Persistently Altered Brain Mitochondrial Bioenergetics After Apparently Successful Resuscitation From Cardiac Arrest

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Background—Although advances in cardiopulmonary resuscitation have improved survival from cardiac arrest (CA), neurologic injury persists and impaired mitochondrial bioenergetics may be critical for targeted neuroresuscitation. The authors sought to determine if excellent cardiopulmonary resuscitation and postresuscitation care and good traditional survival rates result in persistently disordered cerebral mitochondrial bioenergetics in a porcine pediatric model of asphyxia-associated ventricular fibrillation CA.

Methods and Results—After 7 minutes of asphyxia, followed by ventricular fibrillation, 5 female 1-month-old swine (4 sham) received blood pressure–targeted care: titration of compression depth to systolic blood pressure of 90 mm Hg and vasopressor administration to a coronary perfusion pressure >20 mm Hg. All animals received protocol-based vasopressor support after return of spontaneous circulation for 4 hours before they were killed. The primary outcome was integrated mitochondrial electron transport system (ETS) function. CA animals displayed significantly decreased maximal, coupled oxidative phosphorylating respiration (OXPHOSCI+CIII) in cortex (P<0.02) and hippocampus (P<0.02), as well as decreased phosphorylation and coupling efficiency (cortex, P<0.05; hippocampus, P<0.05). Complex I– and complex II–driven respiration were both significantly decreased after CA (cortex: OXPHOSCI, P<0.01, ETSCII, P<0.05; hippocampus: OXPHOSCI, P<0.03, ETSCII, P<0.01). In the hippocampus, there was a significant decrease in maximal uncoupled, nonphosphorylating respiration (ETSCII+CIII), as well as a 30% reduction in citrate synthase activity (P<0.04).

Conclusions—Mitochondria in both the cortex and hippocampus displayed significant alterations in respiratory function after CA despite excellent cardiopulmonary resuscitation and postresuscitation care in asphyxia-associated ventricular fibrillation CA. Analysis of integrated ETS function identifies mitochondrial bioenergetic failure as a target for goal-directed neuroresuscitation after CA. IACUC Protocol: IAC 13-001023. (J Am Heart Assoc. 2015;4:e002232 doi: 10.1161/JAHA.115.002232)

Key Words: acute brain injury • brain • cardiac arrest • electron transport system • mitochondria • neuroprotection

Pediatric cardiac arrest (CA) is an important public health problem, with thousands of children sustaining one of these potentially devastating events each year in the United States. Although survival rates are improving, many survivors have neurologic sequelae, resulting in a major burden for patients, families, and society. The brain is particularly susceptible to ischemic injury because of its high metabolic energy demand and limited intrinsic energy supply.

Alterations in mitochondrial bioenergetics may play a pivotal role in secondary neurologic injury cascades, sometimes also referred to as ischemia–reperfusion injury, initiated by CA in mature animals. Several laboratories report critical alterations in adult brain mitochondrial bioenergetics after ischemia and reperfusion, including diminished mitochondrial mass, population number, decreased function of the electron transport system, reduced antioxidant enzyme activity and content, and damaged and oxidized lipid membrane parameters.

Because the immature brain has unique energetics compared with older animals, the effects of CA on cerebral mitochondrial bioenergetics may be different in immature animals.

To address the paucity of data regarding mitochondrial alterations in the immature brain after CA and cardiopulmonary
resuscitation (CPR), we performed a detailed evaluation of mitochondrial bioenergetics in a large animal model of hypoxic ventricular fibrillation (VF) pediatric CA that results in good survival rates.9,10 We implemented high-resolution respirometry to ascertain integrated global mitochondrial function, as well as relative contributions of individual electron transport complexes by using specific substrate and inhibitor combinations, together with measures of mitochondrial functional content. We hypothesized that despite good survival parameters after CA and excellent CPR with explicit, protocolized postresuscitative care, pediatric animals would have global persistent alterations in cerebral mitochondrial bioenergetics, mitochondrial content, and specific patterns of individual electron complex dysfunction.

