Regional lung metabolic profile in a piglet model of cardiopulmonary bypass with circulatory arrest

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Abstract

Introduction Acute lung injury is common following cardiopulmonary bypass and deep hypothermic circulatory arrest for congenital heart surgery with the most severe injury in the dorsocaudal lung. Metabolomics offers promise in deducing mechanisms of disease states, providing risk stratification, and understanding therapeutic responses in regards to CPB/DHCA related organ injury.

Objectives Using an infant porcine model, we sought to determine the individual and additive effects of CPB/DHCA and lung region on the metabolic fingerprint, metabolic pathways, and individual metabolites in lung tissue.

Methods Twenty-seven infant piglets were divided into two groups: mechanical ventilation + CPB/DHCA (n = 20) and mechanical ventilation only (n = 7). Lung tissue was obtained from dorsocaudal and ventral regions. Targeted analysis of 235 metabolites was performed using HPLC/MS–MS. Data was analyzed using Principal Component Analysis (PCA), Partial Least Square Discriminant Analysis (PLS-DA), ANOVA, and pathway analysis.

Results Profound metabolic differences were found in dorsocaudal compared to ventral lung zones by PCA and PLS-DA (R2 = 0.7; Q2 = 0.59; p < 0.0005). While overshadowed by the regional differences, some differences by exposure to CPB/DHCA were seen as well. Seventy-four metabolites differed among groups and pathway analysis revealed 20 differential metabolic pathways.

Conclusion Our results demonstrate significant metabolic disturbances between dorsocaudal and ventral lung regions during supine mechanical ventilation with or without CPB/DHCA. CPB/DHCA also leads to metabolic differences and may have additive effects to the regional disturbances. Most pathways driving this pathology are involved in energy metabolism and the metabolism of amino acids, carbohydrates, and reduction–oxidation pathways.

Keywords Acute lung injury · Cardiopulmonary bypass · Kynurenine metabolism · Congenital heart disease surgery · Metabolomics · Pathway analysis

1 Introduction

Approximately 1% of children born in the United States and Europe are diagnosed with congenital heart disease (Benjamin et al., 2018). Of those, 25% have critical congenital heart disease that requires surgery or catheter intervention within the first year of life (Benjamin et al., 2018). Unfortunately, the morbidity rate for these surgeries remains high with frequent complications leading to prolonged hospital length of stay (Pasquali et al., 2014). For many of these surgeries, cardiopulmonary bypass (CPB) is used to maintain perfusion and deep hypothermic circulatory arrest (DHCA) or selective cerebral perfusion may be used if a bloodless field is required. However, CPB and DHCA directly
contribute to postoperative inflammation and multiorgan injury (Kozik & Tweddell, 2006).

Acute lung injury (ALI) is among the lesser understood complications of CPB/DHCA but is an important determinant of morbidity and mortality (Apostolakis et al., 2010). Post-CPB ALI occurs in over 35% of cases (Liu et al., 2012; Shi et al., 2014) and can lead to prolonged mechanical ventilation and hospital stays (Shi et al., 2008). In severe cases, the mortality rate is approaches 33% (Khemani et al., 2019). The lungs, particularly those of neonates and infants (Kozik & Tweddell, 2006), are uniquely vulnerable to injury during CPB. The pulmonary vasculature has substantially diminished circulation with only bronchial artery supply during CPB, ventilation is often stopped or decreased during CPB, the lungs filter the venous circulation thus receiving all inflammatory products circulating from CPB, lung capillaries are small and trap aggregates of activating substances and cells in the lung, and large pools of neutrophils reside in the lungs that are immediately present to contribute to injury (Apostolakis et al., 2010; Kozik & Tweddell, 2006). This constellation leads to increased pulmonary vascular resistance, reduced surfactant activity, atelectasis, and pulmonary edema (Apostolakis et al., 2010; Kozik & Tweddell, 2006; Stephens et al., 2013). Furthermore, supine positioning leads to regional differences in lung ventilation and perfusion, which results in differences in injury patterns in the dorsal regions compared to ventral regions (Altemeier et al., 1985; Carvalho et al., 1985; Freitas et al., 2016). This heterogeneous injury contributes to overall lung inflammation, hypoxemia, and intrapulmonary shunting (Tojo et al., 2015). Particularly, little is known on the early metabolic changes induced by mechanical ventilation or CPB/DHCA nor if these changes may be localized regionally.

