning to the probe from the tissue. The collected light is separated by wavelength into 2048 bins, measured simultaneously. For oximetry, the blue-to-yellow (476-584 nm) portion of the visible spectrum is then used to solve the light scattering and the concentration of each of the major forms of hemoglobin (deoxyhemoglobin, oxyhemoglobin, and optionally methemoglobin and carboxyhemoglobin), using first differential spectroscopy and least-squares fitting to known hemoglobin spectra. Tissue hemoglobin is estimated as deoxyhemoglobin + oxyhemoglobin, and the tissue hemoglobin oxygen saturation (S\textsubscript{vgO2}) is determined as oxyhemoglobin/deoxyhemoglobin + oxyhemoglobin. The oximetry measurements are continuous, with each measurement typically requiring 5 to 50 ms, depending on the intensity of the reflected light.\textsuperscript{13} The use of VLS technology in human tissue ischemia detection has been validated.\textsuperscript{13,14} Benaron and colleagues\textsuperscript{13} evaluated the ability of VLS to detect hypoxemia and ischemia in human subjects. Clinical hypoxemia has been evaluated in healthy human subjects using 3 types of probes: buccal, esophageal, and rectal VLS probes. Hy-
poxemia was induced by subjecting the volunteers to a mixture of air and helium, while pulse oximetry was monitored at the left index finger. Clinical regional ischemia was induced by increasing tourniquet pressure at the base of the forefinger or by exerting direct pressure. Clinical global ischemia was evaluated using buccal VLS probes in patients undergoing implantation of an automatic implantable defibrillator. VLS probes were found to be sensitive to hypoxemia, regional, and global ischemia.13

Cardiopulmonary Bypass Circuit and Experimental Protocol

After placement of VLS probes, the animals were turned supine, the carotid artery and jugular vein were isolated at the neck, and a 2-mm flow probe was placed around the left carotid artery to measure continuous carotid flow during the experiment. Cerebral blood flow was determined by doubling the measured carotid flow because both carotid arteries are perfused in the pig during RLFP.

After aseptic skin preparation, the animals were draped in a sterile fashion and a midline sternotomy was performed. The heart and great vessels were exposed, and after heparinization (400 IU/kg), the innominate artery was cannulated with a 10F arterial cannula and an 18F straight 2-stage venous cannula inserted into the right atrial appendage (Medtronic Bio-Medicus, Minneapolis, Minn), and CPB was initiated.

The CPB circuit consisted of a roller pump, a membrane oxygenator (Medtronic, Minimax Plus, Medtronic, Minneapolis, Minn), and sterile quarter-inch tubing. The circuit was primed with blood previously harvested from a donor pig mixed with crystalloid prime solution (Normosol R, Abbott Laboratories, North Chicago, III), to maintain hematocrit no lower than 30%. In addition, methylprednisolone (Solu-Medrol, Pfizer, New York, NY) (30 mg/kg), heparin 2500 units, mannitol (0.5 g/kg), and sodium bicarbonate (20 mL) were added to the priming solution.

CPB was initiated with the aid of vacuum-assisted venous drainage. Additional fentanyl (10 µg/kg) and pancuronium (0.1 mg/kg) were administered to the piglet, and 1% isoflurane was continued on the pump. Core cooling was commenced at a pump flow of 200 mL/kg/min using pH stat arterial blood gas manage-

Statistical Analysis

Statistical analysis was performed using GraphPad InStat version 3.00 for Windows 95 (GraphPad Software, San Diego Calif). SgvO2 recorded at deep and superficial brain sites for specific RLFP flows were charted and compared using one-way analysis of variance for multiple comparisons followed by Dunnett’s post hoc test.

Results

In 6 of the 8 animals, we obtained both deep and superficial SgvO2 measurements. We were unable to acquire reliable SgvO2 in 1 animal, and the data were not used for analysis. The superficial brain SgvO2 probe malfunctioned in 1 animal. Inaccurate measurements were subsequently noted to be the result of blood clots at the tip of the measuring catheter occluding light transmission and absorption.