Materials and Methods

All procedures were approved by The Children’s Hospital of Philadelphia’s Institutional Animal Care and Use Committee. Four-week-old female piglets (N=9, 8 to 10 kg), which have neurodevelopment comparable to that of a human toddler, were used for the study.11,12 Piglets were designated into 2 cohorts: (1) the CA group (n=5) killed 4 hours after CPR and return of spontaneous circulation (ROSC), ≈5 hours after the initiation of anesthesia; and (2) anesthetized sham animals (n=5) with no CA and no CPR killed 5 hours after the induction of anesthesia.

Animal Preparation

A porcine model was chosen because of the similarities between swine and humans in regard to anterior–posterior chest depth and compression characteristics (eg, chest stiffness), because the neuroanatomy and neurodevelopment are similar to those of a human toddler, and because of improved survival with the use of blood pressure–targeted CPR in an asphyxia-associated VF CA.10,11,13,14 To keep the population as homogeneous as possible, we did not investigate male animal responses based on prior work.15 All piglets were anesthetized and mechanically ventilated on a mixture of room air and titrated isoflurane (=1.0% to 2.5%) with a tidal volume of 10 to 12 mL/kg, positive end-expiratory pressure of 6 cm H2O, and titration of rate to maintain end-tidal carbon dioxide at 38 to 42 mm Hg (NICO, Novametrix Medical Systems Inc). High-fidelity, solid-state, micromanometer-tipped catheters (MPC-500; Millar Instruments) were used to measure continuous aortic and right atrial pressures, respectively. A bipolar pacing catheter (Edwards Lifesciences) was used to induce VF. All catheter placements were confirmed with the use of fluoroscopy. Unfractionated heparin 200 U/kg was provided to prevent catheter clotting. All animals received a 20 mL/kg 0.9% normal saline bolus intravenously before we obtained baseline measurements to replace overnight fasting deficits. Animals were prepared in a similar fashion as detailed in our previous publications.16,17

Experimental Protocol: CA Animals

Our group has established that hemodynamically targeted CPR titrated to arterial blood pressure improves short-term survival compared with standard CPR targeted to a fixed depth of compressions and a fixed frequency of vasoactive medications after hypoxic and normoxic VF CA.15,16 This goal-directed approach optimizes coronary perfusion pressure (CPP) and cerebral perfusion pressure and presumably coronary and cerebral blood flow during CPR. In addition, our well-established protocol used in this experiment was designed to mimic resuscitation occurring in an intensive care unit after a respiratory deterioration with an initial documented rhythm of VF (Figure 1).10,16 We selected 7 minutes of asphyxia, plus 10 minutes of CPR, for a total 17 minutes of CA because this insult is clinically relevant and because our prior experience suggested that the majority of these animals would be successfully resuscitated so that we could examine mitochondria 4 hours after the injury.15,16 This model maintains important clinical relevance because most in-hospital pediatric arrests occur in invasively monitored intensive care unit patients.17,18 Ten minutes of CPR was performed before the first defibrillation attempt to mimic the median duration of clinical CPR in the in-hospital CPR setting.10,18,19

Asphyxial and Resuscitation Periods: CA Animals

We selected 7 minutes of asphyxia, plus 10 minutes of CPR for a total 17 minutes of CA because this insult is both clinically relevant and our prior experience suggested that the majority of these animals would be successfully resuscitated so that we could examine mitochondria 4 hours after the injury.17 This model maintains important clinical relevance because most in-hospital pediatric arrests occur in invasively monitored intensive care unit patients.18,19 Endotracheal tube clamping was confirmed by absence of exhaled CO2 (Time 0). After 7 minutes of tracheal tube clamping with severe arterial hypoxemia, VF was induced and maintained by electrical pacing to ensure that the animal would not have ROSC from CPR alone before 10 minutes of CPR.20,21 Once VF was confirmed, the tracheal tube was unclamped and CPR was started. Compressions were provided with a target rate of 100 min−1 and ventilations at 6 min−1 with 100% oxygen. To ensure rigorous adherence to CPR rate goals, a metronome was used. Brief, planned interruptions in CPR of 4 seconds every 2 minutes mimicked pulse checks/rhythm analyses. Animals received arterial blood pressure–targeted care.

