Molecular Signature of HFpEF
Systems Biology in a Cardiac-Centric Large Animal Model

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HIGHLIGHTS
- Early cardiac mitochondrial dysfunction is mediated by transcriptional down-regulation of the mitochondrial proteome.
- Comprehensive metabolic remodeling is conserved throughout HFpEF progression and includes increased amino acid and lipid species, indicative of impaired oxidative metabolism.
- Transcriptional and metabolic remodeling of skeletal muscle suggests cardiac signaling as a mediator of peripheral tissue maladaptation.
- Unbiased systems-level analysis provides new mechanisms underlying HFpEF development.

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In this study the authors used systems biology to define progressive changes in metabolism and transcription in a large animal model of heart failure with preserved ejection fraction (HFpEF). Transcriptomic analysis of cardiac tissue, 1-month post-banding, revealed loss of electron transport chain components, and this was supported by changes in metabolism and mitochondrial function, altogether signifying alterations in oxidative metabolism. Established HFpEF, 4 months post-banding, resulted in changes in intermediary metabolism with normalized mitochondrial function. Mitochondrial dysfunction and energetic deficiencies were noted in skeletal muscle at early and late phases of disease, suggesting cardiac-derived signaling contributes to peripheral tissue maladaptation in HFpEF. Collectively, these results provide insights into the cellular biology underlying HFpEF progression. (J Am Coll Cardiol Basic Trans Science 2021;6:650–672) © 2021 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
FIGURE 1 1-Month Aortic Banding Reveals a Cardiac RNAseq Signature of Transcriptional and Post-Transcriptional Regulation, Mitochondrial Dysfunction, and Inflammation

A. Heart: 1-mo. Banded vs 1-mo. Sham

B. 1-mo. Banded vs 1-mo. Sham

C. Biological Process

D. Molecular Function

Epigenetics
DNA Repair
Metabolism/Mitochondria

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in HFpEF due to the lack of a reliable model. Here, we interrogated the early cardiac and skeletal muscle transcriptional and metabolic signatures in a feline banding model to aid our understanding of HFpEF pathogenesis and identify primary mechanisms driving disease progression and potential therapeutic targets.

**METHODS**

**ANIMAL PROCEDURES.** All animal procedures were approved by the Temple University School of Medicine Institutional Animal Care and Use Committee. We used a total of 18 male shorthair kittens (aged 2 months [1.3 kg]) (Liberty Research Inc), that underwent either aortic constriction (n = 10, with customized preshaped bands or a sham procedure (n = 8). Before surgery, baseline phenotyping was performed, and animals were randomly assigned to their respective groups. Animals were sedated with ketamine (25 mg/kg) and acepromazine (0.1 mg/kg), intubated, and mechanically ventilated (Narkomed 2b, Dräger). Surgical (plane) anesthesia was maintained throughout the procedure with 1%-2% isoflurane mixed with 100% oxygen. Under sterile conditions, a 2- to 4-cm skin incision was made over the 3rd and 4th intercostal space and extended through the intercostal muscles. The pericardium was opened, and the aorta was dissected from the pulmonary artery. The preshaped band was placed around the ascending aorta and gently tied down without causing significant constriction of the aorta. Animals received heat support during anesthesia and recovery. In contrast to immediate pressure overload, this approach results in a slow progressive pressure overload during growth, as the cats grow into the aortic band. Cats were group housed in spacious rooms that were enriched with scratching posts, climbing towers, and a hammock where they could freely move and were provided with food and water ad libitum. No animals were excluded from study.

**TRANSTHORACIC ECHOCARDIOGRAPHY.** Echo was performed with a Vivid q Vet Premium BT12 (GE Healthcare) using a 12S-2S sector probe at baseline and 1 month post-surgery, and 4 months post-surgery, as previously described (16,17). Animals were sedated with an intramuscular injection of alfaxalone (1 mg/kg), butorphanol (0.5 mg/kg), and midazolam (0.5 mg/kg). Alfaxalone is a neuroactive steroid molecule with properties of a general anesthetic. Alfaxalone does not cause an increase in heart rate, which is occasionally observed with ketamine. Ivabradine (0.3 mg/kg) was administered intravenously before echo to decrease the heart rate to around 150 beats/min to ensure reliable diastolic assessment (21). Echocardiographic measurements were subsequently performed offline in a blinded fashion with EchoPAC SW v201 plugin (GE Healthcare). Anterior and posterior wall thickness was measured at end-diastole from the parasternal short-axis view at the level of the papillary muscle. The average of the anterior and posterior wall was reported. The LA aortic root ratio (LA/Ao) was measured in early ventricular diastole using the first frame after aortic ejection using the right parasternal short-axis view (22). LA volume (LAV) and LA area were calculated as described in humans (23) from a right parasternal long-axis view, using the Simpson’s rule at end-systole and end-diastole. LA function, assessed by LA volumes, was calculated according to atrial function studies in humans (24). Changes in LAV expressed as ejection fraction (LA-EF = [LAVmax LAVmin]/LAVmax) (25). LV end-diastolic diameter and fractional shortening were measured in b-mode from a right parasternal short-axis view. Echocardiographic results for the 4-month time point can be found here (16). Tissues from this previous published cohort were used in the current transcriptomic and metabolomic experiments.

**HEART TISSUE COLLECTION AND PROCESSING.** Hearts were removed, rinsed, and weighed. The aorta was cannulated, and the coronary arteries were cleared by perfusion with cold Krebs-Henseleit buffer. Before tissue fixation, small pieces of the anterior-lateral LV were collected and immediately flash frozen for transcriptomic and metabolomic profiling. For mitochondrial assays, fresh tissue was used immediately following isolation. The hearts were then gravity perfused with 10% formalin at mean arterial pressure (100 mm Hg). Fixed hearts were immersed overnight in 10% formalin and then

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**FIGURE 1** Continued

Differential expression and gene enrichment analysis in hearts subjected to RNAseq 1-month post-banding were compared with sham controls. (A) Volcano plot of differentially expressed RNA transcripts indicating the number of genes increased and decreased in expression (fold change $\geq 1.5$; FDR $\leq 0.3$); (B) heatmap of the 99 genes found to be differentially expressed. Gene ontology (GO) analysis revealing the (C) biological processes and (D) molecular functions of those genes found to be differentially expressed. n = 3 sham and 5 banded. FDR — false discovery rate.
FIGURE 2  Cardio-Metabolic Remodeling Is a Characteristic Feature of Early HFpEF Development

A  Heart: 1-mo. Banded vs 1-mo. Sham

B  1-mo. Banded  1-mo. Sham

tetradecanolate  coenzyme A  thiamin diphosphate
adenosine 5′-triphosphate  NADH
N6-carboxymethyllysine  erucate
mannonate  3-aminoisobutyrate
ornithine  S-methylcysteine sulfoxide
Trans-4-hydroxyproline  N-delta-acetylnorleucine
3,4-dihydroxybutyrate  asparagine
proline  methionine
N-acetylgulosaminylasparagine  Oleoyl ethanolamide
stachydrine  2-hydroxyadipate
uracil  serine
erythronate  pro-hydroxy-pro
erythronate  pro-hydroxy-pro
glycine  p-cresol sulfate
UDP-N-acetylglucosamine
isoceine
valine  leucine
arachidonate  Sphingomyelin
eicosapentaenoate  DPA (22:5n3)
DPA (22:5n3)
docosahexaenoate
N1 N8-acetylperimidine
2-hydroxybutyrate
Gamma-glutamyl-epsilon-lysine
imidazole propionate
tartrate hydroxymalonate
Ascorbic acid
threonate
1-linoleoylglycerol
2-linoleoylglycerol
1-oleoylglycerol
1-palmitoleoylglycerol
1-arachidonoylglycerol
3-hydroxy-3-methylglutarate

C  Metabolite Abundance  Increased  Decreased

D  Metabolite Set Enrichment Overview (p≤0.1)

E  Variable of Importance (VIP) Scores
stored in 70% ethanol for up to 1 week before being processed. The heart was cut on a short-axis plane, starting at the apex and continuing up to the base, and then short-axis sections were cut in half (lateral and septal wall) and embedded in paraffin wax blocks. Tissue sections (5-μm) from 2 different levels from each sample were slide-mounted (AML Laboratories).

