BRIEF COMMUNICATION

Mouse Model of Heart Failure With Preserved Ejection Fraction Driven by Hyperlipidemia and Enhanced Cardiac Low-Density Lipoprotein Receptor Expression

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BACKGROUND: The pathways of diastolic dysfunction and heart failure with preserved ejection fraction driven by lipotoxicity with metabolic syndrome are incompletely understood. Thus, there is an urgent need for animal models that accurately mimic the metabolic and cardiovascular phenotypes of this phenogroup for mechanistic studies.

METHODS AND RESULTS: Hyperlipidemia was induced in WT-129 mice by 4 weeks of biweekly poloxamer-407 intraperitoneal injections with or without a single intravenous injection of adeno-associated virus 9–cardiac troponin T–low-density lipoprotein receptor (n=31), or single intravenous injection with adeno-associated virus 9–cardiac troponin T–low-density lipoprotein receptor alone (n=10). Treatment groups were compared with untreated or placebo controls (n=37). Echocardiography, blood pressure, whole-body plethysmography, ECG telemetry, activity wheel monitoring, and biochemical and histological changes were assessed at 4 to 8 weeks. At 4 weeks, double treatment conferred diastolic dysfunction, preserved ejection fraction, and increased left ventricular wall thickness. Blood pressure and whole-body plethysmography results were normal, but respiration decreased at 8 weeks (P<0.01). ECG and activity wheel monitoring, respectively, indicated heart block and decreased exercise activity (P<0.001). Double treatment promoted elevated myocardial lipids including total cholesterol, fibrosis, increased wet/dry lung (P<0.001) and heart weight/body weight (P<0.05). Xanthelasma, ascites, and cardiac ischemia were evident in double and single (p407) groups. Sudden death occurred between 6 and 12 weeks in double and single (p407) treatment groups.

CONCLUSIONS: We present a novel model of heart failure with preserved ejection fraction driven by dyslipidemia where mice acquire diastolic dysfunction, arrhythmia, cardiac hypertrophy, fibrosis, pulmonary congestion, exercise intolerance, and preserved ejection fraction in the absence of obesity, hypertension, kidney disease, or diabetes. The model can be applied to dissect pathways of metabolic syndrome that drive diastolic dysfunction in this lipotoxicity-mediated heart failure with preserved ejection fraction phenogroup mimic.

Key Words: heart failure ■ hyperlipidemias ■ mice ■ stroke volume

With a high and increasing incidence and few effective treatment options, heart failure with preserved ejection fraction (HFpEF) has been designated the greatest unmet medical need in cardiovascular disease.1 HFpEF accounts for ≥50% of HF diagnoses worldwide, and its complex cause reflects a multisystem as opposed to monolithic disease.1 Animal models, which are required to dissect the underlying pathophysiology and identify therapeutic targets for HFpEF, must be similarly multifactorial and reflect
individual human classes of the disease. Tied with hypertension as the third most important independent risk factor after age and sex, obesity-driven HfPEF represents a distinct pathophysiological phenotype wherein direct cardiac lipotoxicity may confer cardiac dysfunction. The precise mechanisms involved in diastolic dysfunction driven by excess accumulation of myocardial lipids are unclear and animal models are few. Lipoprotein lipase (LPL) is decreased (Figure 1A) and low-density lipoprotein receptor (LDLR) increased in patients with HfPEF. Here we describe a novel, reproducible mouse model of rapid-onset HfPEF wherein hyperlipidemia conferred by pharmacologic inhibition of LPL by poloxamer-407 (P407), a selective LPL inhibitor, and cardiac-specific overexpression of the LDLR drive a full spectrum of classic symptoms that are independent of hypertension, diabetes, or kidney disease.

METHODS

The data that support the findings of this study are available from the corresponding author upon reasonable request. All procedures involving animals were approved by the Institutional Animal Care and Use Committee at the University of Miami, conforming to National Institutes of Health guidelines (IACUC protocol 20–118). Wild type (WT) mice on 129/J background were subjected to biweekly intraperitoneal injections of poloxamer-407 with or without a single intravenous injection of $1 \times 10^{12}$ VGS adenovirus 9–cardiac troponin T–LDLR to direct human LDLR overexpression selectively to the heart. Controls received single, placebo, or no treatment. At 4 weeks and 8 weeks post-treatment, echocardiography, whole body plethysmography, tail-cuff blood pressure recording, etc.