Metabolomics is an emerging field that examines changes in patterns of metabolite production to deduce mechanisms of disease states, provide risk stratification, and understand therapeutic responses (Johnson et al., 2016). Metabolites are small molecules that are the end products of metabolism and reflect protein activity (Mussap et al., 2013). They drive the essential functions of the cell including signaling, energy production and storage, and reduction-oxygenation reactions (Mussap et al., 2013). Patterns of metabolite expression are related to activity of specific biochemical pathways (Bowler et al., 2017; Johnson et al., 2016). In the study of the lung metabolome, indirect samples of lung metabolic activity such as urine, blood, exhaled breath particulate tissues, and bronchoalveolar lavage fluid (BALF) are most often used (Bowler et al., 2017); however, the interpretation of metabolite levels in these samples is less clear than direct samples of lung tissue (Johnson et al., 2016). There are far fewer studies that use lung tissue for metabolomic profiling.

In this study, we used a porcine model of early infant CPB/DHCA and performed targeted metabolomic profiling on lung tissue to compare key metabolic changes induced by (1) CPB/DHCA versus mechanical ventilation only controls (MV) and (2) dorsocaudal versus ventral lung regions. We hypothesized that CPB/DHCA would induce broad metabolic changes in the lung compared to the MV group and that the dorsocaudal region of the lung would demonstrate unique metabolic disturbances compared to the ventral region of the lung. We also sought to identify key metabolic pathways and individual metabolites that drove differences between these groups.

2 Materials and methods

Please refer to the online supplement for full details.

2.1 CPB/DHCA piglet model

The animal protocol used was approved by the Institutional Animal Care and Use Committee of the University of Colorado (Protocol Number: 107715(02)1D) in accordance with the Guide for the Care and Use of Laboratory Animals and the ARRIVE guidelines. Twenty-seven infant female piglets (5–10 kg) were included in an animal protocol. Water was supplied ad libitum and the feeding (Teklad 8753, Envigo, Madison, WI, U.S.A.) schedule was based on an activity factor of 1.2, for slight growth/gain and maintenance, 100 g feed twice daily (200 g/day).

The animals were divided into two groups, CPB/DHCA \((n = 20)\) and MV controls \((n = 7)\). The surgical methods have previously been published (Davidson et al., 2019). Briefly, peripheral CPB was achieved under isoflurane anesthesia. During DHCA, the animals were cooled to 22 °C for 75 min. After rewarming and CPB separation, the piglets were provided ICU care for 4 h then euthanized. For the MV group, animals were invasively mechanically ventilated under anesthesia with the same ventilator parameters and medications for anesthesia and euthanasia for 7 h without undergoing CPB/DHCA. Groups were then subdivided based on location of lung tissue biopsy to yield 4 sets of samples: ventral CPB/DHCA, dorsocaudal CPB/DHCA, ventral MV, and dorsocaudal MV. In our piglets in a supine position, the ventral lung region is considered gravity independent and the dorsal lung region is considered gravity dependent.

2.2 Histology

Right upper and lower lobe lung tissue was collected from each animal and fixed overnight in 10% formalin, paraffin-embedded, and sectioned at 4 μm. Serial sections were stained with hematoxylin–eosin (H&E) and evaluated for severity of lung injury by a blinded pathologist per the ATS recommendations (Matute-Bello et al., 2011).
2.3 Wet/dry ratio

Separate tissue samples (2×2 cm) from the right upper and lower portions of the lungs were weighed and left to dry for 5–7 days at 65 °C until constant weight was achieved to measure wet/dry ratios.

2.4 RNA preparation, RT, and real-time PCR

Total RNA was isolated from lung tissue (right upper and lower lobe tissue samples snap frozen in liquid N₂) using the RNeasy Mini Kit (Qiagen, Santa Clarita, CA) as described in the manufacturer’s protocol. RNA concentrations were quantified at 260 nm, and the purity and integrity were determined using a NanoDrop. RT and real-time PCR assays were performed to quantify steady-state mRNA levels of IL-6. cDNA was synthesized from 0.5 µg of total RNA. Predeveloped TaqMan primers and probes (Applied Biosystems, Foster City, CA) were used for detection. Reporter dye emission was detected by an automated sequence detector combined with ABI Prism 7300 Real Time PCR System (Applied Biosystems, Foster City, CA). Real-time PCR quantification was performed with TaqMan b-actin controls and relative mRNA expression calculated using the $2^{-\Delta\Delta CT}$ method.

2.5 Tissue harvest and sample preparation

Collected lung tissue samples were extracted according to a published protocol (Davidson et al., 2018). Briefly, frozen samples were ground with tissue grinder in 500 µL ice-cold methanol, sonicated for 10 min and centrifuged at 14,000 g for 10 min at 4 °C. The precipitant was reextracted with additional 500 µL ice-cold methanol, the supernatants were combined, centrifuged, and dried in a SpeedVac concentrator (Savant, ThermoFisher, Waltham, MA). Samples were reconstituted with 60 µL water/methanol (80:20, volume/volume).