**HEART HISTOLOGY.** Paraffin-embedded LV samples fixed at 1 month post-banding were stained with Masson’s Trichrome (Sigma-Aldrich) to determine percentage of fibrosis. Cytoplasm and muscle fibers stain red, whereas collagen (fibrosis) stains blue. The stained slides were imaged at 10× magnification (Nikon Eclipse Ni, NIS Elements software), stitched into 2 × 2 larger images, and analyzed using ImageJ software (National Institutes of Health) with color threshold analysis. The percentage of fibrotic tissue was determined as the collagen-positive tissue of the total stained LV tissue. In order to examine the LV fibrosis gradient, the subendocardium and subepicardium were imaged and quantified separately. All analyses were performed in a blinded fashion. Previously reported fibrosis quantifications for the 4-month time point were used for correlations with metabolomics data in the current study (16).

**RNA SEQUENCING.** Anterior-lateral LV, immediately flash frozen in liquid nitrogen following excision to most accurately capture the cardiac transcriptome as it is in vivo, was subjected to RNAseq analysis. Skeletal muscle isolated from the gastrocnemius was likewise flash frozen in liquid nitrogen. Total RNA was isolated using a fibrous tissue RNA isolation kit (Qiagen). The TrueSeq stranded mRNA library prep kit was used to enrich polyA mRNAs via poly-T mRNA library prep kit was used to enrich polyA mRNAs via poly-T. Tissue sections were dissociated, and to recover chemically diverse metabolites, proteins were then precipitated with methanol (final concentration 80% v/v) under vigorous shaking for 2 minutes (Glen Mills GenoRinder 2000) followed by centrifugation. For quality assurance and control, a pooled matrix sample was generated by taking a small volume of each experimental sample to serve as a technical replicate throughout the data set. Extracted water samples were used direct metagenomic profiling of bacterial communities.

**METABOLIC ANALYSIS.** Anterior LV tissue was immediately flash frozen in liquid nitrogen following excision to most accurately capture the cardiac metabolome as it is in vivo and subjected to unbiased metabolomics. Skeletal muscle isolated from the gastrocnemius was likewise flash frozen in liquid nitrogen. Samples were prepared by Metabolon using their automated MicroLab STAR system (Hamilton Company). First, tissue homogenates were made in water at a ratio of 5 μL per mg of tissue. For quality control, several recovery standards were added before the first step in the extraction process. To remove protein, small molecules bound to protein or trapped in the precipitated protein matrix were dissociated, and to recover chemically diverse metabolites, proteins were then precipitated with methanol (final concentration 80% v/v) under vigorous shaking for 2 minutes (Glen Mills GenoG-Rinder 2000) followed by centrifugation. For quality assurance and control, a pooled matrix sample was generated by taking a small volume of each experimental sample to serve as a technical replicate throughout the data set. Extracted water samples were used direct metagenomic profiling of bacterial communities.
FIGURE 3 Correlations Between the Cardio-Metabolic Profile and Mitochondrial Function Highlight an Energetic and Functional Decline During Early Cardiac Remodeling

Cardiac Mitochondrial Function

A

Sham  Banded

Pyruvate Maltose + ADP  Oligo  FCCP

Complex I OCR (pmol/min/μg protein)

Time (min)

Sham  Banded

Succinate  Rot  + ADP  State 3

Complex II OCR (pmol/min/μg protein)

Time (min)

Correlation of the Cardiac Metabollome with Functional & Structural Endpoints

F

G

H

I

J

Correlation Coefficient

Complex I - State 3

6'-ADP-ribose
2,6-dihydroxybenzoic acid
methionine sulfoxide
N-acetylglucosamine
arachidonic acid
thiamin diphosphate
DHAP
benzoylecgonine
cytochrome
oleoyl-lysoPC(18:1/18:2)
N-acetyl-glucosamine
1,2-dihydroxyacetone-P
1,3-diacetate
2-hydroxybutyrate
N-acetylglutamine
N-methylpyruvate
acetoacetate 20:4(n6)
1-methylhistidine
methylmalonate
spermidine

guanosine 5'-diphosphate ATP
5-methyl-2-deoxycytidine
N-acetylglutamate
1-methylhistamine
1-oleoylglcerol 18:1
homocitrulline
aspartate
phenylacetylglycine
phenol sulfite
trans-4-hydroxyproline
2-lysoPC(18:2)
3,4-dihydroxyproline
3-sulfur-L-ascorbine
threonine
dehydropyrroline
sorbitol
1-lysine 18:2
fumarate
hydroxyproline
ascorbic acid 3-sulfate
5-aminolevulinate

Correlation Coefficient

Complex I - FCCP

Complex II - State 3

hypotaurine
imidazole propionate
N-formylmethionine
3-hydroxy-3-methylglutarate
1-lysoPC(18:2)
N-acetylthreonine
trigonelline
N-methylhistidine
methionine
1-lysine 18:2
5-methyl-2-deoxycytidine
N-acetylglutamate
arachidonic acid 20:4
2-methylbutylcarbamate
5-methylmalonate
N-delta-acetyldiamine
5-dodecanoate 12:1(n7)
arachidonate 20:4
1-arachidonoylglcerol 20:4
methylmalonate
5-methylmalonate
N-acetyllysine
5-methylmalonate

Correlation Coefficient

Complex II - RCR

Lateral LV Fibrosis

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served as process blanks. A cocktail of standards known not to interfere with the measurement of endogenous compounds was spiked into every analyzed sample, allowing instrument performance monitoring and aiding chromatographic alignment.