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defined as a compression depth titrated to a target systolic blood pressure of >90 mm Hg and, starting at minute 9 of the protocol, vasopressor administration (epinephrine 0.02 mg/kg and vasopressin 0.4 U/kg) if and only if the CPP was <20 mm Hg. After 10 minutes of CPR (minute 17 of the protocol), the initial 200-J biphasic waveform defibrillation attempt was provided. Resuscitation continued until sustained ROSC was attained in the CA animals or until minute 27 of the protocol (an additional 10 minutes of resuscitation post initial defibrillation attempt). If ROSC was attained, the animals then received explicit, protocolized post-CA care (see later).

Post-CA Care: CA Animals

After ROSC, animals received explicit protocolized intensive care, including (1) titration of inspired oxygen concentration to maintain oxygen saturation at 92% to 96%; (2) titration of ventilation to maintain end-tidal carbon dioxide between 38 and 42 mm Hg; and (3) intravenous infusions of dopamine (up to 20 μg/kg per minute) to maintain mean arterial pressure >55 mm Hg. Anesthesia was maintained with inhaled isoflurane (≈1.0%). Blood pressure, end-tidal carbon dioxide, oxygen saturation, temperature, and anesthesia were maintained by using the identical explicit post-ROSC protocol as the treatment group.

Sample Acquisition for Mitochondrial Assessments

At 4 hours post ROSC in CA animals (5 hours post induction of anesthesia) and 5 hours post induction of anesthesia in sham animals, a bilateral craniectomy was performed to expose the brain. Tissue was rapidly extracted from 2 regions of interest (a 2-cm² region of left frontal cortex and both hippocampal regions) while simultaneously receiving a sodium pentobarbital overdose (150 mg/kg). Tissue was removed in <10 seconds and placed in ice-cold isolation buffer (320 mmol/L sucrose, 1.21 g/L tris base, and 2 mmol/L EGTA), where blood, blood vessels, or necrotic tissue was dissected and disposed from the cortex and hippocampus. In addition, subcortical white matter was isolated from the cortex for study. Tissue was then gently homogenized on ice in MiR05 (110 mmol/L sucrose, 0.5 mmol/L EGTA, 3.0 mmol/L MgCl₂, 60 mmol/L K-lactobionate, 10 mmol/L KH₂PO₄, 20 mmol/L taurine, 20 mmol/L HEPES, and 1.0 g/L fatty acid–free BSA) by using a 5-mL Potter-Elvehjem Teflon-glass homogenizer (5 strokes over 30 seconds, 3 sets with a pause of 30 seconds between each
set) to a concentration of 1 mg wet weight tissue/10 μL MiR05 buffer. We then compared each measure per milligram of tissue (pmol O$_2$/s×mg) and per citrate synthase activity (pmol O$_2$/s×mg×CS) in sham and CA animals, region by region.

