The use of Invos™ somatic oximetry to measure variations in placental tissue oxygenation in laboring healthy term parturients with epidural analgesia: an observational study

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Purpose: Near-infrared spectroscopy provides a non-invasive continuous real-time monitoring of tissue oxygen saturation. As uterine contractions during labor may be associated with a transient uteroplacental hypoperfusion, this prospective, observational study investigates the ability of near-infrared spectroscopy to detect variation in uteroplacental oximetry during uterine contractions.

Patients and methods: Four Invos™ oximetry probes (Medtronic®, Minneapolis, MN, USA) per subject were applied on the placental surface (PLA), the abdomen (MYO), the forearm (ARM) and the leg (LEG), of twenty healthy laboring parturients with epidural analgesia. Measurements of mean tissue oxygen saturation and area under the curve (AUC) were made during 60 minutes. The primary outcome was the difference of the AUC measurements between the PLA probe and the MYO probe.

Results: The AUC values for the PLA and MYO probes were not different. The mean saturation values recorded by the PLA probe were not different from the other probes.

Conclusion: The Invos monitor was unable to detect variations in uteroplacental saturation during labor in healthy parturients.

Keywords: near-infrared spectroscopy, uteroplacental perfusion, Invos™ monitor, tissue oxygenation

Introduction
Obstetric decision-making algorithms to evaluate the fetus well-being and assess the indication for cesarean delivery could benefit from the near-infrared spectroscopy (NIRS), as a continuous, non-invasive, real-time monitoring of fetal oxygen delivery. The rationale behind its use follows the idea that the uterine contractions during labor cause a cyclic, transient reduction in myometrial and uteroplacental blood perfusion, and could therefore serve as an in vivo human clinical model of transient uteroplacental hypoperfusion which could be detected by variations in tissue oxygen saturation.

The primary objective of this novel concept is to evaluate the feasibility and the usefulness of the NIRS to detect the variations in the uteroplacental oxygen saturation during the contraction.

Patients and methods
After approval by the Maisonneuve-Rosemont Hospital Research Ethics Committee (approval # 12117) and written informed consent received, a convenience sample of
20 laboring healthy (American Society of Anesthesiologists physical status II) term parturients were included, between April and July 2013, according to the following inclusion criteria: normal pregnancy, anterior placenta, planned vaginal birth, term pregnancy (≥37 weeks of gestation), and desire for epidural analgesia. Patients were excluded if they presented with exclusion criteria: contraindication to neuraxial analgesia, utero-placental insufficiency as diagnosed by the attending obstetrician, abruptio placentae, abnormal placentation, twin pregnancy, planned cesarean delivery or unplanned cesarean delivery, morbidly obese patient and refusal to participate. Demographic data such as age, height, weight, parity, color of the skin and position of the placenta were collected at the clinic.

In the labor ward, the attending anesthesiologist performed a standardized epidural at the L2–L3 or L3–L4 interspace upon maternal request. Maternal blood pressure, heart rate and pulse oximetry were collected every 5 minutes until the end of the study. The fetal heart rate as well as the frequency and length of uterine contractions were measured with a cardiotocographic device every 5 minutes. A control ultrasound was performed by an experiment investigator to delineate the location of the placenta and measure the distance, oxytocin administration) and AUC measurements.

An analysis of variance for repeated measures test with Bonferroni’s multiple comparison test was applied, when data were normally distributed. Clinical trial registry: Clinicaltrial.gov (NCT01834599).

Results

Subject characteristics are shown in Table 1. Table 2 reports AUC results for each probe. There was no difference between the AUC values for the PLA and MYO probes at room air, nor on 100% oxygen.

Breathing 100% oxygen did increase the mean tissue oxygen saturation values from breathing room air as measured by all of the probes. The lowest mean tissue oxygen saturation of tissues in a 1.5 cm³ sample located ~1.7 cm under the skin’s surface.2 Extensive reviews of the mechanisms underlying Invos NIRS technology are provided in several articles.3–5 The Invos monitor provides cumulative tissue oxygen saturation every 5 seconds below a baseline threshold – the area under the curve (AUC) measure (mm%) – for 3 of the 4 probes (PLA, MYO, ARM). The rationale for using the AUC measure in this study is that during uterine contractions the transiently reduced oxygen delivery-to-extraction ratios could be detected as an increase in the AUC values for these organs in comparison with non-myometrial or non-uteroplacental tissue, such as skeletal muscle.