The extract was divided into fractions for analysis by reverse phase ultraperformance liquid chromatography (UPLC)-tandem mass spectrometry (UPLC-MS/MS) with positive ion mode electrospray ionization (ESI), by reverse phase UPLC-MS/MS with negative ion mode ESI, and by hydrophilic interaction UPLC-MS/MS with negative ion mode ESI. Samples were placed briefly on a TurboVap (Zymark) to remove the organic solvent. All methods used a Waters ACQUITY UPLC and a Thermo Scientific Q-Exactive high resolution/accurate mass spectrometer interfaced with a heated electrospray ionization (HESI-II) source and Orbitrap mass analyzer operated at 35,000 mass resolution. The sample extract was reconstituted in solvents compatible with each MS/MS method. Each reconstitution solvent contained a series of standards at fixed concentrations to ensure injection and chromatographic consistency. One aliquot was analyzed using acidic positive ion conditions, chromatographically optimized for hydrophilic compounds. In this method, the extract was gradient eluted from a C18 column (Waters UPLC BEH C18-2.1 × 100 mm, 1.7 μm) using water and methanol, containing 0.05% perfluoropentanoic acid and 0.1% formic acid. For more hydrophobic compounds, the extract was gradient eluted from the aforementioned C18 column using methanol, acetonitrile, water, 0.05% perfluoropentanoic acid, and 0.01% formic acid. Aliquots analyzed using basic negative ion optimized conditions were gradient eluted from a separate column using methanol and water, containing 6.5 mmol/L ammonium bicarbonate (pH 8). The last aliquot was analyzed via negative ionization following elution from a hydrophilic interaction column (Waters UPLC BEH Amide 2.1 × 150 mm, 1.7 μm) using a gradient consisting of water and acetonitrile with 10 mmol/L ammonium formate (pH 10.8). The MS analysis alternated between MS and data-dependent MS² scans using dynamic exclusion. The scan range covered 70-1000 mass to charge ratios (m/z).

Raw data were extracted, peak-identified, and processed using Metabolon’s proprietary hardware and software. Compounds were identified by comparison to library entries of purified, authenticated standards or recurrent unknown entities, with known retention times/indices (RI), m/z, and chromatographic signatures (including MS/MS spectral data). Biochemical identifications were based on 3 criteria: RI within a narrow RI window of the proposed identification, accurate mass match to the library ± 10 ppm, and the MS/MS forward and reverse scores between experimental data and authentic standards. Proprietary visualization and interpretation software (Metabolon) was used to confirm the consistency of peak identification among the various samples. Library matches for each compound were checked for each sample and corrected, if necessary. Area under the curve was used for peak quantification.

Original scale data (raw area counts) were analyzed using MetaboAnalyst 4.0 software. Metabolites with >50% of the values missing were omitted from the analysis, and missing values were imputed by introducing values with one-fifth of the minimum positive value of each variable. An interquartile range filter was used to identify and remove variables unlikely to be of use when modeling the data. The data were log-transformed (base log-10) and standardized (mean-centered and divided by the standard deviation of each variable). Univariate (eg, volcano plots) and multivariate (eg, principal component analysis) analyses were then performed. Unadjusted P values (p ≤ 0.1) were used to determine significance. Fold changes and P values for all metabolites found to be significantly different are listed in Supplemental Tables 1-8. The raw metabolomics data from individual cats were used for correlative comparison (pairwise analysis; Pearson’s correlations) with mitochondrial functional and tissue fibrosis.

**MITOCHONDRIAL FUNCTION.** Heart and skeletal muscle mitochondria were isolated from fresh tissue and subjected to respiratory function assays using the Seahorse XF96 (Agilent), as described previously (17,26). Briefly, ~100 mg of anterior-lateral LV was washed 5× with cold buffer A (220 mmol/L mannitol, 70 mmol/L sucrose, 5 mmol/L MOPS, 1 mmol/L EDTA;
FIGURE 4 Changes in Metabolism, Cell Signaling, and Inflammation Define the Cardiac Transcriptomic Signature Following the Progression to HFpEF

A

Heart: 4-mo. Banded vs 4-mo. Sham

59 decreased  109 increased

RNAseq FDR (log2)

Log2(FC)

B

4-mo. Banded  4-mo. Sham

Relative Expression

C

Biological Process

regulation of metabolic process

response to abiotic stimulus

negative regulation of signal transduction

vesicle-mediated transport

positive regulation of immune system process

exocytosis

negative regulation of developmental process

negative regulation of cell differentiation

neurotroph activation

granulocyte activation

glycoylation

glycoprotein biosynthetic process

glycoprotein metabolic process

positive regulation of leukocyte migration

D

Molecular Function

signaling receptor binding

molecular transducer activity

receptor ligand activity

receptor regulator activity

cell adhesion molecule binding

growth factor receptor binding

cytokine activity

G protein-coupled receptor binding

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2.5 mmol/L, followed by homogenization using a glass-col homogenizer in 2 ml of buffer A containing 0.2% fatty acid-free bovine serum albumin. Homogenate was then subjected to centrifugation at 800g for 10 minutes followed by supernatant collection and centrifugation at 10,000g for 10 minutes. The pellet containing mitochondria was then resuspended in 1 ml of fresh buffer A (without bovine serum albumin) and centrifuged at 10,000g, with this step repeated once. The washed mitochondrial pellet was then resuspended in 150 μL of respiration buffer (120 mmol/L KCl, 25 mmol/L sucrose, 10 mmol/L HEPES, 1 mmol/L MgCl2, 5 mmol/L KH2PO4; pH 7.2 with potassium hydroxide) and kept on ice.

To determine mitochondrial function, samples were diluted to a concentration of 5 μg (protein) in 50 μL of respiration buffer per well and centrifuged onto XFP96 microplates at 500g for 3 minutes at 4°C. State 3 respiration in response to substrates was measured after injection of pyruvate + malate (5.0 mmol/L + 2.5 mmol/L, final concentrations) or succinate + rotenone (10 mmol/L + 1 μmol/L, final concentrations) to assess complex I and II rates, respectively. Fatty acid oxidation rates were assessed by injection of palmitoyl-L-carnitine (50 μmol/L). The oxygen consumption rates recorded after injection of oligomycin (1 μg/mL), an inhibitor of ATP synthase, served as a measure of state 4 respiration. Following state 4 respiratory measurements, injection of FCCP, a mitochondrial uncoupler, provided electron transport chain (ETC) complex maximal respiratory capacity. Respiratory control ratios, state 3/state 4, were calculated as a measure of the coupling of oxygen consumption to ATP production.

**STATISTICAL ANALYSIS.** Statistical analysis was performed using GraphPad Prism 9 (GraphPad Software), MetaboAnalyst, and R version 3.6. Statistical parameters including the value of n (number of cats), the definition of center, precision (mean), and dispersion (SEM or SD) measures are reported. A P value of ≤0.05 was considered statistically significant. For the metabolomics data set, an unadjusted P value of ≤0.1 was considered statistically significant. For direct comparisons, statistical significance was calculated by unpaired or paired Student’s t-test for between or within group comparisons, respectively. Details on the statistical methods employed for the metabolomics and RNA-sequencing (RNA-seq) data sets can be found within their respective methods sections.

**RESULTS**

**ONE-MONTH AORTIC BANDING REVEALS IMPAIRMENTS IN TRANSCRIPTIONAL AND POST-TRANSCRIPTIONAL REGULATION, MITOCHONDRIAL DYSFUNCTION, AND INFLAMMATION.** Following 1-month aortic banding, LV function was largely unchanged with no significant difference in LV end-diastolic, end-systolic diameter, or fractional shortening (Supplemental Figures 1A to 1F). Modest LA remodeling was observed, with a small, but significant, increase in LA end-systolic volume. No differences in LA EF or LA systolic diameter, as a ratio to the aortic root (LA/Ao; a measure of atrial dilation), were observed (Supplemental Figures 1G to 1L). Significant cardiac hypertrophy 1 month post-banding indicated by echocardiographic measurements of end-diastolic LV wall thickness and increased heart weight to body weight ratio were observed (Supplemental Figures IM to IO). Tissue fibrosis, a common feature of HFpEF cardiac pathophysiology, which, we previously reported, is present 4 months post-aortic banding (16,17), was noted 1 month post-banding (Supplemental Figures 1P to 1R), suggesting fibrotic initiation precedes the development of HFpEF-related functional decline. Collectively, echocardiographic results confirm early cardiac remodeling in the form of LV hypertrophy and interstitial fibrosis in the absence of systolic or diastolic functional decline.