**Mitochondrial Assessments: High-Resolution Respirometry**

A final concentration of tissue homogenate suspended in MiR05 solution was analyzed at a 1 mg/mL concentration, at a constant 37°C, and the rate of oxygen consumption was measured by using a high-resolution oxgyraph and expressed in pmol/[s×mg of tissue homogenate] (OROBOROS, Oxygraph-2K, and DatLab software; all from OROBOROS Instruments). All experiments were performed at 75 to 220 μmol/L oxygen, and reoxygenation was performed routinely before addition of the complex IV electron donor as described later. The oxgyraph was calibrated daily: oxygen concentration was automatically calculated from barometric pressure and MiR05 oxygen solubility factor was set at 0.92 relative to pure water. A substrate, uncoupler, inhibitor titration protocol (Figure 2) previously used for rodent brain tissue was further developed and optimized in previous studies in porcine brain tissue. Respiratory capacities with tissue22 was further developed and optimized in previous studies in porcine brain tissue. 23 Respiratory capacities with electron flow through both complex I and complex II, as well as the convergent electron input through the Q-junction (complexes I and II), were evaluated sequentially by using succinate and NADH-linked substrates. Plasma membranes were permeabilized with the detergent digitonin to allow non-membrane-permeable substrates and ADP access to the subpopulations of mitochondria trapped within synaptosomes. Further, to achieve similar results in brain tissue homogenates and isolated brain mitochondria, with a combination of both subpopulations, Pecinova and colleagues demonstrated that digitonin is necessary in brain homogenate preparations. Without the addition of digitonin, oxidative phosphorylation capacity would likely be greatly underestimated. Thus, in preliminary experiments, a careful digitonin dose titration was completed, in the presence of exogenously administered cytochrome c, which did not induce any significant effect on respiration with the digitonin dose used in the present study, indicating intact integrity of the outer mitochondrial membrane (data not shown). The same concentration of digitonin was used for both sham and injured tissue. Routine mitochondrial respiration was established by the concomitant addition of malate (5 mmol/L) and pyruvate (5 mmol/L), followed by ADP (1 mmol/L) and glutamate (5 mmol/L), to measure the oxidative phosphorylation capacity of complex I (OXPHOS$_{Cl}$ [where Cl is complex I]), driven by the NADH-related substrates. Sequential additions followed: Succinate (10 mmol/L) was added to stimulate maximal phosphorylating respiration capacity via convergent input through complexes I and II (OXPHOS$_{Cl+ClI}$ [where ClI is complex III]). Oligomycin, an inhibitor of ATP-synthase, induced mitochondrial respiration independent of ATP production across the inner mitochondrial membrane, commonly referred to as LEAK respiration (LEAK$_{Cl+ClI}$) or state 4$_o$. Maximal convergent nonphosphorylating respiration of the ETS (ETS$_{Cl+ClI}$) was evaluated by titrating the protonophore, carbonyl cyanide p-(trifluoromethoxy)phenylhydrazone until no further increase in respiration was detected. Rotenone was added to inhibited complex I–driven respiration and remaining measured complex II–driven respiration; the ETS capacity through complex II alone (ETS$_{ClI}$) was measured. The complex III inhibitor antimycin-A (1 μg/mL) was added to measure the residual oxygen consumption that is independent of the ETS. This residual value was subtracted from each of the measured respiratory states to report ETS function. Finally, complex IV activity was determined by the addition of ascorbate (0.8 mmol/L) and N,N,N,N',N'-tetramethyl-p-phenylenediamine (0.5 mmol/L), an electron donor to complex IV. Due to the high level of auto-oxidation of N,N,N,N',N'-tetramethyl-p-phenylenediamine, sodium azide (10 mmol/L), an inhibitor of complex IV, was added, and the remaining chemical background was subtracted to assess complex IV activity. The substrate, uncoupler, inhibitor titration protocol was identical for shams and injured tissue. Tissue harvest to completion of respirometry measurements were completed in all animals in <1.5 hours. Varying the order of the substrate, uncoupler, inhibitor titration protocol additions did not affect individual respiratory state values. Integrated ETS analysis with internal

![Image](https://via.placeholder.com/150)
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normalization was generated using flux control ratios (FCRs) calculated by dividing each respiratory state by the maximal uncoupled mitochondrial respiration (ETS_{CI+CII}).