No significant difference was found between deep and superficial brain tissue oxygenation during RLFP before DHCA. After DHCA, the superficial brain displayed lower
CSP

RLFP flow of 20 mL/kg/min deep brain SgvO2 tended to decline before DHCA. Flows of 10 mL/kg/min caused severe desaturations in the deep and superficial (cortical) brain. Flow reduction before DHCA did not demonstrate significant differences among the caudate nucleus, thalamus, and superficial brain. In contrast, tissue oxygenation (SgvO2) in the superficial brain regions after DHCA was lower at all pump flow rates. This may reflect a higher oxygen extraction by superficial brain (cortical areas). Alternatively, perfused tissue oxygenation because of failure of autoregulation, as reflected by lower SgvO2 values even at pump flows of 30 to 40 mL/kg/min, may be a contributing factor. The clinical implication of these observations is that flows of 30 to 40 mL/kg/min, which are often used in clinical practice, may be inadequate to perfuse all regions of the brain when intermittent brain perfusion is used during DHCA.

In pigs the innominate artery gives rise to both carotid arteries and to the right subclavian artery, whereas in humans the innominate artery gives rise to a carotid artery and a subclavian artery; thus, the flow delivered through the innominate artery is not the same in the human and in the piglet. The direct correlation of carotid flow with pump flow and deep cerebral tissue oxygenation could have significant clinical impact for the management of RLFP in human subjects. In our experiment the carotid flow was found to be approximately one third of the total pump flow. Because both carotids are perfused in the piglet during RLFP, cerebral blood flow was determined by doubling the measured carotid flow. Values reflect tissue oxyhemoglobin percentage. Values are expressed as means and standard error of means. RLFP, Regional low-flow perfusion; DHCA, deep hypothermic circulatory arrest.

**TABLE 1. Flow-related changes in brain oxygenation during regional low-flow perfusion and deep hypothermic circulatory arrest**

<table>
<thead>
<tr>
<th>SgvO2 (%)</th>
<th>Baseline</th>
<th>Cooling</th>
<th>RLFP 40</th>
<th>RLFP 30</th>
<th>RLFP 20</th>
<th>RLFP 10</th>
<th>DHCA</th>
<th>RLFP 10</th>
<th>RLFP 20</th>
<th>RLFP 30</th>
<th>RLFP 40</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deep 1 (caudate)</td>
<td>37.5 ± 17.5</td>
<td>73 ± 5.6</td>
<td>70.6 ± 7</td>
<td>52.1 ± 5.6</td>
<td>40.5 ± 6.2</td>
<td>19.3 ± 11*</td>
<td>5.8 ± 1</td>
<td>18.3 ± 6.6*</td>
<td>33.1 ± 13*</td>
<td>54.5 ± 10</td>
<td>69.4 ± 8.5</td>
</tr>
<tr>
<td>Deep 2 (thalamus)</td>
<td>47.2 ± 7.5</td>
<td>56 ± 4.1</td>
<td>50 ± 6.3</td>
<td>44 ± 6.3</td>
<td>32.8 ± 8.7</td>
<td>23 ± 7*</td>
<td>4 ± 2.4</td>
<td>14.2 ± 7.4*</td>
<td>31.6 ± 10.4*</td>
<td>46.2 ± 11.2</td>
<td>46.4 ± 11</td>
</tr>
<tr>
<td>Superficial</td>
<td>53 ± 6</td>
<td>75 ± 4</td>
<td>69 ± 7.2</td>
<td>64.3 ± 8</td>
<td>54 ± 13</td>
<td>32 ± 20</td>
<td>3.6 ± 3.6</td>
<td>13 ± 8</td>
<td>17 ± 11</td>
<td>27 ± 16</td>
<td>38 ± 22</td>
</tr>
<tr>
<td>Cerebral blood flow</td>
<td>28 ± 4.3</td>
<td>24 ± 5</td>
<td>24 ± 5.8</td>
<td>21 ± 6</td>
<td>14 ± 4.6</td>
<td>9.5 ± 3.3</td>
<td>0</td>
<td>5 ± 1.3</td>
<td>10 ± 1.7</td>
<td>15 ± 2.2</td>
<td>19.7 ± 2.2</td>
</tr>
</tbody>
</table>