To identify potential molecular pathways and targets contributing early in HFpEF-like disease, we isolated RNA from the LV and performed next-gen RNA-seq. We observed changes in the expression of 99 genes post-banding, with 83 of these down-regulated (fold change [FC] ≥1.5 and FDR ≤0.1) (Figures 1A and 1B). GO analysis of differentially expressed genes identified processes associated with transcriptional regulation (eg, gene expression,
FIGURE 5  Metabolic Analysis of the Heart 4 Months Post-Banding Reveals a Sustained Remodeling of Amino Acid Metabolism and a Greater Reliance on Aerobic Glycolysis for Energy Provision

Metabolic phenotyping of hearts 4 months post-aortic banding compared with age-matched shams: (A) Volcano plot analysis showing metabolites with a fold change (FC) >1.25-fold; P < 0.1; (B) heatmap of the top 50 significantly different metabolites; (C) metabolic network visualization of metabolites in 4-month banded hearts as compared with sham control hearts. Colored circles indicate a statistically significant difference with the size of the circle indicative of the degree of change: 1. gamma-glutamyl amino acids; 2. inosine purines; 3. cytidine pyrimidines; 4. adenosine purines; 5. uracil pyrimidines; 6. long-chain polyunsaturated fatty acids; 7. poly-unsaturated fatty acid species; 8. phospholipids; 9. mono- and dicarboxylic fatty acids; 10. monoacylglycerols; 11. TCA cycle; 12. vitamin B6 metabolism; 13. ascorbate metabolism; 14. Met, Cys, SAM metabolism; 15. lysine metabolism; 16. Leu, Ile, Val metabolism; 17. His metabolism; 18. Ala, Asp metabolism; 19. Gly, Ser, Thr metabolism; 20. Trp metabolism; 21. glycolytic intermediates; 22. amino sugars; (D) enrichment overview of the metabolic pathways significantly affected. Only pathways with a P < 0.1 shown; (E) partial least squares-discriminant analysis variables of importance projection indicative of the metabolites most responsible for group separation. n = 5 sham and 5 banded. AA = amino acids; Carbs = carbohydrates.
nucleic acid and RNA metabolic processes) among the top enriched pathways (Figure 1C). Several epigenetic modifiers were found to be down-regulated in the banded heart (eg, BRD4, KMT2E, KDM5A), suggesting chromatin reorganization early in disease. Molecular function enrichment identified nucleic acid binding proximal promoter DNA-binding transcription activator activity, as well as RNA polymerase II regulation, as system-wide alterations 1 month post-banding (Figure 1D). Several genes involved in the DNA repair pathway (eg, RIF1, MHBOX2, JMY) were down-regulated (Figure 1C), which may underlie the HFpEF hypertrophic phenotype (27).

Several biosynthetic metabolic processes (eg, macromolecule, nucleic acid, and RNA metabolic processes) were highly enriched (Figure 1C); however, interestingly, few genes directly related to metabolism were differentially expressed. We did observe a significant decrease in genes associated with the mitochondrial ETC, such as MT-ND6 and ATP5F1B, which are critical core subunits of complex I and ATP synthase (Figure 1B). Furthermore, CHCHD3, an inner mitochondrial membrane scaffold protein required for mitochondrial cristae structural integrity, was down-regulated, which is reported to impair oxygen consumption and ATP production (28) (Figure 1B). Although intermediary metabolic pathways did not differ at the transcriptional level, several GO terms associated with enzymatic function/activity, including protein kinase and phosphoprotein binding, were significantly enriched (Figure 1D), possibly suggesting alterations in the post-translational regulation of cellular metabolism. Immune response pathways were also identified (eg, T-cell differentiation, activation of immune response) (Figure 1C), suggesting inflammatory signaling may be an early driver of HFpEF progression. Indeed, systemic inflammation brought upon by the numerous associated comorbidities has been recognized in HFpEF through increased microvascular dysfunction, nitrosative stress, and insulin resistance (29). Lastly, GO term analysis indicated cellular responses to nitrogen and organonitrogen compounds, processes resulting in changes in secretion, enzyme production, and gene expression as a result of nitrogen compound stimulus. Recently, differences in arterial and coronary sinus metabolomic profiling of patients undergoing cardiac ablation revealed a large net negative nitrogen balance (30), indicative of active proteolysis. Our results provide a potential transcripational link to account for these metabolite changes, and the aforementioned referenced study provides a small measure of clinical relevance for the model used here.

CARDIOMETABOLIC REMODELING IS AN EARLY SIGNATURE HFpEF DEVELOPMENT. Because critical mitochondrial ETC components were down-regulated, we hypothesized that derangements in myocardial metabolism likely occur early in the disease process. To evaluate changes in metabolism, we quantified the steady-state abundance of myocardial metabolites in 1-month banded cats versus sham controls. Volcano plot analysis revealed that of the 477 metabolites detected, 82 metabolites were significantly different (FC $\geq 1.25; P \leq 0.1$) (Figure 2A). Heatmap analysis of the top 50 metabolites and partial least squares-discriminant analysis clustering indicated group-dependent patterning (Figure 2B, Supplemental Figure 2A). Visualization of the metabolic networks revealed significant changes in the amino acid superfamily, specifically 1-carbon metabolism (methionine, cysteine, s-adenosylmethionine; subfamily 14), branched chain amino acids (BCAAs; leucine, isoleucine, valine; subfamily 16), and glycine, serine, threonine metabolism (subfamily 19) as pathways highly affected early in HFpEF development (Figure 2C, Supplemental Figure 2B). We also observed an increased abundance of BCAAs (eg, isoleucine, valine, leucine) (Figure 2B), and pathway enrichment analysis further highlighted amino acids as significantly changed early in HFpEF (Figure 2D). Interrogation of the metabolites with the highest variable of importance score revealed glycine and the amino acid gamma-glutamyl-epsilon-lysine as defining group separation (Figure 2E). Interestingly, because glycine constitutes 33% of collagen and gamma-glutamyl-epsilon-lysine is involved in covalently cross-linking extracellular matrix (ECM) proteins in fibrotic tissue (31), this may suggest the activation of fibrotic pathways.