Citrate Synthase Determination

On completion of the high-resolution respirometry measurements, chamber contents were frozen at −80°C for subsequent citrate synthase (CS) activity quantification (Citrate Synthase Assay Kit, CS0720; Sigma). CS activity (given in micromoles per milliliter per minute) was used as a marker of brain metabolism and surrogate for mitochondrial content or mass secondary to its location within the mitochondrial matrix and its ability to catalyze the condensation of oxaloacetate and the acetyl coenzyme-A, yielding citrate and coenzyme-A in the first step of the tricarboxylic acid cycle. Currently, to our knowledge, there is limited research on biomarkers of mitochondrial content (CS activity, cardiolipin content, mitochondrial DNA content, complexes I to IV protein, and complexes I to IV) that validates these surrogates against morphologic measures after acute brain injury, such as transmission electron microscopy. In addition, after ROSC post-CA care, vital signs were arterial systolic blood pressure 99±0.6 mm Hg, arterial diastolic blood pressure 64±0.5 mm Hg, CPP 39±0.8 mm Hg, end-tidal carbon dioxide 40±0.1 mm Hg, and pulse oximetry 94±0.4%. There was no statistical difference in mean vital signs between CA and sham animals in the post-CA period.

Results

Physiologic and Hemodynamic Variables

In the CA animals, average PaO₂ after 6.5 minutes of endotracheal tube–clamped asphyxia (Time 6.5) was 11±3 mm Hg, and after 4.5 minutes of CPR (Time 11.5), it was 110±14 mm Hg. Hemodynamic measurements (ie, arterial systolic blood pressure, diastolic blood pressure and CPP) during CPR in the CA animals demonstrated adherence to experimental protocol goals (Figure 3). All 5 CA animals studied survived to 4 hours post ROSC. Median number of vasopressor doses given during the CPR period was 5.5 (range 4 to 6). All animals achieved sustained ROSC on first defibrillation attempt. No additional vasopressor support was needed post ROSC. In addition, after ROSC post-CA care, vital signs were arterial systolic blood pressure 99±0.6 mm Hg, arterial diastolic blood pressure 64±0.5 mm Hg, CPP 39±0.8 mm Hg, end-tidal carbon dioxide 40±0.1 mm Hg, and pulse oximetry 94±0.4%. There was no statistical difference in mean vital signs between CA and sham animals in the post-CA period.
Citrate Synthase

There were no differences in CS activity between regions in sham animals (Figure 4). Further analysis displayed no differences in CS activity post CA in the cortex (16.8 [14.3 to 17.9] μmol/mL per minute) compared with sham (14.5 [12.8 to 15.6] μmol/mL per minute; P = 0.15). However, post CA, the mitochondrial content as measured by CS activity in the hippocampus decreased significantly, by nearly 30%, compared with sham (13.9 [10.7 to 15.2] versus 16.9 [15.0 to 18.3] μmol/mL per minute; P < 0.04). The results of all measures of respiratory capacity within the mitochondrial ETS were also normalized to CS activity for each animal (Table 1).

Cerebral Mitochondrial Respiration After Hypoxic VF Pediatric CA

Complex I–driven (OXPHOSCI) respiration and complex II–driven respiration (ETSCI) were both significantly decreased after CA, in the cortex (OXPHOSCI: P < 0.01, ETSCI: P < 0.05) and hippocampus (OXPHOSCI: P < 0.03, ETSCI: P < 0.01) when compared per milligram of tissue (Table 1). Reductions in both complex I– and complex II–driven respiration compared with sham animals remained significant when the data were normalized for CS (OXPHOSCI: cortex P < 0.04, hippocampus P < 0.01; ETSCI: cortex P < 0.05, hippocampus P < 0.01) (Table 1) and FCR (OXPHOSCI: cortex P < 0.01, hippocampus P < 0.05; ETSCI: cortex P < 0.05, hippocampus P < 0.01) (Figure 5A). Consequently, maximal coupled, oxidative phosphorylating respiration (OXPHOSCI+ETSCI) stimulated by both complex I and complex II substrates, was significantly decreased in the cortex (P < 0.02) and hippocampus (P < 0.02) per milligram of tissue and remained significant when normalized to CS activity (cortex P < 0.01, hippocampus P < 0.01) and FCR (cortex P < 0.05, hippocampus P < 0.02) (Figure 5B).