Deep and superficial brain tissue saturations (SgvO2) during cooling, RLFP, and DHCA. Cerebral blood flow was determined by doubling the measured carotid flow. Values reflect tissue oxyhemoglobin percentage. Values are expressed as means and standard error of means. RLFP, Regional low-flow perfusion; DHCA, deep hypothermic circulatory arrest; SgvO2; capillary oxygen saturation. *P < 0.05 compared to baseline.
bral flows were determined by doubling the measured carotid flow. Cerebral flows of 10 ± 1.7 to 14 ± 4.6 mL/kg/min and 5 ± 1.3 to 9.5 ± 3.3 mL/kg/min correlated with pump flows of 20 mL/kg/min and 10 mL/kg/min, respectively. On the basis of our data, during RLFP, unilateral carotid flows higher than 14 to 20 mL/kg/min seem to be adequate in maintaining cerebral tissue oxygenation.

The normal cardiac output in a neonate at normothermia is 200 mL/kg/min, and the brain takes 20% of the normal cardiac output; therefore, cerebral blood flow at normothermia is approximately 40 mL/kg/min. By using the Q10 relationship that links metabolic rate to temperature, it is easy to estimate brain blood flow requirements at various degrees of hypothermia. These calculations are consistent with the results of our present study.

Relatively little is known about cerebral blood flow at 18°C in the neonate. Most of the data have been obtained from animal models of total body perfusion at low temperatures and extrapolated to humans. In a group of children undergoing cardiac surgery, Kern and colleagues clini-

cally demonstrated that a reduction of 45% to 70% in pump flow at 18°C to 20°C significantly reduced cerebral blood flow and CMRO₂ but did not change O₂ extraction, suggesting that at deep hypothermia (despite a significant re-
duction in pump flow rates) cerebral blood flow and cerebral oxygen supply exceed cerebral metabolic needs.

With the use of NIRS in 6 neonates undergoing RLFP, Pigula and colleagues demonstrated that to maintain baseline cerebral saturation, regional perfusion had to be maintained at 20 mL/kg/min. Children undergoing DHCA alone showed significantly greater decreases in cerebral oxygen saturations (−33.5 ± 14.6 vs −0.8 ± 5.2, P = .02) and change in cerebral blood volume index (−19.2 ± 14.3 vs −1.4 ± 2.7, P = .003) compared with neonates supported with RLFP. However, our observations demonstrate that even flows greater than 20 mL/kg/min after DHCA may be inadequate to maintain cerebral oxygenation.

Because it averages values obtained from large tissue volumes, NIRS technology may be unable to detect SgvO₂ gradients within the brain. We propose that VLS technology, at least experimentally, may offer significant improvements over NIRS in that it measures smaller tissue volumes and can detect subtle changes in saturation.

**Study Limitations**

Although the VLS technology is reliable and easy to use, and other noninvasive probes have been used in human studies, the present study could be undertaken only in an animal model because of the invasive nature of the probes used. Therefore, this study cannot be validated in humans. To validate our findings and advance the field, other accurate, readily available, easy to use, noninvasive technologies need to be refined.

Deep and superficial brain SgvO₂ were measured using different VLS probes (disc vs needle probes), which may account for some of the differences in measurements. Nevertheless, both disc and needle probe measurements correlated well during cooling, showing a difference only during rewarming post-DHCA, suggesting that the data are valid and accurate.

**Conclusions**

In a neonatal piglet model of RLFP at 18°C, we demonstrated that pump flow rates of 30 to 40 mL/kg/min and cerebral flows of 14 to 24 mL/kg/min provide adequate deep and superficial brain oxygenation. Although there were no significant differences between deep and superficial brain SgvO₂ during RLFP, after DHCA significant changes between superficial and deep brain oxygenation were noted that varied with pump flow. On the basis of the available data we believe that RLFP flows in human subjects should be kept between 30 and 40 mL/kg/min, because that results in carotid flows of approximately 15 to 20 mL/kg/min. Future studies should focus on optimizing other variables used to control RLFP, such as RLFP pressures, temperatures, and the “safe” duration of RLFP.

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