Several lipid species, such as polyunsaturated (subfamily 6), mono- and di-carboxylic fatty acids (subfamily 9), and monoacylglycerols (subfamily 10), were found to be elevated in banded hearts. Acyl-carnitines, used for β-oxidation, did not appear to be changed based on metabolic network visualization (Figure 2C). However, z-score analysis indicated that some species of acyl-carnitines trended to increase in abundance in banded hearts (Supplemental Figure 3C), suggesting decreased utilization or increased synthesis. The citric acid cycle was also significantly impacted (Figure 2D), which, when coupled with our transcriptomics data and the accumulation of acyl-carnitines and BCAAs, is suggestive
FIGURE 6 Skeletal Muscle-omics Profiling Indicates an Energetic Deficit in Early HFpEF Progression

A

RNAseq

Skm: 1-mo. Banded vs 1-mo. Sham

15 decreased 6 increased

RNAseq FDR (log 10)

-2 logs(FC) 0 1 2

B

1-mo. Banded 1-mo. Sham

RNA Processing Mitochondrial Dynamics Metabolism

Relative Expression

C

Metabolomics

Skm: 1-mo. Banded vs 1-mo. Sham

7 decreased 12 increased

-10(logp) -5 0 5 10 15

D

1-mo. Banded 1-mo. Sham

Relative Concentration

Creatine mannuronate galactonate

3-hydroxybutyroylglycine

Phosphate propionylcarnitine

1-methyl-4-imidazoleacetic acid

Pyrraline 1-linolenoyl-2-linolenoyl-GPC

2'-O-methylcytidine ararabonate/xylitol

Uridine p-cresol sulfate

3,1-stearyl-2-docosahexaenoic acid-GPC

Cysteine/glycine

Tricosanoyl sphingomyelin

Adenosine 5'-monophosphate (AMP)

Sarcosine ascorbate

Phosphoethanolamine

Guanine (N1) + (N8)-acetylspermidine

Eicosapentaenoate (EPA; 20:5n3)

3-hydroxyisobutyrate

E

Complex I OCR (pmol/min/mg protein)

State 3

State 4

F

Complex I OCR (pmol/min/mg protein)

State 3

State 4

G

Complex I RCR

H

Complex I OCR (pmol/min/mg protein)

State 3

State 4

I

Complex II OCR (pmol/min/mg protein)

State 3

State 4

J

Complex II RCR

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of an overall decrease in oxidative metabolism. Significant enrichment in pathways associated with lipid biosynthesis (eg, cardiolipin, glycerolipid, and phospholipid) (Figure 2D) were also observed, indicating changes, not only in catabolism, but also in anabolism. Lastly, nucleotide metabolites associated with purine and pyrimidine metabolism were differentially impacted (Figure 2C). Collectively, these results highlight metabolic remodeling of amino acid, lipid, and nucleotide metabolism early in HFpEF-like disease progression, and likely point to an impairment in oxidative metabolism.

**METABOLIC REMODELING CORRELATES WITH CARDIAC MITOCHONDRIAL DYSFUNCTION.** Mitochondrial dysfunction is a hallmark characteristic of established HFpEF and HFrEF (32,33); however, whether mitochondrial dysfunction precedes or contributes to cardiac dysfunction in HFpEF remains unclear. With our omics analyses suggesting impaired mitochondrial metabolism, we isolated mitochondria from LV tissue biopsies of sham and 1-month banded cats to interrogate state 3 (substrate-mediated), state 4 (oligomycin inhibition of ETC complex V), and maximal respiratory rates (ATP; P = 0.003; r = 0.888; GTP; P = 0.016; r = 0.805) and complex II RCR (guanosine 5'-diphosphate [GDP]; P = 0.007; r = 0.848) positively correlated with these same nucleotide phosphates (Figures 3G and 3I). A significant negative correlation between ascorbate and complex II RCR (P = 0.048, r = −0.711) (Figure 3I), which may be indicative of oxidative stress and contribute to diminished coupling of mitochondrial oxygen consumption to ATP synthesis, was also observed. Conversely, numerous monoacylglycerols (eg, 1-dihomo-linoleylglycerol) and diacylglycerols (DAGs; eg, oleoyl-linoleoyl-glycerol) negatively correlated (P < 0.05) with mitochondrial functional parameters (Figures 3F, 3G, 3H, and 3I). Collectively, these results, along with the observed transcriptional down-regulation of ETC-associated components (MT-ND6, ATP5F1B, and CHCHD3), indicate that mitochondrial dysfunction and impairments in energy provision are early features in HFpEF development.

We also observed strong negative correlations with LV lateral free wall fibrosis and the abundances of glutamine (P = 0.027; r = −0.765), thiamin diphosphate (P = 0.014; r = −0.811), and metabolites involved in the glutathione cycle (eg, 5-oxoproline; P = 0.021; r = −0.784; gamma-glutamylglutamate; P = 0.005; r = −0.869) (Figure 3J). Pro-hydroxyproline was also significantly increased in banded hearts (Figure 2B). Given that several of these metabolites are in ECM remodeling, it is likely they contribute to initiation of the fibrotic response and resultant tissue fibrosis (34-39). The high correlations between the abundance of various metabolites and phenotypic indicators of cardiac remodeling indicate changes in metabolism are not merely consequential, but likely effectors and drivers of disease.

**TRANSCRIPTIONAL AND METABOLIC REMODELING DURING HFpEF PROGRESSION INDICATE A SHIFT FROM OXIDATIVE TO GLYCOLYTIC METABOLISM.** We examined whether the transcriptional and metabolic changes occurring 1 month post-banding persist following the development of HFpEF characteristics (16,17) and whether this could provide mechanistic

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**Figure 6 Continued**

Transcriptomic and metabolomic analysis of skeletal muscle (SkM) following 1-month aortic banding compared with sham controls. (A) Volcano plot of differentially expressed RNA transcripts indicating the number of genes increased and decreased in expression (FC ≥1.5; FDR ≤0.1); (B) heatmap of differentially expressed genes; (C) volcano plot analysis showing metabolites with a FC ≥1.25-fold, P ≤ 0.1; (D) metabolite heatmap profile. n = 3 sham 1-month and 5 banded 1-month. Abbreviations as in Figures 2 and 3.
Remodeling of Skeletal Muscle 4 Months Post-Banding Is Defined by a Defective Oxidative Metabolism

**FIGURE 7**

**A**

**RNAseq**

Skm: 4-mo. Banded vs 4-mo. Sham

**B**

4-mo. Banded  vs 4-mo. Sham

**C**

**Biological Process**

| (dets) -log10 (p-value) |
|-------------------------|
| 0 | 1 | 2 | 3 | 4 |

**D**

**Metabolomics**

Skm: 4-mo. Banded vs 4-mo. Sham

**E**

**Epigenetics**

**Disease**

**Metabolism**

**Continued on the next page**
insight to disease progression. Using tissue from a historical cohort previously subjected to 4-month aortic banding (16), RNA-seq identified 59 significantly down-regulated and 109 significantly up-regulated genes (FC ≥1.5 and FDR ≤0.1) (Figures 4A and 4B), with GO analysis revealing regulation of metabolic process to have the highest number of associated genes (Figure 4C). Down-regulation of ACADVL, which catalyzes the first step of β-oxidation, an increase in PDK1, which inactivates pyruvate dehydrogenase to reduce pyruvate oxidation, and an increase in TIGAR, which influences the activity of PFK, suggest a metabolic preference for aerobic glycolysis over oxidative metabolism (Figure 4B). Other metabolic genes altered in expression are related to glycerolipid synthesis (PTDSS1), protein glycosylation (B3GNT7), and serine metabolism (VSTM2A). These transcriptional changes indicate a higher degree of metabolic regulation later in HFpEF, as compared with the 1-month time point where down-regulation of ETC machinery was the only observed metabolic-linked change. In line with this idea, numerous pathways linked to cell signaling (eg, cell surface receptor and GPCR signaling pathways, cyclic-nucleotide-mediated signaling, and negative regulation of signal transduction) were found to be significantly enriched in both biological process and molecular function GO analysis (Figures 4C and 4D).