Maximal uncoupled, nonphosphorylating respiration (ETSCI+ETSCII), initiated by the protonophore carbonyl cyanide p-(trifluoromethoxy)phenylhydrazone after ATP synthase inhibition with oligomycin, was significantly reduced in the hippocampus (P < 0.01) per all measures: milligram of tissue and normalized to CS, while cortical ETSCI+ETSCII post CA was significantly reduced only per CS (P < 0.03) (Table 1). Complex IV respiration was significantly decreased in the hippocampus per milligram of tissue and normalized to CS activity (P < 0.02) but did not display a significant difference in either the cortex or the hippocampus post CA via FCR. Finally, LEAK (LEAKCI+ETSCI), or state 4 respiration, did not display a significant alteration post CA compared with sham in either region, except when normalized to FCR in the cortex and hippocampus, where there was a significant increase (P < 0.03) (Figure 5B).

Respiratory ratios evaluating phosphorylation coupling efficiency were calculated for sham and CA animals (Table 2). The respiratory control ratio for OXPHOSCI+ETSCI (OXPHOSCI+ETSCI/LEAK) was calculated to determine the efficiency of coupled phosphorylating respiration, a global measure of overall mitochondrial function, and was significantly decreased in the cortex (P < 0.05) and the hippocampus (P < 0.05) in CA compared with sham.

Discussion

These data establish that pediatric animals with ROSC after severe hypoxia, followed by VF and excellent CPR for 10 minutes, survive; however, they have diffuse cerebral mitochondrial bioenergetics dysfunction 4 hours after ROSC compared with sham controls. The pattern of mitochondrial bioenergetics dysfunction differs from previously described findings in mature animals, as we observe significant decreases in both complex I– and complex II–driven respiration, suggesting an important limitation in the immature brain’s ability to maintain or even accelerate oxidative phosphorylation in the first few hours after CA and challenges
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The notion that these animals should expect good neurologic recovery. This persistent cerebral mitochondrial bioenergetic dysfunction could play an important role in the development of secondary neurologic injury and reduce the likelihood of long-term recovery in the at-risk immature brain after survival from prolonged CA.

Our biochemical analysis of complex interdependent pathways of electron flow through the ETS revealed an overall injury pattern with a significant decrease in both complex I– and complex II–driven respiration in the brain 4 hours after CA despite maintenance of adequate CPP and cerebral perfusion pressure for successful clinical ROSC. Our method-
ology extends conventional bioenergetic protocols that use only complex I NADH–linked substrates (pyruvate+malate+glutamate) with the addition of the complex II substrate, succinate (pyruvate+malate+glutamate+succinate), to maximize oxidative phosphorylation capacity (ATP production) by simulating the in situ tricarboxylic acid cycle and maintaining key substrates for convergent flow of electrons through complexes I and II. Consequently, the reductions in convergent pathways of complexes I and II translated into significant reductions in oxidative phosphorylation post CA, likely placing the immature brain at significant risk due to a reduced ability to increase its ATP production and maintain bioenergetic output. Rosenthal and colleagues reported, in another large animal model of CA with dogs, that mitochon-
drial respiratory inhibition recovered to sham levels by 2 hours after a 10-minute VF CA. Our data suggest a more prolonged period of mitochondrial respiratory inhibition; however, this may be due to model differences where animals underwent a prolonged period of hypoxia before VF CA to simulate pediatric CA. Other confounders complicating the comparison with Rosenthal and colleagues’ work is the developmental age of the animal, as well as the techniques used to obtain mitochondrial respiration.

Complex I–driven respiratory dysfunction is a persistent characteristic of early mitochondrial dysfunction within 6 to 8 hours after traumatic and ischemic injury in the brain and in other organ systems such as the heart. Interestingly, our data show that both complex I– and complex II–driven respiration decreased in the immature brain after CA and resuscitation. This differs from measurements in mature rat brains exposed to forebrain ischemia and 6 hours of reper-
fusion, which did not show significant inhibition of complex II–driven respiration with succinate. It should be noted that this model may be distinctly different from the ischemia–

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<th>Table 1. Respiration of Brain Tissue Homogenates From Sham and Cardiac Arrest Cohorts</th>
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Mitochondrial respiration of brain tissue homogenates from the cortex and hippocampus (Hippo) side in 2 cohorts: sham and 4 hours post CA. Respiration is expressed per mg of tissue (pmols O2/s×mg), and per citrate synthase activity (pmols O2/s×CS). Presented as median (25th to 75th percentile range). CA indicates cardiac arrest; CS, citrate synthase; ETS, electron transport system; OXPHOSCI, oxidative phosphorylation capacity of complex I.