2.6 Mass spectrometry

Multiple reaction monitoring (MRM) of 184 metabolites using a positive/negative ion-switching high-performance liquid chromatography–tandem mass spectrometry (5500 QTRAP HPLC-MS/MS (Yuan et al., 2012)) was used for analysis. Yuan et al. have provided reproducibility and extraction recovery testing in FFPE-embedded normal lung and kidney tissue as well as in kidneys from acute myeloid leukemia patients and lung tissues from patients with lymphangioleiomyomatosis lung disease. Our samples were immediately frozen and processed quickly using the published protocol thus no negative impact regarding metabolites stability or recovery should be expected.

2.7 Spectra analysis

MultiQuant (v2.1.1., Sciex, Foster City, CA) software was used for data processing of 235 unique metabolites in plasma. For between-sample normalization, the intensity values for each sample were summed, and the median value of the sums across all samples were determined. Tune and quality control samples were evenly distributed during the batches. The intensity values of each sample were then scaled such that the sum of the scaled intensities equaled the median value of all samples. Normalized intensity values were then log₂ transformed to reduce the influence of extreme values and to meet the homogeneity of variance assumption.

2.8 Statistics

Relative peak intensities were analyzed using MetaboAnalyst 4.0 (http://www.metaboanalyst.ca), which is run on the statistical package R (v3.4.3) (Chong et al., 2019). Where applicable, metabolites are named and grouped as listed on the Human Metabolome Database (HMDB) (http://www.hmdb.ca) (Wishart et al., 2018). Principle component analysis (PCA) was performed first to see how data clustered in an unsupervised analysis. Partial least-squares discriminant analysis (PLS-DA) was then used to identify the features that discriminated between the preassigned groups. Variable importance of projection (VIP) was used to identify the most influential variables to the discrimination in PLS-DA. A one-way analysis of variance (ANOVA) was performed to determine which metabolites differed significantly among groups. P-value was adjusted for false discovery rate (FDR) of 0.05. Fisher’s Least Significant Difference (LSD) post-hoc test was performed to determine if the relative peak intensity was significantly elevated by region (dorsocaudal region versus ventral region), CPB/DHCA versus MV, or additive. Metabolites with an additive effect demonstrated effects of both CPB/DHCA and lung region in at least one group. Pathway analysis was performed using GlobalANCOVA (Hummel et al., 2007) for Metabolite Set Enrichment Analysis (MSEA) and Relative-Betweeness centrality for topographic analysis. SMPDB (https://www.smpdb.ca) (Frolkis et al., 2010; Jewison et al., 2014) was used for the pathway library. Descriptions from this database were used to categorize pathways based on common features. Cut-off for statistical significance for pathway analysis were a Holm-Bonferonni adjusted p-value < 0.000556 and an impact factor ≥ 0.1. All artwork was created with MetaboAnalyst 4.0 with labels and size edited with Apple Preview version 11.0.
3 Results

3.1 Samples and initial metabolite data preparation

There were 20 animals who underwent CPB/DHCA and 7 animals who underwent MV. After the samples from the ventral and dorsocaudal regions were obtained, the final samples included the following: (1) in the MV group, 6 samples from the ventral region and 7 samples from the dorsocaudal region and (2) in the CPB/DHCA group, 18 samples from the ventral region and 20 samples from the dorsocaudal region.

On gross inspection, the lungs from all animals in both the CPB/DHCA and MV control groups showed significant discoloration in the dorsocaudal region (see Supplemental Fig. S1). Histologic examination of the dorsocaudal lung compared to the ventral lung demonstrated a marked difference in atelectasis as well as increased septal thickening and the scattered presence of intra-alveolar inflammatory cells in both the CPB/DHCA animals and MV controls. (Supplemental Fig. S2). The dorsocaudal region demonstrated a higher wet-to-dry ratio compared to the ventral lung within both the MV (median 4.9 [IQR 4.7, 5] versus 4.2 [IQR 3.8, 4.4]; \( p = 0.02 \)) and CPB/DHCA group (median 4.4 [IQR 3.9, 4.8] versus 3.5 [IQR 3.2, 3.8]; \( p < 0.0001 \)). There was no significant difference in wet-to-dry ratio between the CPB/DHCA and MV groups. Collectively these findings suggest a different microenvironment in the dorsocaudal region and support the concept of regional metabolomic analysis. Additionally, IL-6 mRNA expression was significantly elevated in the dorsocaudal lung of CPB/DHCA animals compared to MV controls (17-fold increase; \( p = 0.01 \)) but not in the ventral lung, demonstrating the potential for synergistic effects between lung region and exposure to CPB/DHCA that should also be accounted for during metabolomic analysis.

Two hundred and thirty-five metabolites were initially targeted. After removing unstable metabolites or metabolites that were below the limits of quantification for the large majority of samples, 175 metabolites remained.