Genes associated with cyclic monophosphates (eg, PDZD3, PKIA, OSTN) were differentially expressed, as were PRKAB2 (AMPK subunit) and PPP2R2C (protein phosphatase 2 subunit). AMPK is a primary sensor of the energy state of a cell, and these data suggest that both transcriptional and post-translational modifications (eg, phosphorylation, dephosphorylation, glycosylation) are controlling intermediary metabolism later in disease.

Additional biological processes associated with cellular remodeling (eg, negative regulation of developmental process and cell differentiation) and the immune response (eg, positive regulation of immune system process, neutrophil and granulocyte activation) were significantly enriched in our data set (Figure 4C). Additional genes of interest found in our data set were SCN10A, a sodium-voltage channel responsible for action potential propagation, and SYNPO2L, an actin-associated protein linked to heart conduction disease, which were significantly increased in expression (Figures 4B and 4C).

Metabolomics at 4 months post-banding revealed a smaller degree of change in fewer metabolites (67 metabolites had a FC ≥1.25; P ≤ 0.1) as compared with differences 1 month after banding (Figures 5A, 5B, and 5C). Citric acid cycle and the Warburg effect were highly enriched due to changes in metabolite abundance (Figure 6B). Z-score analyses support the Warburg effect, the favoring of glycolysis over oxidative phosphorylation under aerobic conditions, because glycolytic intermediates were overall reduced, whereas numerous acyl-carnitine species (eg, palmitoylcarnitine) trended to increase in abundance (Supplemental Figures 3A to 3C). We also observed a global decrease in citric acid cycle intermediates (Figures 5B and 5C), further indicative of reduced oxidative metabolism. Surprising to us, assessment of mitochondrial function 4 months post-banding revealed a normalization of the mitochondrial dysfunction observed at 1 month (Supplemental Figure 4). Interpreting these results with our transcriptomic data sets suggests metabolic dysfunction in HFpEF progression switches from an initial state of dysfunctional mitochondrial machinery (ie, down-regulation of MT-ND6, ATP5F1B, and CHCHD3) to mitochondrial proteomic compensation and alterations in the transcriptional and post-transcriptional regulation of key metabolic nodes (eg, ACADVL, PDK1, AMPK).

Among the most conserved changes observed over the course of HFpEF progression was the extensive metabolic remodeling of the amino acid superfamily (Figures 5B, 5C, and 5D, Supplemental Figure 3C). Unsurprisingly, cardiac fibrosis was more significant 4 months post-banding than at 1 month based on Masson’s trichrome staining (16,17). Amino acids critical in ECM remodeling and collagen formation (eg, glutamine, glycine, serine) were decreased in abundance and displayed negative correlations with tissue fibrosis like our findings 1 month post-banding (Figures 5B, 5C, 5D, and 5E, Supplemental Figures 3D and 4), suggesting sustained changes in amino acid metabolism underlie the temporal accumulation of interstitial fibrosis during HFpEF progression. BCAAs

**Figure 7 Continued**

Transcriptomic and metabolomic analysis of skeletal muscle (Skm) following 4-month aortic banding compared with sham controls. (A) Volcano plot of differentially expressed RNA transcripts indicating the number of genes increased and decreased in expression (FC ≥ 1.5; FDR ≤ 0.1). (B) Heatmap of differentially expressed genes (FC ≥ 1.5; P ≤ 0.1; total of 163 genes). (C) Gene ontology (GO) analysis of the biological processes of the differentially expressed genes; (D) volcano plot analysis showing metabolites with a FC ≥ 1.25-fold, P ≤ 0.1; (E) metabolite heatmap profile. n = 5 sham 4-month and 5 banded 4-month. Abbreviations as in Figures 2 and 3.
were again increased in abundance with high variable
of importance scores (Figures 5B, 5C, 5D, and 5E,
Supplemental Figure 3D). Collectively, the metabolic
changes occurring 1 month and 4 months post-
banding reveal a conserved signature of amino acid
metabolism and a shift in a reliance on aerobic
glycolysis as opposed to oxidative metabolism,
consistent with the pathological heart (18,19).

CARDIOCENTRIC DRIVERS OF SKELETAL MUSCLE
REMODELING IN HFpEF PROVIDES INSIGHT INTO
POTENTIAL MECHANISMS OF SYSTEMIC DISEASE
PROGRESSION. Exertional dyspnea and exercise
tolerance are 2 of the major clinical manifestations
of HFpEF (14). To our knowledge, no studies have
investigated whether a primary cardiac insult con-
tributes to peripheral tissue dysfunction and disease
progression in an animal model of HFpEF. We iso-
lated skeletal muscle from the same felines banded
for 1 month and 4 months to examine skeletal muscle
remodeling at the systems level. RNA-seq of
gastrocnemius 1 month post-banding revealed 21
genes differentially expressed (Figures 6A and 6B),
which were associated with RNA editing and pro-
cessing (ADAR, NARS1, NSRP1, STAU2), de novo pu-
rine biosynthesis (ATIC), and mitochondrial fusion
(MIGA1). Metabolically, skeletal muscle was modestly
affected as only 19 metabolites were signi-
cficantly different. However, we noted a reduction in
creatine and free phosphate accompanied by an in-
crease in AMP (Figures 6C and 6D), suggesting an en-
ergic deficit. An increase in 3-hydroxyisobutyrate, a
catabolic intermediate of valine metabolism and a
strong marker of systemic and muscle insulin resis-
tance (40,41), was also increased in abundance.
Assessment of skeletal muscle mitochondrial func-
tion revealed reduced complex I and complex II state
3 respiratory rates (Figures 6E, 6F, 6G, 6H, 6I, and 6J),
suggesting early cardiac remodeling is suf-
cient to
promote peripheral tissue remodeling.
Transcriptomic analysis 4 months post-banding
revealed 163 differentially expressed genes, with
numerous metabolic processes, development and cell death/autophagy players identified (Figures 7A, 7B, 7C). Of those related to metabolism, several fatty acid-associated genes were highly enriched, with ACADVL specifically down-regulated, consistent with the heart. An increase in numerous acyl-carnitine species (Figures 7D and 7E) and the expression of regulators of glucose metabolism (TIGAR, PFKFB4, GYS2) (Figure 7B) may indicate a fiber-type switch from oxidative to glycolytic, which we have previously reported in this model (17). Consistent with these results, yet unlike our assessment of cardiac mitochondrial function, which was normalized at the 4-month time point, skeletal muscle mitochondrial dysfunction persisted with depressed rates of complex I and complex II respiration (Figure 8). We also observed down-regulation of PRDM8 and up-regulation of SS18—genes associated with chromatin and histone modification—as well as NRXN2, TRIM72, and NOTCH1—genes associated with muscular dystrophy—which provide mechanistic insights and potential targets for HFpEF skeletal muscle disease progression.

**DISCUSSION**

In this study, we used unbiased approaches to define the transcriptional and metabolic changes that occur in a large animal model that recapitulates the cardiopulmonary phenotype of HFpEF. Importantly, this model allowed for the assessment of changes in HFpEF-like disease driven by cardiac dysfunction. Our results reveal distinct changes in cardiac metabolic pathways and transcriptional programs that could shed novel insights into disease progression and provide new therapeutic targets for testing.

