Short paper

Transcriptome and metabolome after porcine hemodynamic-directed CPR compared with standard CPR and sham controls

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Abstract
Objective: The effect of cardiac arrest (CA) on cerebral transcriptomics and metabolomics is unknown. We previously demonstrated hemodynamic-directed CPR (HD-CPR) improves survival with favorable neurologic outcomes versus standard CPR (Std-CPR). We hypothesized HD-CPR would preserve the cerebral transcriptome and metabolome compared to Std-CPR.

Design: Randomized pre-clinical animal trial.
Setting: Large animal resuscitation laboratory at an academic children’s hospital.
Subjects: Four-week-old female piglets (8–11 kg).
Interventions: Pigs (1-month-old), three groups: 1) HD-CPR (compression depth to systolic BP 90 mmHg, vasopressors to coronary perfusion pressure 20 mmHg); 2) Std-CPR and 3) shams (no CPR). HD-CPR and Std-CPR underwent asphyxia, induced ventricular fibrillation, 10–20 min of CPR and post-resuscitation care. Primary outcomes at 24 h in cerebral cortex: 1) transcriptomic analysis (n = 4 per treatment arm, n = 8 sham) of 1727 genes using differential gene expression and 2) metabolomic analysis (n = 5 per group) of 27 metabolites using one-way ANOVA, post-hoc Tukey HSD.

Measurements and main results: 65 genes were differentially expressed between HD-CPR and Std-CPR and 72 genes between Std-CPR and sham, but only five differed between HD-CPR and sham. Std-CPR increased the concentration of five AA compared to HD-CPR and sham, including the branched chain amino acids (BCAA), but zero metabolites differed between HD-CPR and sham.

Conclusions: In cerebral cortex 24 h post CA, Std-CPR resulted in a different transcriptome and metabolome compared with either HD-CPR or sham. HD-CPR preserves the transcriptome and metabolome, and is neuroprotective. Global molecular analyses may be a novel method to assess efficacy of clinical interventions and identify therapeutic targets.

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Keywords: Cardiac arrest, Cardiopulmonary resuscitation, Metabolomics, Transcriptome, RNA sequence

Introduction:

Pediatric in-hospital cardiac arrests (CA) affect thousands of children each year, and permanent brain injury is common amongst those that survive to hospital discharge.1,2 Very little is known about alterations in RNA or metabolite profiles in cerebral cortex after CA. Describing the effect of distinct cardiopulmonary resuscitation (CPR) strategies on RNA and metabolite profiles may serve as an additional way of evaluating CPR efficacy and uncover novel pathways for future study.

Our group has previously shown that in a pediatric porcine model of asphyxia-associated CA, a hemodynamic-directed CPR strategy (HD-CPR) resulted in improved cerebral mitochondrial bioenergetics...
and greater 24-hour survival with favorable neurologic outcome compared to a guideline-based CPR strategy (Std-CPR). Among survivors, we have also previously shown improved intra-arrest hemodynamics and neurologic scoring 24 h post ROSC after HD-CPR compared to Std-CPR. To understand the effect of HD-CPR and Std-CPR on transcriptomics and metabolomics in the immature brain, we performed a quantitative RNA and metabolite analysis of cerebral cortex 24 hours after return of spontaneous circulation (ROSC) following asphyxia-associated ventricular fibrillation in a pediatric porcine model of CA with these two distinct CPR strategies. We hypothesized that among survivors of CA, HD-CPR would better preserve the cerebral cortical transcriptome and metabolome than Std-CPR when compared to sham animals that did not undergo CPR.

**Materials and methods**

**Experimental protocol**

**Cohort description**

These experiments were performed as a component of a previously reported pre-clinical trial in which animals were randomized to Std-CPR versus HD-CPR and complied with ARRIVE guidelines.

In that study, 7 of 10 animals treated with HD-CPR and 5 of 12 animals treated with Std-CPR survived to 24 hours. Available specimens were used from these survivors without knowledge of pre-arrest and intra-arrest characteristics or neurologic outcomes. Tissue samples available and sufficient for analysis were chosen, which yielded five samples in each treatment arm for metabolomic analysis and four samples in each treatment arm for transcriptomic analysis. Additional sham animals were available from concurrent studies with identical sham groups. Investigators performing the downstream analyses remained blinded to the characteristics and neurologic outcomes of the animals.

**Experimental protocol**

All animal procedures were approved by the Institutional Animal Care and Use Committee at the Children’s Hospital of Philadelphia (CHOP) and were conducted in strict accordance with the NIH Guide for the Care and Use of Laboratory Animals. Detailed description can be found in prior publications.