3.2 Unsupervised and supervised metabolomic fingerprinting

To begin the analysis, unsupervised PCA was performed to see how the data segregated independent of sample group identification. The analysis revealed clear differentiation between regions of the lung based on the metabolic fingerprint. As shown in Fig. 1A, the data differentiated along both PC1 (21.8%) and PC2 (15.6%) demonstrating distinct metabolic patterns between the ventral and dorsocaudal regions.
of the lung. The marked regional differences in metabolites in the lung made it difficult to determine differentiation based on exposure to CPB/DHCA. PLS-DA was therefore performed to assess if CPB/DHCA group differentiation could be further elicited and if so, what variables contributed most to this differentiation.

PLS-DA demonstrated discrimination between ventral and dorsocaudal regions clearly along the first two components as shown in Fig. 1B. The first two components also begin to show discrimination between CPB/DHCA and MV groups. Cross-validation showed good reproducibility \( R^2 = 0.7; Q^2 = 0.59; p < 0.0005 \). The discrimination is especially apparent in the dorsocaudal regions compared to the ventral regions.

Variable importance in projection (VIP) was then performed to identify which metabolites contributed most to the observed distribution between 4 groups. Fifty-three metabolites had a VIP score greater than 1 and thus contribute to the data clustering. See Supplemental Table S1 for a complete list of these metabolites. Twenty-five metabolites with a VIP score greater than 1.5 are shown in Fig. 2 along with a heat map of relative peak intensities in each group. To verify the statistical significance of the metabolites interpreted as contributing to discrimination by VIP and to quantitatively compare metabolites among groups, an ANOVA was performed. Using an FDR of <0.05, 74 significant metabolites were found. All but 1 metabolite with a VIP score greater than 1 were also shown to be significant between groups using ANOVA. To determine specific between-group differences, post-hoc testing was performed. Results are listed in Table 1, demonstrating metabolites elevated by lung region and by CPB/DHCA. Table 1 also lists metabolites that appeared to have an additive effect by both lung region and CPB/DHCA. Please refer to the supplemental Table S2 for complete detail. The following sections highlight significant metabolites and associated pathways by lung region and by CPB/DHCA. HMDB ID numbers for these significant metabolites are listed in supplemental Table S3.

### 3.3 Metabolic differences by lung region

As listed in Table 1, 61 metabolites differed by lung region with 38 metabolites elevated in the ventral region and 23 metabolites elevated in the dorsocaudal region. As an example of this pattern, Fig. 3A demonstrates the normalized relative intensities of glyceric acid, which was elevated in the dorsocaudal region of both MV and CPB/DHCA animals. Pathway analysis was performed which demonstrated 18 metabolic pathways that contributed to the metabolic differences between ventral and dorsocaudal regions. Pathwayst...
| Metabolites by region | Metabolites by CPB/DHCA | Metabolites with additive effects |
|-----------------------|-------------------------|----------------------------------|
| Elevated in ventral region | Elevated in CPB/DHCA | 2-ketobutyric acid |
| 1-methylnicotinamide | 2-aminobenzoic acid | 3-phosphoglyceric acid |
| 2-ketobutyric acid | 2-ketobutyric acid | 4-hydroxybenzoic acid |
| 3-methylphenylacetic acid | 3-phosphoglyceric acid | Allantoin |
| 3-methylthiopropionic acid | 4-Hydroxybenzoic acid | Cis-aconitic acid |
| 3-phosphoglyceric acid | 4-hydroxyphenylpyruvic acid | Citric acid |
| 5'-methylthioadenosine | Acetoacetic acid | Glycerophosphocholine |
| 6-phosphogluconic acid | Allantoin | Isocitric acid |
| Alpha-ketoisovaleric acid | Cis-aconitic acid | Kynurenic acid |
| Ascorbic acid | Citric acid | L-lactic acid |
| Betaine aldehyde | D-erythrose 4-phosphate | L-lysine |
| Citric acid | D-glucose | L-kynurenine |
| Fumaric acid | Fructose-6-phosphate | Malondialdehyde |
| Glutaric acid | Glycerophosphocholine | Phosphoserine |
| Glutathione | Indoleacrylic acid | Pyruvic acid |
| Hydroxyisocaproic acid | Isocitric acid | S-adenosylhomocysteine |
| Isocitric acid | L-alanine | Thiamine |
| Imidazoleacetic acid | L-lactic acid | Ureidosuccinic acid |
| L-asparagine | L-lysine | |
| L-aspartic acid | L-proline | |
| L-cysteine | Malondialdehyde | |
| L-lactic acid | Pyruvic acid | |
| L-malic acid | Quinolinic acid | |
| L-tryptophan | S-adenosylhomocysteine | |
| Maleic acid | Sedoheptulose 1,7-bisphosphate | |
| N-acetyl-D-glucosamine | Succinic acid | |
| N-acetyl-glutamate | Thiamine | |
| Nicotinic acid | Ureidosuccinic acid | |
| Phosphoenolpyruvic acid | Decreased in CPB/DHCA | |
| Phosphoserine | L-kynurenine | |
| Pyrophosphate | Phosphoserine | |
| Pyruvic acid | Uridine | |
| S-adenosylhomocysteine | | |
| S-adenosylmethionine | | |
| Shikimate 3-phosphate | | |
| Thiamine | | |
| Trigonelline | | |
| Elevated in dorsocaudal region | | |
| 1-methyladenosine | | |
| 1-methylguanosine | | |
are shown in Fig. 4A and the complete list of pathways are shown in Table 2. Please see supplemental Table S4 for more detailed statistical data. Affected pathways included those involved in the metabolism of amino acids, betaine, carbohydrates, energy, nitrogen, and vitamins. Glutathione metabolism and spermidine/spermine biosynthesis were also affected by lung region.