**METABOLIC RELIANCE FOR ENERGY PROVISION.** Similar to that observed in HFrEF, an early and sustained shift from oxidative to glycolytic metabolism in heart and skeletal muscle during HFpEF progression was observed; however, unlike HFrEF, mitochondrial function was normalized later in disease. At 1 month post-bandng, mitochondrial respiratory rates were positively correlated with the abundance of high-energy phosphates (eg, ATP), suggesting an overall energetic deficit. There was also a strong negative correlation with DAGs, the accumulation of which occurs under conditions of reduced beta-oxidation (42). DAG levels are increased in the hearts of HFpEF patients concomitant with a decrease in total fatty acids (43). Interestingly, mice with cardiac-specific deletion of diacylglycerol O-acyltransferase 1, the enzyme converting DAGs to triacylglycerols, display cardiac hypertrophy with preserved systolic function (44), a metabolic and physiological phenotype similar to HFpEF. However, whether this mouse model presents with additional features of HFpEF (eg, diastolic dysfunction, impaired pulmonary mechanics) was not investigated.

Mechanistically, mitochondrial dysfunction is likely a result of reduced expression of key subunits required for complex I (MT-ND6), ATP-synthase (ATP5F1B), and efficient electron transfer/super-complex formation (CHCHD2), indicating deficient metabolic machinery at 1 month. At 4 months, these ETC subunits are no longer down-regulated, and mitochondrial function is normalized, likely suggesting compensation and a later issue with fuel selection or utilization. In support of this hypothesis, the down-regulation of ACADVL and up-regulation of TIGAR and PDK1 implies cardiac metabolism likely becomes more regulated by key rate-limiting steps of metabolism. Post-translational modifications of key metabolic enzymes also likely infer metabolic control at this later time point. For example, protein acetylation of ε-amino groups of lysine residues can greatly affect protein function (45,46). Treatment with a pan-HDAC inhibitor (SAHA) decreased the acetylation of proteins involved in mitochondrial fatty acid oxidation and interestingly enhanced acetylation of enzymes involved in the malate-aspartate shuttle and Krebs cycle (17). Succinate dehydrogenase A hyper-acetylation has previously been shown in HF, which results in reduced complex II-driven respiration (47). Collectively, these reports suggest that changes in mitochondrial function and oxidative metabolism are likely driven by post-translational modifications of proteins involved in energy metabolism and the ETC, the regulation of which could be transiently lost during the mitochondrial isolation process and potentially explain why we did not see mitochondrial dysfunction at 4 months. Although previously published studies in animal models and humans at later stages of HFrEF have shown reduced expression of numerous genes and proteins associated with the ETC and cellular metabolism (48-50), our data are more in line with Lai et al (18), who recently reported minimal transcriptional regulation of metabolically linked enzymes 1 month post-aortic banding in mice. Importantly, this progression from deficient machinery to a coordinated substrate preference appears to differ from that of HFrEF, providing a unique feature of HFpEF.

Murine models of HFrEF and human HF patients also display BCAA accumulation because of decreased utilization (51). Decreased oxidation of BCAAs also reduces pyruvate oxidation through the inhibition of pyruvate dehydrogenase via diminished hexosamine
biosynthetic pathway-mediated protein O-GlcNAcylation (52), thereby reducing overall oxidative metabolism. Our results are supportive of these findings because we observed an increase in BCAA abundance at both early and late stages of disease and a significant enrichment in glycosylation biological processes in our transcriptomics data. Heart failure also results in the transcriptional down-regulation and inhibitory phosphorylation of the BCAA catabolic enzyme branch-chain alpha-ketoacid dehydrogenase (BCKD) (51), activation of which increases BCAA catabolism preventing adverse cardiac remodeling in models of pressure overload (51) and ischemia/reperfusion (52). Although we did not observe a transcriptional change in BCKD, it is possible that phosphorylation was altered. Whether increasing BCAA catabolism provides similar beneficial effects in HFpEF has yet to be tested and warrants investigation.

In the skeletal muscle, reduced mitochondrial respiration at 1 month was associated with a reduced energy state and the down-regulation of ATIC, an important enzyme for purine synthesis. Skeletal muscle AMP levels were increased, a potential feedback mechanism to limit purine biosynthesis in the face of a reduced energetic state. Unlike cardiac mitochondria that normalized at 4 months, skeletal muscle mitochondrial function remained depressed and was associated with the down-regulation of ACADVL and up-regulation of PFKFB4 and TIGAR, again suggesting a switch from oxidative to glycolytic machinery, such as what is occurring in the heart.

What is most interesting here is that the cardiocentric development of HFpEF affects peripheral tissue remodeling providing clues toward the mechanisms underlying progressive exercise intolerance in HFpEF patients.

Amino Acids. Amino acids were among the most conserved and broadly affected changes in our hearts at both early and late HFpEF. Of the many changes, the increase in ascorbate at 1 month likely indicates oxidative stress as high levels of ascorbate can deplete GSH and NADPH, potentially contributing to mitochondrial dysfunction. Furthermore, increased oxidative stress is known to yield DNA strand breaks and adducts resulting in DNA damage. Our transcriptomics data revealed a down-regulation of DNA repair genes RIF1, HMBOX1, and JMY, and enrichment of biological processes associated with DNA replication, suggesting changes in these important cellular processes.

Cardiac fibrosis is strongly associated with clinical HFpEF (53). Studies in HFpEF myocardial biopsies display significant fibrosis and increased collagen volume fraction (54,55). Remarkably, a positive correlation between myocardial fibrosis and diastolic dysfunction (56) was observed in HFpEF patients, as was myocardial collagen volume with impaired myocardial relaxation (57) and elevated LV diastolic pressures (54,58). Our metabolomics data showed a strong negative correlation between glutamine and LV fibrosis. Recently, comparison of arterial versus coronary sinus blood metabolomics revealed a significantly greater release of glutamine into the coronary sinus in human failing hearts (30). Given the notion that cardiomyocytes use very little glutamine for anaplerosis (59), it is plausible that the observed decrease in glutamine abundance is due to increased fibroblast utilization, which is required for differentiation and collagen biosynthesis (34-37). Although we observed a decrease in cardiac glutamine, we believe this is due to differences in consumption versus production rates in the vascular and tissue microenvironment, consistent with other cardiac tissue metabolomics studies (19). HFpEF patients also show increased collagen cross-linking and up-regulation of the crosslinker lysyl-oxidase (57); whether our observed increase in the cross-linking metabolite gamma-glutamyl-epsilon-lysine is similarly associated is unknown. Although no studies have investigated the role of these cross-linkers in HFrEF, trials using crosslink breakers showed decreased LV stiffness in HFpEF patients; however, no efficacy in HFrEF was noted with the results indicating increased cardiac dilation (60,61).

Although the transcriptional signatures failed to reveal direct biological processes associated with fibrosis, the large inflammatory response at both time points provides a tangible link to tissue fibrosis. Perivascular fibrosis is a consequence of systemic and cardiac inflammation by infiltrating immune cells producing probiotic proteins (eg, TGFβ) (62). Although this type of fibrosis is observed in both HFrEF and HFpEF, it is significantly greater in HFpEF (63). HFpEF patients also have increased serum levels of monocyte chemoattractant protein 1 (MCP1) (64), which is reduced in HFrEF (65), possibly suggesting a distinct inflammatory-mediator of the fibrotic response in HFpEF. In our data set, T-cell activation and differentiation were enriched, and genetic deletion or antibody blockade of T-cell expression of the glycoprotein OX40 ligand (OX40L) has been shown to reverse tissue fibrosis (66).