*P<0.05 significantly different from corresponding sham region. Percent change from sham indicated in parentheses. For de

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Figure 5. Mitochondrial FCR normalized by ETS capacity (ETS\textsubscript{CI+CII}) measured 4 hours post CA. A, CA resulted in a significant decrease in complex I–driven respiration (OXPHOS\textsubscript{CI}) in both regions: cortex and hippocampus. Complex II–driven respiration (ETS\textsubscript{CI}) in CA tissue is significantly decreased bilaterally, compared with corresponding sham tissue. B, Maximal coupled, phosphorylating respiration (OXPHOS\textsubscript{CI+CII}), stimulated by both complex I and complex II substrates, was significantly decreased post CA in both regions compared with shams. State 4\textsubscript{o} (LEAK\textsubscript{CI+CII}) displayed a significant increase 4 hours post CA compared with shams in the cortex and the hippocampus. For definitions of respiratory states and substrates, please see Figure 1. *P<0.05. Boxplots: Horizontal lines represent the median FCR, with the boxes representing the 25th and 75th percentiles, the whiskers representing the 5th and 95th percentiles, and the dots representing the outliers. CA indicates cardiac arrest; ETS, electron transport system; FCR, flux control ratios; OXPHOS\textsubscript{CI+CII}, oxidative phosphorylation capacity of complexes I and II.
reperfusion injury that occurs after CA. Despite this difference, several studies suggest that complex II–driven respiration is critical to mitochondrial respiration and salvaging ATP generation in the first several hours of reperfusion after cerebral ischemia.39,40 Future experiments will focus on measurements of substrate quantification and enzyme activity for succinate-dependent respiration. Thus, we provide preliminary support for the hypothesis that complex II–targeted substrate resuscitation may present a plausible candidate target for therapeutic intervention given its central role in cerebral energy production after hypoxic–ischemic injury.

Concomitant reduction in both complex I and II respiration in this study reveals a pattern of injury that suggests specific opportunities for mitochondrial-targeted interventions for brain injury after CA, beyond substrate supplementation. Mitochondria are a major source of reactive oxygen species generation in mature animals after reperfusion of the brain after hypoxia,41 and complex I is an important source of cerebral reactive oxygen species in ischemia and reperfusion.42,43 Niatsetskaya et al reported that inhibition of complex I with pyridaben was beneficial and led to a significant reduction in oxidative injury and cerebral infarct volume after hypoxic ischemia in neonatal mice.33 It is possible that acute attenuation of complex I respiration after CA in immature animals may be beneficial despite reductions in ATP production because of the potential to limit reactive oxygen species generation by attenuating reverse electron transport during reperfusion.33 However, chronic reductions in complex I function have been implicated in exacerbating chronic inflammation and cellular destruction in disease states such as Parkinson’s disease, and may play a prominent role in persistent bioenergetic dysfunction in other chronic neurodegenerative processes, such as Alzheimer’s disease, and, potentially, chronic encephalopathy triggered by acute brain injury, such as CA.44,45 Future research should focus on temporal variations in bioenergetic demands and mechanistic alterations that will inform manipulation of the mitochondrial respiration in survivors after CA to limit ongoing neurologic injury.