3.4 Metabolic differences by CPB/DHCA

As listed in Table 1, 29 metabolites were significantly higher in the lungs of CPB/DHCA animals. As an example of this pattern, Fig. 3B demonstrates the normalized relative intensities of acetoacetatic acid, which was elevated in both the ventral and dorsocaudal regions of CPB/DHCA animals. Pathway analysis demonstrated 2 pathways that contribute to the metabolic differences caused by CPB/DHCA alone. These included 1 pathway involved in amino acid metabolism (phenylalanine and tyrosine metabolism) and 1 pathway involved in energy metabolism (ketone body metabolism). Ketone body metabolism was unique to differences caused by CPB/DHCA while phenylalanine/tyrosine metabolism was affected by both lung region and CPB/DHCA.

Table 1 (continued)

| Metabolites by region | Metabolites by CPB/DHCA | Metabolites with additive effects |
|-----------------------|-------------------------|----------------------------------|
| 4-hydroxybenzoic acid |                         |                                  |
| Allantoin             |                         |                                  |
| Citraconic acid       |                         |                                  |
| Creatine              |                         |                                  |
| Creatinine            |                         |                                  |
| Glucose-6-phosphate   |                         |                                  |
| Glyceric acid         |                         |                                  |
| Glycerophosphocholine |                         |                                  |
| Hexose phosphatea     |                         |                                  |
| Kynurenic acid        |                         |                                  |
| L-arginine            |                         |                                  |
| L-carnitine           |                         |                                  |
| L-kynurenicine        |                         |                                  |
| L-lysine              |                         |                                  |
| L-tyrosine            |                         |                                  |
| Malondialdehyde       |                         |                                  |
| Methylcysteine        |                         |                                  |
| Oxidized glutathione  |                         |                                  |
| Pyroglutamic acid     |                         |                                  |
| Thymine               |                         |                                  |
| Ureidosuccinic acid   |                         |                                  |

Metabolites with significantly elevated relative peak intensities as determined by analysis of variance post-hoc testing. Metabolites are listed by region and by cardiopulmonary bypass/deep hypothermic circulatory arrest (CPB/DHCA). Metabolites elevated in the ventral or dorsocaudal regions are listed as such. Metabolites with additive effects are those that appeared to have an additive effect of both region and CPB/DHCA. aHexose-phosphate represents all hexose sugar phosphates combined, as there were too many isomers to distinguish using the extraction methods used.
3.5 Metabolites demonstrating additive effects from both lung region and CPB/DHCA

As listed in Table 1, 18 metabolites showed additive effects from both lung region and CPB/DHCA. As an example, Fig. 3C shows the relative normalized levels of citrate, demonstrating effects of both lung region and CPB/DHCA. These 18 metabolites are involved in a number of pathways including amino acid, carbohydrate, lipid, propanoate, pyrimidine, and vitamin metabolism as well as reduction–oxidation reactions and energy production.

4 Discussion

4.1 Key findings

Using a large animal model of CPB/DHCA, we performed targeted metabolomic profiling to directly investigate the influence of CPB/DHCA on the global and regional metabolic changes in porcine lung tissue. Our results demonstrate a dramatic difference in the metabolic fingerprint of the dorsocaudal lung region compared to the ventral lung region in both animals that underwent CPB/DHCA and control animals that underwent MV. To a lesser extent, lung tissue from the CPB/DHCA animals showed a unique metabolic fingerprint compared to MV animals. Key drivers of these metabolic changes were metabolites involved in essential cell function pathways including energy, carbohydrate, and nitrogen metabolism, amino acid synthesis and degradation, vitamin metabolism, and reduction–oxidation reactions. Due to the abundance of significant results that have important prognostic and therapeutic potential, a comprehensive review of all pathways is beyond the scope of this article. Specific pathways will be discussed based on their statistical significance, contribution to the understanding of injury mechanisms, and potential to be therapeutic targets.