Interestingly, glycoprotein biosynthesis was also enriched 4 months post-banding in our model, which can potentially modulate collagen properties. Although the observation of fibrosis in our HFpEF model was not surprising, unique to our results is the concept that early fibrotic initiation may be causal to HFpEF.
development, suggesting therapeutic strategies aimed at targeting the metabolic and transcriptional networks contributing to fibroblast activation and function may improve HFrEF outcomes.

The amino acid 3-hydroxyisobutyrate was found to be elevated at both 1-month and 4-month time points in skeletal muscle and could be important for 2 reasons. First, increased levels of 3-hydroxyisobutyrate are known to reduce insulin sensitivity in oxidative, but not glycolytic, muscle (67). Because HF is associated with skeletal muscle fiber type switching, and our data are suggestive of an increased reliance on glycolytic metabolism, increased 3-hydroxybutyrate likely further exacerbates the energetic deficit indicated by our metabolomics and mitochondrial function results. Second, we observed changes in the expression of PRDM8 and SS18, genes associated with chromatin and histone modification. 3-Hydroxybutyrate has been shown to act as a histone deacetylase inhibitor in muscle (68). This could suggest a metabolic signal to alter, not only histone acetylation, but protein acetylation as well, which we have shown is occurring in HF (17). Targeting of 3-hydroxybutyrate by increasing its oxidation or reducing its synthesis could prove therapeutically beneficial in HFrEF and requires investigation.

**Novel Targets for HFrEF.** The identification of pathways that distinguish HFrEF from HFrEF are critical for the development of targeted therapeutic strategies; however, the specific mechanisms differentiating these 2 forms of HF are still unknown. Electrocardiography revealed an association between abnormal P waves and HFrEF, but not HFrEF (69). P-wave abnormalities can occur due to atrial hypertrophy and could be indicative of atrial fibrillation. Importantly, the prevalence of atrial fibrillation positively correlates with EF%, but, paradoxically, also with worse cardiovascular outcomes (ie, crude mortality, and hospitalizations remained significant in patients with HFrEF, not HFrEF [70]). Interestingly, we identified SYNPO2L and SCN10A, genes involved in electrical conduction and associated with atrial fibrillation (71,72), as increased in expression. These genes could provide novel therapeutic targets due to the prevalence of atrial fibrillation in HFrEF.

Understanding the pathophysioloogy of exercise intolerance, a key clinical feature of HFrEF, is critical for improving patient outcomes. In skeletal muscle biopsies from HFrEF patients, mitochondrial dysfunction was observed and was related to a reduction in oxidative muscle fibers, mitochondrial density, citrate synthase activity, and impaired mitochondrial fusion (73,74). Use of phosphorous magnetic resonance spectroscopy also revealed rapid depletion of high-energy phosphates during exercise and delayed repletion during recovery, which was significantly worse in HFrEF versus HFrEF patients (75). Our data confirm impaired skeletal muscle bioenergetics because mitochondrial function and high-energy phosphate metabolites were negatively affected, likely a result of fiber-type switching in our model (17). Furthermore, the down-regulation of MGA1, a regulator of mitochondrial fusion, provides potential mechanistic insight into dysfunctional mitochondrial dynamics and changes in the mitochondrial network. Skeletal muscle myopathy also occurs in HF, yet are likely intrinsic to HFrEF and not a secondary consequence or an epiphenomenon (76).

Our results revealed altered expression of NRXN2, TRIM72, and NOTCH1, all of which are associated with muscular dystrophy, potentially providing novel targets to address muscle atrophy in HFrEF. Because the current belief is that HFrEF is a systemic disease, our model also suggests the possibility of cardiocentric mechanisms or possibly cardiac-derived circulating factors in mediating skeletal muscle pathogenesis.

**Study Limitations.** A powerful aspect of systems-based approaches is their utility in providing broad, unbiased insights into disease progression. For example, Gjoneska et al (77) studied a mouse model of Alzheimer disease by integrating omics datasets to profile transcriptional and chromatin dynamics across early to late pathology. Surprisingly, our metabolomic and transcriptomic profiling failed to reveal completely distinct signatures of disease progression in heart or skeletal muscle, which we believe is a limitation of the model system. In the progressive banding model, kittens are banded at 2 months of age, with tissue collection occurring at 1 month and 4 months post-banding. By 6 months of age, the cats have undergone significant growth and development, which obfuscates the ability to resolve differences induced by banding (ie, 2 independent variables leading to the inability to effectively resolve either one). This hypothesis is strengthened by our metabolomic and transcriptomic analyses of 4-month sham versus 1-month sham heart and skeletal muscle, which were found to display greater changes than aortic banding (Supplemental Figures 6 and 7). Although comparing the effects of banding across the 2 time points does yield significantly more changes, it is difficult to tease out which of these are due to disease progression or an artifact of aging (Supplemental Figures 8 and 9), which is why here we compared against age-matched controls. Although this is a limitation of our HFrEF model, our data do indicate distinct metabolic and
transcriptional signatures that are consistent and different at early and late states of disease. Furthermore, recent interventional studies performed in this model have shown improved cardiopulmonary function and outcomes (17), suggesting model validity and translational utility.

An additional limitation of our study is the exclusive utilization of male kittens. We certainly acknowledge the importance of understanding potential sex differences as they relate to the molecular pathways of HF, and our current work in the Houser lab is aimed at investigating potential sex differences in our model.

CONCLUSIONS

In summary, the results presented here demonstrate the power of integrating omics and functional datasets to define complex signatures of cardiac disease states and provide distinct metabolic and transcriptional changes in a model of slow-progressive pressure overload, which closely recapitulates the cardiopulmonary phenotype of HFpEF. Importantly, due to the lack of comorbidities associated with this HFpEF model, the pathological remodeling of skeletal muscle suggests a cardiac-centric contribution to systemic HFpEF progression and pathology. Collectively, these findings reveal novel metabolic and transcriptional features which may allow for the discovery of therapeutic targets to treat and potentially reverse HFpEF-like disease.

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PERSPECTIVES

COMPETENCY IN MEDICAL KNOWLEDGE: This study used a recently developed feline aortic banding model that recapitulates many of the multifactorial features of HFpEF and shows that mitochondrial and metabolic dysfunction precede the development of the traditional cardiopulmonary features. The systems-based approach employed here identified distinct and conserved changes in amino acids and energy metabolism suggesting metabolic derangements in these pathways as potentially causative in HFpEF disease progression. Transcriptional signatures of early and late HFpEF in our model revealed transcriptional signatures of processes necessary for cellular remodeling. The identification of these metabolic and transcriptional pathways may provide novel therapeutic approaches to prevent and possibly reverse HFpEF disease progression.

TRANSLATIONAL OUTLOOK: Additional preclinical in vivo studies are needed to determine the targetable metabolic and transcriptional pathways that may have therapeutic potential in HFpEF. For example, the promotion of BCAA oxidation in preclinical models of HFrEF has proven beneficial, warranting investigation in models of HFpEF. Identification of numerous genes with potential impacts on arrhythmias, skeletal muscle atrophy, epigenetics, and cellular metabolism could provide targets for therapeutic intervention to treat HFpEF.
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**KEY WORDS** heart failure, mitochondria, metabolomics, preserved ejection fraction, systems biology, transcriptomics

**APPENDIX** For supplemental figures and tables, please see the online version of this paper.