The primary outcome for this feasibility study is the difference of the AUC measurements between the PLA probe and the MYO probe. Secondary outcomes are the difference in mean tissue saturation values for each probe, and associations between clinical factors (color of skin, skin-to-placental distance, oxytocin administration) and AUC measurements.

Table 1 Subjects characteristics

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>27.6 (4.6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMI (kg/m²)</td>
<td>28.9 (2.6)</td>
</tr>
<tr>
<td>Gestational age (weeks)</td>
<td>40 (39–40)</td>
</tr>
<tr>
<td>Primiparity</td>
<td>10 (50)</td>
</tr>
<tr>
<td>Skin color</td>
<td>17 (85)</td>
</tr>
<tr>
<td>Black</td>
<td>3 (15)</td>
</tr>
<tr>
<td>S-P distance (cm)</td>
<td>7 (35)</td>
</tr>
<tr>
<td>S-P distance &gt;1.7 cm²</td>
<td>5 (25)</td>
</tr>
</tbody>
</table>

Notes: *Mean (standard deviation), *Median (interquartile range). %N (%).
Abbreviation: S-P, skin-to-placenta.

Table 2 AUC measurements, for probe PLA, MYO and ARM, while patients were breathing room air and 100% oxygen

<table>
<thead>
<tr>
<th>Probe</th>
<th>Room air</th>
<th>Oxygen</th>
<th>P-value</th>
</tr>
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<tbody>
<tr>
<td>PLA (mm%)</td>
<td>60.40 (31.38)</td>
<td>54.55 (23.44)</td>
<td>0.4311</td>
</tr>
<tr>
<td>MYO (mm%)</td>
<td>56.80 (44.00)</td>
<td>44.50 (30.99)</td>
<td>0.2270</td>
</tr>
<tr>
<td>ARM (mm%)</td>
<td>127.70 (63.15)</td>
<td>132.70 (64.19)</td>
<td>0.6568</td>
</tr>
</tbody>
</table>

Notes: AUCs were presented as mean (SD). An analysis of variance for repeated measures test with Bonferroni’s multiple comparison test was applied.
Abbreviations: AUC, area under the curve; PLA, placental surface; MYO, the abdomen; ARM, the forearm; LEG, the leg.
Measure placental tissue oxygenation with the Invos™

Discussion

Our study found lack of variation in tissue saturation with uterine contractions for the placenta and the myometrium.

Our hypothesis was that the PLA probe would measure placental blood flow in the intervillous space, where fetal-maternal gas exchange takes place. We suggested that this would have proven more likely in subjects where the distance between the skin and the myometrial-placental edge, as measured by the ultrasound examination, was <1.7 cm.

This study had specific technological limitations, such as the skin-to-placenta distance of 1.7 cm or less in 65% of the subjects, the presence of bony structures, which is not taken into consideration in the calculation algorithms included in the Invos monitor, and the fact that the Invos was originally created to function on tissues where the blood network is predominantly venous, which contrasts with the unique anatomy of the placenta as a large arterio-venous shunt. The findings of our study were negative and it is thus difficult to know whether it is because there is truly no change in uteroplacental blood flow, or whether the Invos technology is simply unable to accurately determine the uteroplacental blood oxygen saturation.

Yet, assuming the accuracy of the Invos, other explanations for our negative results could be proposed. First, reduced tissue vascular resistance, induced by the epidural sympathetic blockade, leading to an increase in regional blood flow would result in an increase in tissue saturation.

Second, Brar et al suggested that fetoplacental blood flow is uninterrupted during uterine contractions, this is supported by Fairlie et al who showed that there was no change in the fetal umbilical artery pulsatility index as labor progressed in healthy parturients with a normal fetal heart rate, and despite a reduction in uteroplacental blood flow with uterine systole, placental oxygen delivery exceeds fetal oxygen extraction. So, under such excessive blood flow, it is possible that small variations in the oxygen delivery-to-extraction ratio may cause negligible shifts in placental pO2 and tissue saturation.

Our study should be repeated in laboring pre-eclamptic women, especially when intrauterine growth restriction is present or when labor is complicated by fetal bradycardia. In these parturients, uteroplacental blood flow may be significantly impaired and the oxygen delivery-to-extraction ratio may be pathological, translating into wider shifts in placental pO2 between uterine contractions. These patients may constitute a better model for transient uteroplacental hypoperfusion, which may be detected by the Invos monitor.

Acknowledgment

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Disclosure

The authors report no conflicts of interest in this work.

References