4.2 Isolated regional differences in lung tissue metabolic fingerprint

The metabolic fingerprint of the dorsocaudal lung samples differed dramatically from the ventral lung samples in both control animals and the CPB/DHCA group, with the majority of pathways demonstrating additive effects by lung region and CPB/DHCA exposure (discussed in subsequent sections). Several metabolic pathways appear to be influenced primarily by lung region, with the most notable being glutathione (GSH) metabolism. Our findings suggest a difference in oxidative states between lung regions, with elevation of upstream reactants GSH and L-cysteine in the ventral regions and elevation of oxidized glutathione (GSSG) and pyroglutamic acid in the dorsocaudal regions. Other studies have shown that the balance among GSH, L-cysteine, and GSSG are altered in BALF and serum of humans with lung injury (Bernard et al., 1997; Kellner et al.,...
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2017), and our model adds to these findings by identifying the dorsocaudal lung as the potential source of GSSG. This finding has important translational implications as GSH metabolism is being investigated for its therapeutic potential in lung diseases. For example, repletion of both GSH with the precursor N-acetylcysteine (NAC) and L-cysteine with precursor pro-cysteine decreases the duration of ALI in human subjects (Bernard et al., 1997). In mice undergoing CPB/DHCA, increasing glutathione peroxidase activity reduced markers of multi-organ damage and ischemia/reperfusion injury in the lung (Steinbrenner et al., 2016). GSH supplementation has also been used to treat dyspnea in patients with COVID-19 (Horowitz et al., 2020). Our findings of regional oxidation of glutathione in the dorsocaudal lung suggest that increased oxidative stress in the posterior lung of supine patients could contribute to the increased lung injury seen in this area. Given the promising uses of GSH-related therapy for inflammation-induced lung injury, targeted GSH or other antioxidant therapy to the posterior lung could represent a novel mitigation strategy for post-CPB lung inflammation.

4.3 Isolated CPB/DHCA effects on lung metabolome

CPB/DHCA also influenced the lung metabolome, but to a lesser extent than lung region. The main pathway that showed significant effects from CPB/DHCA exposure but not from lung region was ketone body metabolism. Specifically, we found that CPB/DHCA significantly altered ketone body metabolism on pathway analysis and the ketone body acetoacetic acid was elevated in CPB/DHCA compared to MV. Although the liver is known to be the major source of ketone bodies following CPB/DHCA (Nomoto et al., 1992; Ozawa et al., 1983; Poggetti et al., 1992), the lungs themselves also have the potential to produce ketone bodies. The lungs use fatty acid beta oxidation during stress states, leading to ketone body production (Liu & Summer, 2019). Furthermore, unique isotypes of the rate limiting enzyme for ketone body production, HMG-CoA lyase, exist in the lung (Arnedo et al., 2019) but thus far there are limited studies on their role in lung injury. Our study is the first to suggest that the lungs may participate in ketone body synthesis following CPB/DHCA. As ketone body metabolism plays a protective role in stress states (Arnedo et al., 2019), and is being proposed as a potential therapy to prevent ARDS in patients with COVID-19 (Bradshaw et al., 2020), our findings should...
be further investigated to evaluate the prognostic and therapeutic applications of ketone body metabolism in CPB/DHCA patients.

While our study is the first to examine tissue-level metabolic changes after CPB, Maltesen et al. did examine the effects of CPB on metabolomic changes in BALF from a cohort of adults with COPD who were undergoing cardiac surgery (Maltesen et al., 2018). The authors showed similar metabolomic changes to those found in our study, namely elevated metabolites involved in glycolysis, urea cycle, protein catabolism, urea metabolism, purine metabolism, and nitric oxide synthesis. It is unclear how much lung region could have contributed to these findings as the sampling of BALF in this study was obtained only from the right middle lobe. Furthermore, Maltesen’s study used adult patients with the confounding variable of chronic lung disease. These limitations demonstrate the importance of using complementary, clinically relevant translational models to study the complex metabolic changes in cardiac surgery patients. Specifically, pediatric porcine models may more accurately reflect pediatric congenital heart surgery patients, since pigs have similar lung metabolome, anatomy, histology, and immune systems, and thus serve as good models of humans in physiological and mechanistic studies (Judge et al., 2014; Mair et al., 2014).