In contrast to a previous study using mature rodent models of VF and CPR,46 we did not find significant global alterations in LEAKCI+CII, or state 4o, respiration early after ROSC (with the exception of results normalized to FCR in the hippocampus). In the context of respiratory control ratios, OXPHOSCI+CII (state 3) respiration/LEAKCI+CII (state 4o), the reductions in coupled respiration seem to be driven by significant reductions in OXPHOSCI+CII post CA rather than significant alterations in LEAKCI+CII, or state 4o. Therefore, we suspect that the global reductions in mitochondrial respiration, in both the cortex and the hippocampus, were not due to significant uncoupling and increased permeability of the inner membrane at this time point. Our data suggest that mitochondrial permeability transition pores do not seem to be open at 4 hours after CA and add to an important understanding of the cerebral bioenergetic time-course after CA. Mitochondrial permeability transition activation is likely an early and potentially reversible event in ischemia–reperfusion injury, leading to mitochondrial swelling and release of cytochrome c that may contribute to global reductions in mitochondrial respiration seen in our study at 4 hours post CA.47,48

Finally, decreased CS activity in the hippocampus may parallel significant oxidative injury in the at-risk selectively vulnerable hippocampus after CA.28 Importantly, our data reveal a substantial reduction of CS activity, by nearly 30%, within the hippocampus 4 hours after CA. This is presumably relevant because CS is generally assumed to be the rate-limiting enzyme of the tricarboxylic acid cycle and, possibly, a biomarker of mitochondrial content. Future studies will be performed to design and assess other markers of mitochondrial content, such as mitochondrial DNA, which may add to the assessment of surrogates for evolving mitochondrial content alterations after injury.

**Limitations**

Time-course likely plays a significant role in alterations of mitochondrial function after CA. Evaluation of mitochondrial function over a more comprehensive time interval post injury will further improve our understanding of bioenergetic alterations post CA. In an attempt to limit heterogeneity, we also limited our investigations only to female animals at this time; however, future investigations should involve male animals to determine

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**Table 2. Respiratory Ratio of Brain Tissue Homogenates From Sham and Cardiac Arrest Cohorts**

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<th>Respiratory Ratio</th>
<th>Sham (n=5)</th>
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<td></td>
<td>Cortex</td>
<td>Hippo</td>
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<td>OXPHOSCI+CII/LEAKCI+CII</td>
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The respiratory control ratio OXPHOSCI+CII/LEAKCI+CII is calculated and presented as median (25th to 75th percentile range). OXPHOSCI+CII indicates the oxidative phosphorylation capacity of complexes I and II.

*P<0.05 significantly different from corresponding sham region. Percent change from sham is indicated in parentheses.
the applicability of these findings across sexes in the immature brain. Other limitations are that this is a descriptive study that has yet to be correlated with neurologic outcome measures. Additionally, we have reported that physiologic directed resuscitation improves neurologic outcomes; however, the Swine Cerebral Performance Category scales at 24 hours are a gross measure of both short- and long-term neurologic dysfunction. In fact, good neurologic outcome was defined by whether an animal could stand, but could still possess an unsteady gait, be able to eat, and exhibit a slow response to environmental stimuli. Based on this gross scale, there could still be significant neurologic injury. In addition, bioenergetic dysfunction has been shown to correlate with neurologic injury in other models; including human studies. Our current study generates several hypothesis-driven questions that future research will address, including the development of and correlation to short- and long-term neurologic outcomes. Finally, the use of an inhaled volatile anesthetic, isoflurane, may affect mitochondrial respiration. However, both the sham and experimental groups received the same concentrations and duration of isoflurane, therefore, the differences are likely to remain meaningful despite the anesthetic.

Conclusions

We have evaluated the mitochondrial bioenergetic response in the pediatric brain after CA and conclude that there are significant alterations in cerebral mitochondrial respiration and mitochondrial content in the pediatric brain 4 hours post ROSC in a large animal model despite excellent CPR, protocolized postresuscitative care, and improved traditional survival rates. Significant alterations in complex I– and complex II–driven mitochondrial respiration may offer opportunities for targeted therapeutic interventions in the immature brain to support post-CA bioenergetic demand by using substrate-directed resuscitation.

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Disclosures

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