Briefly, one-month-old female swine, *Sus scrofa domesticus*, were anesthetized with isoflurane and mechanically ventilated and underwent seven minutes of asphyxia by endotracheal tube clamping during which fentanyl was administered. Ventricular fibrillation (VF) was induced to ensure a standard 1-minute period of CPR that would only be terminated if ROSC was restored for 24 hours. At 24 hours post-CA, animals were re-anesthetized, a craniectomy was performed, and cortical tissue was rapidly extracted and snap frozen in liquid nitrogen and then stored at −80 degrees Celsius. Animals were humanely euthanized while under general anesthesia. Frontoparietal cortex was used for downstream outcome metrics by members of the scientific team blinded to treatment status.

**RNA sequencing**

RNA sequencing was performed for 16 animals (HD-CPR n = 4, Std-CPR n = 4, sham n = 8). RNA was extracted using a Qiagen Rneasy (Qiagen, Hilden, Germany) automated RNA extraction robot and reverse transcribed using a high-capacity cDNA reverse transcription kit with RNase inhibitor (Cat. 4387406 Life Technologies, NY). Ribosome rRNA Removal Kit, Human/Mouse/Rat probe (illumina, San Diego, CA, USA) was used for rRNA depletion, followed by library preparation using NEBNext Ultra RNA Library Prep Kit for Illumina (NEB, Ipswich, MA, USA). Sequencing was performed using an Illumina HiSeq instrument for sequencing in 2 × 150 paired end configuration.

Reads were aligned to the SusScrofa11.1 reference genome using STAR.

We used featureCounts from the subread package to quantify transcripts at the gene level. DESeq2 v1.26.0 and R package v3.6.3 were used for downstream analyses. A false discovery rate (FDR) of 0.05 and 2-fold change was used to identify differentially expressed genes.

**Metabolite analysis**

Metabolomic analysis was performed for 15 animals (HD-CPR n = 5, Std-CPR n = 5, sham n = 5). Analysis prioritized pathways that feed into the TCA cycle and may impact cerebral cortex bioenergetics, as oxidative phosphorylation dysfunction after CA in the immature brain may contribute to post-ROSC neurologic injury. Amino acid concentrations were determined with an Agilent 1260 Infinity HPLC system, utilizing pre-column derivatization with O-phthalaldehyde. Lactate levels were determined enzymatically. Organic acid concentrations were determined by isotope dilution approach using GC-MS system. Briefly, an aliquot of the brain sample (100 ul) was spiked with a mixture of 13C-labeled organic acids of known concentrations. Then, GC-MS measurement of 13C isotopic abundance in each sample was performed and concentrations of organic acids in the sample were calculated. Protein level was determined by Coomassie blue (ThermoFisher Scientific).

Processing and analysis of metabolite data was done using the MetaboAnalystR (PMID: 29955821) package in R. Metabolite data was normalized using median row-normalization, log-norm transformation, and Pareto scaling, which yielded a probability distribution closest to Gaussian. Individual metabolite effect was determined by one-way ANOVA corrected by FDR with post-hoc Tukey honestly significant difference testing. Treatment arms were compared using Pearson correlation of sample medians to measure similarity between groups.

**Porcine cerebral performance category**

Porcine Cerebral Performance Category (CPC) was independently determined 24 h after ROSC by two blinded, trained investigators. The scoring system is as follows: 1 is normal (no difficulty standing, walking, etc.); 2 is mild disability (can stand, but unsteady, slow to respond, drinking, not eating); 3 is severe disability (awake but not
responding, cannot stand, walk, eat, drink); 4 is comatose; and 5 is dead.17,18

Data analysis
A priori planned analyses were to compare metabolome and transcriptome data among the 2 treatment groups (HD-CPR versus Std-CPR) and shams to address potential benefits of HD-CPR on outcomes and the potential mechanism of the outcome benefits in translational models before extending to clinical trials.

Results

RNA sequencing analysis
RNA expression in cerebral cortex at 24 h post-CA demonstrated little separation between HD-CPR and sham treatment arms (Fig. 2a). Of 1727 genes analyzed, only five were differentially expressed between HD-CPR and sham animals, while 65 genes differed between the Std-CPR and sham groups and 72 between Std-CPR and HD-CPR (Fig. 2b). Std-CPR increased expression of multiple genes related to immune function when compared to either of the other two groups, including adhesion molecules (ICAM1), chemokines and cytokines (CCL2, PLEK, TNFSF18, CCL3L1), genes critical to neutrophil function (CSF2RA, NCF2), regulatory molecules important in preventing autoimmunity (CD274) and transcription factors integral to innate immunity (ELF4) (Fig. 2c, 2d). HD-CPR increased expression of ASPA, which hydrolyzes N-acetylaspartate to maintain white matter compared to both Std-CPR and sham (Fig. 2e).