### 4.4 Additive effects of CPB/DHCA and lung region

Lung region and CPB/DHCA together also have substantial additive effects on specific metabolites and associated pathways. Glycolysis, the citric acid cycle (CAC), and the tryptophan-kynurenine pathway are all affected in an additive fashion and demonstrate the importance of considering lung region in the investigation of CPB/DHCA effects on the lung metabolome. For example, in the ventral region but not dorsocaudal region CPB/DHCA may have increased energy producing glycolytic and CAC flux based on elevation of 3-phosphoglyceric acid (a metabolite in the energy generating pathway of glycolysis) (Liu & Summer, 2019), and pyruvate (the end product of aerobic glycolysis which is

| Pathways affected by lung region | Pathway | Matched metabolites/total metabolites | P-value | Impact |
|---------------------------------|---------|--------------------------------------|---------|--------|
| Amino acid metabolism           | Arginine and proline metabolism | 18/48 | 1.18E-07 | 0.50259 |
| Aspartate metabolism            | 13/34 | 5.07E-07 | 0.89583 |
| Glutamate metabolism            | 16/45 | 8.86E-09 | 0.29996 |
| Glycine and serine metabolism   | 21/50 | 6.32E-09 | 0.15486 |
| Homocysteine degradation        | 5/7 | 1.63E-08 | 0.5 |
| Methionine metabolism           | 18/39 | 9.17E-09 | 0.34418 |
| Phenylalanine and tyrosine metabolism | 9/25 | 1.43E-07 | 0.46903 |
| Betaine metabolism              | 10/18 | 1.26E-05 | 0.70336 |
| Carbohydrate metabolism         | Gluconeogenesis | 13/30 | 1.26E-04 | 0.30686 |
| Glycolysis                      | 9/20 | 4.18E-04 | 0.44167 |
| Energy metabolism               | Citric acid cycle | 11/26 | 8.70E-06 | 0.3147 |
| Pyruvate metabolism             | 9/37 | 6.06E-05 | 0.19838 |
| Transfer of acetyl groups into mitochondria | 6/18 | 2.71E-05 | 0.85846 |
| Warburg effect                  | 19/49 | 3.18E-07 | 0.3801 |
| Nitrogen metabolism             | Urea cycle | 13/23 | 1.01E-06 | 0.64611 |
| Reduction-oxidation reactions   | Glutathione metabolism | 5/19 | 2.04E-10 | 0.29287 |
| Spermidine and spermine biosynthesis | 5/14 | 3.32E-09 | 0.51429 |
| Vitamin metabolism              | Nicotinate and nicotinamide metabolism | 8/32 | 3.77E-05 | 0.29083 |

| Pathways affected by CPB/DHCA | Pathway | Matched metabolites/total metabolites | P-value | Impact |
|--------------------------------|---------|--------------------------------------|---------|--------|
| Amino acid metabolism           | Phenylalanine and tyrosine metabolism | 9/25 | 6.82E-05 | 0.46903 |
| Energy metabolism               | Ketone body metabolism | 2/12 | 9.82E-06 | 0.34091 |

Significant pathways contributing to regional effects and cardiopulmonary bypass/deep hypothermic circulatory arrest (CPB/DHCA) are shown. Matched metabolites/Total metabolites reflects the number of metabolites that were sampled out of the total number of metabolites in given pathway. P-value was calculated by Metabolite Set Enrichment Analysis and Impact Factor was calculated by Topographic Analysis. *FDR False Discovery Rate. Adjusted P-value* P-value adjusted by the Holm-Bonferroni correction.
then either converted into lactate or enters the CAC (Liu & Summer, 2019). Lactate and the first three metabolites of the CAC (citric acid, cis-aconitic acid, and isocitrate) were also elevated in both the ventral region and in the CPB/DHCA group. The study by Maltesen et al., also showed alterations in glucose and energy metabolism with CPB (Maltesen et al., 2018), and our study demonstrating that these effects are focused regionally. A study by Eckle et al. further illustrates the importance of considering regional lung metabolome analysis in ALI. The authors showed that in murine MV-induced ALI, hypoxia inducible factors (HIF) were stabilized by alveolar distension, leading to succinate dehydrogenase inhibition, alteration of TCA flux, and stabilization of mitochondrial glycolytic capacity (Eckle et al., 2013). Gravity independent regions have comparatively more ventilation and alveolar distension with less atelectasis than gravity dependent regions (Carvalho et al., 1985; Magnusson et al., 1997; West, 2011). Thus, Eckle’s study shows a potential mechanism for the gravity dependent metabolic changes in a lung with a potentially injurious stimulus. There are therapeutic implications for this as well as HIFs being studied as therapeutic targets for a number of conditions including both ALI and pulmonary hypertension (Lee et al., 2019).