![Figure 1. Heart failure with preserved ejection fraction features with normal blood pressure were shown by the double treatment mice.](image-url)

A. Representative graph comparing lipoprotein lipase (LPL) gene expression in endomyocardial biopsies obtained from healthy human controls (n=24), patients with heart failure with preserved ejection fraction (HfPEF, n=41), and heart failure with reduced ejection fraction (HFrEF, n=30) shows decreased LPL in patients with HfPEF (RNA sequencing data from Hahn et al). B. Representative echocardiography images showing diastolic dysfunction predominantly in the double-treatment group (n=31) as evidenced by prolonged isovolumic relaxation time (IVRT), decreased MV E/A and MV E/E' compared with untreated (n=26), 407+Adenovirus9–cardiac troponin T–Luciferase (AAV9- cTnT- Luciferase, n=11) and Adeno-associatedvirus9–cardiac troponin T–LDLR receptor (AAV9- cTnT-LDLR, n=10) groups. C. Echocardiography revealed preserved ejection fraction (EF%) in all treatment groups. D. Representative graphs showing features of left ventricular hypertrophy; increased left ventricular anterior wall thickness (LVAW), heart weight/body weight (HW/BW), and decreased left ventricular internal diameter (LVID) and volume. IVS indicates interventricular septum. E. Representative graphs reveal normal diastolic and systolic blood pressures (BP) in each group. F. Representative graphs showing daily running activity decreased in all groups; the poloxamer-407+AAV9–cTnT–Luciferase (n=6) group showed the lowest activity (P<0.0001) followed by the double-treatment (n=3; P<0.001) and AAV9–cTnT–LDLR (n=5; P<0.01). For all analyses (except total cholesterol and triglyceride analyses comparing 2 groups by t test), 2-way ANOVA with Tukey–Kramer post hoc correction was used. In all figure panels, means±SD was used. *P<0.05; **P<0.01; ***P<0.001; and ****P<0.0001.
voluntary wheel exercise, ECG telemetry, and histology were performed. Cardiac morphology and function was assessed using the Vevo2100 imaging system (Visual Sonics, Toronto, ON, Canada) with a MS400 linear array transducer (as in our previous work5). Blood pressure was recorded using a noninvasive tail-cuff method (BP-2000 Series II, Visitech Systems, Apex, NC). Mice were trained for 4 consecutive days and data were collected on day 5 (as in our previous work6). Respiratory function was assessed using a Buxco small animal whole body plethysmography system and FinePoint software (Data Science International, New Brighton, MN). Mice were acclimatized for 3 consecutive days and data were collected on day 4 (as in our previous work6). Activity wheel monitoring was assessed using Lafayette Instrument Activity Wheel Monitor Model 86056 (Lafayette Instrument, Lafayette, IN) (as in our previous work7). Mice were acclimatized for 2 days and data were collected over 5 days. Electrocardiogram data were obtained from radiotelemetry HD-X11 ECG devices (Data Science International, St. Paul, MN).

Statistical Analysis
Two-way ANOVA with Tukey–Kramer post hoc correction was used for all 4-group experiments. Student t test was used to analyze results from 2-group experiments. For survival curve analysis, the Log-rank test was used. A significance level of P<0.05 was used and all tests are 2-sided. All data are presented as mean±SD. GraphPad Prism 9 software was used for all the analyses and graph generations.

RESULTS
Elevated myocardial lipids is a prominent feature of patients with HFpEF3 and the gene for LPL in cardiac biopsies is decreased compared with HFrEF or healthy controls.4 Poloxamer-407 induces hyperlipidemia by blocking LPL and increasing plasma triglycerides and LDL cholesterol.5 Our group recently reported that LDLR levels are increased in autopsy specimens of hearts from sudden cardiac death victims with HFpEF and diabetes.5 We hypothesized that hyperlipidemia driven by poloxamer-407 and overexpression of cardiac LDLR in mice mimics the subset of human HFpEF, where metabolic syndrome and diastolic dysfunction are driven primarily by lipotoxicity.3 Quantitative polymerase chain reaction, Western blot, and immunostaining confirmed >3-fold overexpression of LDLR in hearts 4 weeks after injection of adeno-associatedvirus9–cardiac troponin T–LDLR (data not shown). Echocardiography of double-treatment mice at 4 weeks revealed diastolic dysfunction as evidenced by prolonged isovolumic relaxation time, decreased MV E/A and MV E/E′ (Figure 1B) and preserved EF (Figure 1C). Echocardiography also confirmed decreased left ventricular internal diameter (LVID;d and LVID;s), and increased interventricular septum and left ventricular anterior wall (LVAW) thickness (Figure 1D); the latter parameters were also increased in single-treatment groups. Blood pressure remained unchanged by treatments (Figure 1E), exercise intolerance was confirmed by decreased daily running distance in all groups (Figure 1F), and all mice remained nonobese. Telemetry/ECG revealed paroxysmal heart block coincident with lipid accumulation in the atrio-ventricular junction (Figure 2A), also a feature of human HFpEF.9 Picrosirius Red staining confirmed the presence of fibrosis (Figure 2B). Galectin-3 is an early indicator of cardiac fibrosis and ventricular remodeling in patients with HF.10 Galectin-3 was significantly elevated in the hearts of double-treatment mice compared with single-treatment or control groups (Figure 2B). Wheat germ agglutinin staining revealed significantly increased cell size in both double- and single-treatment groups, confirming cardiac hypertrophy (Figure 2C), and pulmonary congestion, a key manifestation of the clinical syndrome of HF, was indicated by increased lung wet/dry ratio (Figure 2D). Myocardial tissue from double-treatment mice contained increased cholesterol and decreased triglycerides, consistent with increased LDL cholesterol uptake (Figure 2E). Total cholesterol was elevated in skeletal muscle and liver; triglycerides were reduced in liver while plasma glucose, blood urea nitrogen, and creatinine remained unchanged (data not shown). Hyperlipidemic mice succumbed to sudden death between 6 and 12 weeks of age (Figure 3A) irrespective of sex, and mice that survived 8 weeks developed more prominent phenotypes while preserving EF (data not shown). With an incidence of ≈35%, sudden death is a leading cause of HFpEF death.1 At 8 weeks, decreased respiration rate was observed in the double-treatment group only (Figure 3B). Necropsy of double-treatment mice revealed pulmonary congestion and ascites, and cardiac ischemia (Figure 3C). Pulmonary hypertension and right ventricular dysfunction are often associated with HFpEF accompanied by hepatic venous congestion and hypoxemia.11 The observed ascites is most likely a consequence of worsening HF and hepatic venous congestion. Unilateral and bilateral skin discoloration and deposits surrounding the eyelids suggest xanthelasmas, a consequence of hyperlipidemia.12 In addition, at 8 weeks, free myocardial cholesterol was significantly increased in this group (Figure 3D).