Metabolite analysis
Metabolite quantities in cerebral cortex at 24 h post-CA demonstrated little separation between HD-CPR and sham treatment arms (Fig. 3a). Of 27 metabolites studied, there were no differences between HD-CPR and sham animals. Std-CPR increased expression of 5 amino acids compared to HD-CPR and sham, including branch chain amino acids (BCAA) isoleucine (p = 0.005), leucine (p = 0.001), valine (p = 0.008) as well as glycine (p = 0.005) and phenylalanine (p = 0.003) (Fig. 3b, 3c). Overall metabolite expression in HD-CPR was more similar to sham (r = 0.12, p = 0.56) than either HD-CPR compared to Std-CPR (r = -0.51, p = 0.006) or between Std-CPR and sham (r = -0.47, p = 0.014). After normalization, positive Pearson correlation indicates similarity and negative correlation indicates differences between groups.

Porcine cerebral performance category
As previously described, at 24 h post ROSC all HD-CPR survivors with metabolome or transcriptome data attained CPC 1 (neurologically normal), whereas 4/5 Std-CPR survivors had CPC 3 (severe disability) and 1/5 had CPC 1.7

Discussion
Our group has previously shown that in a pediatric asphyxia-associated cardiac arrest swine model, HD-CPR leads to improved cerebral mitochondrial respiration and higher rates of 24 h survival with favorable neurologic outcome compared to Std-CPR.3,4 Among survivors, we have also previously demonstrated improved intra-arrest hemodynamics and neurologic scoring 24 h post ROSC after HD-CPR compared to Std-CPR.7 We hypothesized that HD-CPR animals better preserve the cerebral transcriptome and metabolome than Std-CPR in comparison to sham animals that did not undergo CPR. We now show that 24 hours after CA, cerebral cortex from animals that received HD-CPR had a similar transcriptome and metabolome compared to sham animals that did not undergo CA, and both groups were distinct from animals that received Std-CPR. These results, in concert with already established clinical and mitochondrial benefits, establish the benefit of HD-CPR on global molecular changes in the developing brain at 24 hours after CA. Our findings are the first to document alterations in cerebral transcriptomics and metabolomics after distinct cardiopulmonary resuscitation strategies in a highly relevant, pediatric large animal model.

There is no published data on the effect of CA on the transcriptome. Std-CPR results in differential RNA expression of numerous transcripts, many of which are pro-inflammatory compared to HD-CPR and shams. Neuroinflammation mediated by microglial activation plays a pivotal role in post CA brain injury, and immunomodulation has successively been neuroprotective in rodent studies.19,20

Fig. 1 – Schematic of Experimental Protocol. Definitions of abbreviations: CPR cardiopulmonary resuscitation; ETT = endotracheal tube; VF = ventricular fibrillation; Std = Standard, CPR = cardiopulmonary resuscitation; HD = hemodynamic-directed; SBP = systolic blood pressure; AP = anterior–posterior; CoPP = coronary perfusion pressure; ROSC = return of spontaneous circulation.
Trials of immunomodulation in large animals are needed to identify the link between neuroinflammation and poorer neurologic outcomes after CA treated with Std-CPR. We describe BCAA increase after Std-CPR compared to both HD-CPR and sham animals. BCAA are important for protein and neurotransmitter synthesis, though excess is neurotoxic. In adult rats undergoing CA, numerous cerebral amino acids, including BCAAs, are elevated after 30 minutes of reperfusion. Following Std-CPR, BCAA elevation may be a direct mediator of neurologic injury or a neuroprotective response to severe injury. Our findings warrant further mechanistic study of cerebral BCAA after CA associated brain injury. This model is limited in that the RNA and metabolite profiles are only available in survivors at 24 hours after CA. The transcriptome and metabolome may have an evolving time-course, which should be investigated. This study does not evaluate proteomics, which would bridge the gap between the transcriptome and metabolome. Additionally, due to the small number of genes studied, it was not possible to conduct a gene-ontology enrichment analysis.

In the cerebral cortex 24 h after CA, Std-CPR resulted in a different transcriptome and metabolome compared with both HD-CPR and shams. Notably, HD-CPR and sham animals were not substantially different from each other, suggesting that HD-CPR better preserved the native transcriptome and metabolome and appears to be a potent neuroprotective intervention. The effect of CA on immune function and the role of BCAA following CA are promising future directions. Global molecular analyses may be a novel method to assess efficacy of clinical interventions and identify potential therapeutic targets.

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