Metabolites of the tryptophan-kynurenine pathway also showed variance based on both CPB/DHCA and lung region as tryptophan was elevated in the ventral lung, while downstream metabolites kynurenine (KYN) and kynurenic acid (KYNA) were both elevated in the dorsocaudal lung and both altered CPB/DHCA animals. This finding demonstrates potential regional lung sources of KYN and KYNA, both of which have been demonstrated by our group to be elevated in serum from infants undergoing CPB (Davidson et al., 2018). Serum KYNA is associated with worse outcomes in infants following CPB/DHCA for heart surgery (Davidson et al., 2018). As KYN has potent vasodilatory effects on lung vasculature (Nagy et al., 2017; Wang et al., 2015) and the KYN arm of the tryptophan-kynurenine pathway plays important roles in oxidative injury (Lugo-Huitrón et al., 2011; Wang et al., 2015), multiple drugs are under development that target KYN pathway enzymes to modulate inflammation and oxidative injury for cancer and neurodegenerative diseases (Dounay et al., 2015). The availability of these drugs along with the data supplied by the current study that help localize sources of these metabolites are promising for the future of research investigating KYN pathway modulators for treatment of CPB-related lung injury.

4.5 Limitations

This study has several limitations. First, the small number of animals, particularly in the MV group, limited the power of the study; a larger sample size may have yielded more significant metabolites and pathways. This especially was likely to contribute to the low numbers of statistically significant pathways observed when comparing CPB/DHCA vs MV. Similarly, the study lacked a control group that was unexposed to MV. Therefore, we can only comment on regional differences under mechanical ventilation as opposed to the effects of mechanical ventilation compared to spontaneous breathing. Next, the injury was assessed at a single time point, early in the postoperative course. Systemic inflammation and pulmonary dysfunction following CPB/DHCA evolves from the time of CPB/DHCA initiation to 48 h post-surgery with peak inflammation typically between 3 h to 12 h following CPB/DHCA. Thus, studies exploring serial time points may more completely reflect an operative and post-operative course and help capture the most accurate metabolic alterations. Also, due to their similar lung metabolome, anatomy, histology, and immune systems, pigs are known to serve as good models of humans in physiological and mechanistic studies (Benahmed et al., 2014; Judge et al., 2014; Mair et al., 2014). They are not, however, perfect substitutes for humans in clinical trials. Another important limitation is our study used peripheral cannulation, while midline sternotomy with central cannulation is typically used in humans for CPB/DHCA. While this strategy allowed us to isolate the effects of CPB/DHCA from the mechanical effects of an open thorax, it is possible that our model could underrepresent the full extent of injury seen during surgery.

In terms of pathway analysis, pathways were grouped based on similar features to better describe patterns of pathway disturbances. This method, however, may lead to missed pathway categories, overlapping categories, or poorly fit pathways into categories. Lastly, many of the significant metabolites are involved in anapleurotic reactions, meaning they are used in several different metabolic pathways. This makes it difficult to draw mechanistic conclusions; thus, results from this study should be used to generate more targeted studies on individual metabolites, fully mapped metabolic pathways, and the activity of associated enzymes. Also, some metabolites may have different redox stability, such as GSH and GSSG, but processing and analysis of all samples was done uniformly and simultaneously. To best move forward from these limitations and elaborate on the current study, there are several exciting research methods to pursue: tissue metabolic pathways can be further explored with paired analysis of tissue and local arterial/venous blood sampling. Exploring structure–function relationships can accomplished with sub-regional, cellular, or sub-cellular analysis. Pathway modulation and flux studies could further describe mechanisms of metabolic pathway alterations. Taken together, these future studies could be used to design novel biomarkers and therapies for clinical application.
5 Overall conclusions

The current study was the first to directly analyze the metabolome of lung tissue following CPB/DHCA and compare the metabolome of ventral and dorsocaudal lung regions. The results identified a large number of significantly altered metabolites and metabolic pathways, demonstrating broad metabolic disturbances in gravity dependent lung regions during supine mechanical ventilation with or without CPB/DHCA. Exposure to CPB/DHCA also led to metabolic differences and often had additive effects to the regional disturbances. Most perturbed pathways included the metabolism of amino acids, carbohydrates, lipids, steroids, vitamins, and reduction–oxidation pathways.

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Author contributions SJC drafted the article. SJC, LK, JD designed and conceptualized the study. SJC, LK, BF, JD performed data analysis and interpretation. JK developed the metabolite analysis methods and acquired the data. JK and LK analyzed the samples. LK, JR, SO, and JD performed the animal surgery. JR, JJ, RI, SL, SO, and JD developed the model. SO and JD acquired funding. JD mentored the preparation of the manuscript. All authors critically revised the article and approved the article for submission.

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Data availability The metabolomics and metadata reported in this paper are available in the online data supplement.

Code availability Not applicable.

Declarations

Conflict of interest No authors declare any conflict of interest.

Ethical approval The animal protocol used was approved by the Institutional Animal Care and Use Committee of the University of Colorado (Protocol Number: 107715(02)1D) in accordance with the Guide for the Care and Use of Laboratory Animals and the ARRIVE guidelines.

Animal rights All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

Consent to participate Not applicable.

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