DISCUSSION
Together, the results provide a unique model of early-onset, rapidly progressing, hyperlipidemia-driven HFpEF where treated mice display diastolic dysfunction, arrhythmia, cardiac hypertrophy and fibrosis, pulmonary congestion and exercise intolerance, with
preserved EF, absence of hypertension and high incidence of sudden death, independent of kidney disease or diabetes. The model provides an opportunity to dissect the roles of hyperlipidemia driven by suppressed LPL and increased LDLR, both features of human HFpEF in the absence of hypertension or obesity and comorbidities related to diabetes and/or chronic kidney disease. The dramatic response of mice to the double-treatment regimen suggests that LDLR overexpression may exert pathophysiological effects beyond those restricted to LDL/cholesterol transport. In terms of phenotypes, our model bears similarity to a recently presented model of mice exposed simultaneously to high-fat diet and N
\textsuperscript{ω}-nitro-L-arginine methyl ester to simulate metabolic stress and hypertension wherein the mice develop HFpEF over a period of 5 to 15 weeks.\textsuperscript{13} Importantly, this work identified a defective unfolded protein response driven by inducible nitric oxide synthase as central to the HFpEF phenotype. Key differences between our model and the N
\textsuperscript{ω}-nitro-L-arginine methyl ester model include more rapid onset, absence of hypertension, presence of arrhythmias, higher and earlier incidence of sudden death, and absence of the potentially confounding effects of N
\textsuperscript{ω}-nitro-L-arginine methyl ester that do not mimic the pathophysiology of hypertension in human HFpEF (reviewed in\textsuperscript{1}). Also we observed no significant differences between sexes in treated mice, consistent with the possibility that sex-related differences in adipose deposition contribute to HFpEF in obese humans. We do not expect significant coronary artery disease as a factor in sudden death in this model because aortic atherosclerotic lesions in P-407-treated mice require >4 months to mature to mimic human-like pathological features.\textsuperscript{2} Our model represents a unique subtype of HFpEF and presents an opportunity to dissect the pathways of hyperlipidemia and associated lipotoxicity in disease progression in the absence of hypertension, chronic kidney disease, diabetes, or obesity. Both models provide opportunities to further
our understanding of the molecular pathology of metabolic syndrome and hyperlipidemia-induced diastolic dysfunction and HFpEF.

**CONCLUSIONS**

We present a novel murine model wherein diastolic dysfunction and classic symptoms of HFpEF are induced by pharmacological inhibition of combined with cardiac-specific overexpression of LDLR. The absence of obesity, hypertension, or diabetes suggests that lipotoxicity alone is sufficient to drive the phenotype, and this may represent a novel, more restricted HFpEF phenogroup related to but distinct from those of obesity with or without hypertension. The model may be applied to dissect pathways and mechanisms whereby lipotoxicity drives HFpEF and is especially relevant to high-incidence arrhythmia and sudden death.

**ARTICLE INFORMATION**

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**Disclosures**

JH is listed as a co-inventor on patents on GHRH analogs, which were assigned to the University of Miami and Veterans Affairs Department. JH previously owned equity in Biscayne Pharmaceuticals, licensee of intellectual property used in this study. Biscayne Pharmaceuticals did not provide funding for this study. JH reported having a patent for cardiac cell-based therapy. He holds equity in Vestlon Inc. and maintains a professional relationship with Vestlon Inc. as a consultant and member of the Board of Directors and Scientific Advisory Board. JH is the Chief Scientific Officer, a compensated consultant and advisory board member for Longeveron, and holds equity in Longeveron. JH is also the co-inventor of intellectual property licensed to Longeveron. Longeveron LLC and Vestlon Inc. did not participate in funding this work. JH’s relationships are disclosed to the University of Miami, and a management plan is in place.

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